MORC3, a novel MIWI2 association partner, is an epigenetic regulator of piRNA-dependent transposon silencing.

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Abstract

The PIWI (P-element-induced wimpy testis)-interacting-RNA (piRNA) pathway plays a crucial role in the repression of TE (transposable element) expression via de novo DNA methylation in mouse embryonic male germ cells. Various proteins, including MIWI2 are involved in the process. TE silencing is ensured by piRNA-guided MIWI2 that recruits some effector proteins of the DNA methylation machinery to TE regions. However, the molecular mechanism underlying the methylation is complex and has not been fully elucidated. Here, we identified MORC3 as a novel associating partner of MIWI2 and also a nuclear effector of retrotransposon silencing via piRNA-dependent de novo DNA methylation in embryonic testis. Moreover, we show that MORC3 is important for transcription of piRNA precursors and subsequently affects piRNA production. Thus, we provide the first mechanistic insights into the role of this effector protein in the first stage of piRNA biogenesis in embryonic TE silencing mechanism.

Introduction

TEs (transposable elements) constitute about 40% of the mammalian genome \(^1\). Although a few copies of TEs can move in the genome, most of them are incapable of transposing. Some genomic mutations caused by transposon activity may play an important role in genome evolution and contribute to promoting biological diversity. However, the expression of transposons is potentially harmful, because it can disrupt genomic stability and induce damage to germlines, resulting in infertility \(^2\). Among the TEs, LINE1 (long interspersed nuclear element-1) retrotransposons are the most
abundant in the mammalian genome and thousands of LINE1 genes are postulated to be intact and active in the mouse. Retrotransposons including LINE1 are usually silenced by DNA methylation and histone modifications of their promoter regions.

PIWI (P-element-induced wimpy testis) was originally discovered as an essential protein for gametogenesis in *Drosophila*. PIWI family proteins, evolutionally conserved in diverse organisms from *Caenorhabditis elegans* to mammals, are required for germ cell development and piRNA (PIWI-interacting RNA) biogenesis. Embryonic piRNAs are germ cell-specific small noncoding RNAs of 25–31 nucleotides that bind to PIWI proteins and contribute to the silencing of TEs in the germlines. Notably, they are mostly derived from retrotransposons. In mouse gonocytes, MILI (PIWIL2) and MIWI2 (PIWIL4) are essential for spermatogenesis and play critical functions in piRNA biogenesis and *de novo* DNA methylation at the retrotransposon region, respectively.

In addition to PIWI proteins, other proteins like the TDRD proteins (Tudor domain containing proteins), RNA helicases such as MVH and MOV10L1, MITOPLD endonuclease, and PNLDC1 exonuclease have been reported to be involved in the piRNA pathway. The embryonic mouse germline piRNA biogenesis can be divided into primary and secondary pathways. Primary piRNAs are generated from long single-stranded RNAs, which are transcribed from piRNA clusters in the genome. First, long capped and polyadenylated precursors are transcribed. These long piRNA precursor transcripts are then fragmented by MITOPLD to phased precursor piRNAs (pre-piRNAs) that carry uridine at 5’ end (1st U) in 5’→3’ direction. After the pre-piRNAs bind to MILI, the 3’ ends of pre-piRNA are trimmed by PNLDC1. In the secondary piRNA biogenesis pathway, MILI bound piRNA catalyzes a target transcript.
with 10 nt complementarity to create the piRNA by its slicing activity and generates a new secondary piRNA with a prominent bias for carrying adenine at position 10 (10th A) 30. After the secondary piRNA is loaded to MIWI2, the piRNA-bound MIWI2 can be transported into the nucleus and recruit some transcriptional silencing related factors to mediate the DNA methylation of the target retrotransposon loci 31,32.

MORC (originally named from the phenotype of microrchidia) is a nuclear protein family highly conserved in prokaryotic and eukaryotic cells, and the proteins of this family play an important role in gene regulation in multiple organisms 33. In Arabidopsis, AtMORC1 and AtMORC6 play a critical role in TE repression with heterochromatin formation at retrotransposon regions, and the process is independent of DNA methylation and repressive histone modification 34,35. In mice, there are six members of the MORC family, namely, MORC1, MORC2a, MORC2b, MORC3, MORC4, and SmcHD1 36. MORC1, an essential protein in spermatogenesis, has emerged as a novel factor in the silencing of retrotransposons (LINE1 and IAP (intracisternal A-particle)) in male germ cells 37,38. MORC1 plays an epigenetic function and is essential for the establishment of DNA methylation marks at retrotransposon genes in a piRNA pathway-independent manner 39,40. Meanwhile, MORC2b is essential for meiosis and fertility in both sexes. Loss of MORC2b leads to misexpression of genes involved in spermatogenesis but does not cause de-repression of retrotransposons in testes 41.

Transposon silencing in mouse embryonic male germ cells is ensured by MIWI2, which is guided to the appropriate gene locus through base-pairing between piRNAs and nascent RNAs transcribed from genomic TE regions and recruits some effector proteins to the regions. However, the identity of all the PIWI related effector proteins in
the piRNA pathway and the molecular mechanisms involved in this process is currently not known. In this study, we identified and characterized the MORC3 protein as an associating partner of MIWI2 in mouse embryonic testis. We showed that MORC3 is important for transcription of piRNA precursors in the first step of piRNA biogenesis and is required for de novo DNA methylation of a retrotransposon in a piRNA pathway-dependent manner.

Results

MORC3 as a component of the MIWI2 protein complex

To characterize the molecular mechanism of retrotransposon silencing via the piRNA pathway, we searched for the proteins that associate with MIWI2 by immunoprecipitation using a ZF-MIWI2 (zinc finer fused MIWI2) Tg mouse. We previously produced two kinds of Tg mice bearing the ZF proteins which target to the common regulatory region of type A LINE1 elements. One expressed Flag-tagged ZF-MIWI2 fusion protein and the other expressed only Flag-tagged ZF protein in testis.

Immunoprecipitated proteins from the ZF-MIWI2 Tg testes of 21-day-old mice were subjected to SDS–PAGE (sodium dodecyl sulfate-polyacrylamide gel) electrophoresis and silver staining. Comparison of the ZF only and ZF-MIWI2 Tg testes samples showed that one band at around 130 kDa was present in the ZF-MIWI2 Tg but not in the ZF Tg testis samples. LC-MS/MS analysis revealed that the identity of the fragments of 26-peptide of the band was similar to those of the MORC3 protein (303/942 amino acids; 32% coverage) (Fig.1A). Immunoprecipitation and western blotting analysis with the anti-Flag antibody using ZF-MIWI2 Tg testes confirmed the presence...
of MORC3 in the ZF-MIWI2 complex (Fig.1B). To analyze whether the interaction between MIWI2 and MORC3 is direct, we overexpressed PA-tagged MIWI2 and Flag-tagged MORC3 in the HEK 293T human embryonic kidney cell line, in which no piRNA pathway proteins are expressed. As shown in Fig.1C, the association between MIWI2 and MORC3 in the transfected 293T cells strongly suggested a direct interaction between them.

Next, by co-immunostaining, we examined the localization and the possible interaction of MORC3 with MIWI2 and MILI in embryonic testis. Both piP-body and pi-body are important organelles for piRNA production and contain MIWI2 and MILI, respectively. MORC3 was expressed together with MIWI2 in not only the nucleus but also in germinal granules, known as the piP-bodies. However, the cytoplasmic signal of MILI-containing granules or the pi-bodies was not merged with the staining signal of MORC3. Taken together, we conclude that MORC3 is an interaction partner of MIWI2 but not of MILI in the embryonic testis (Fig.1D).

**Subfertility associated with the loss of MORC3**

To examine the function of MORC3 in male germ cells, we generated the MORC3 mutant mouse using the Cre-loxP system. *Tnap* (Tissue non-specific alkaline phosphatase)-Cre Tg mice, in which Cre is efficiently and specifically expressed in germ cells, have been frequently utilized. We checked whether the allele was deleted by PCR (polymerase chain reaction) using the sorted male germ cells of the *Morc3* conditional homozygous and heterozygous mutant mice (Day 12) (supplementary Fig.1A). The male germ cells were purified using the anti-EpCAM antibody, and the purification levels were confirmed by the high methylation status (>90%) of the DMR.
(differentially methylated region) of a paternally imprinted gene, DMR-H19. As shown
in supplementary Fig.1B and 1C, loss of MORC3 expression was ensured both in the
testis of E14.5, a period before the beginning of de novo DNA methylation, and the testis
of 30-day-old mice (Day 30) by immunostaining.

The size, weight, and HE staining of the testes of the MORC3 mutant adult mice
were similar to those of wild type mice. In addition, the sperm numbers in the caudal
epididymis were almost the same (Fig.2A and B). However, the pregnancy rate of the
female mice crossed with the mutant mice (25%) was significantly lower than that of
female mice crossed with the wild mice (85%) (Fig. 2C and supplementary Fig.1D).
Although the litter size of the pregnant female mice was not significantly different
(p=0.08 by t test), the results indicated that the loss of MORC3 leads to subfertility of
the male mice from the standpoint of pregnancy rate.

De-repression of retrotransposons in the MORC3 mutant mice

Based on the previous reports on the effect of the other MORC proteins on
transposon silencing, we examined the expression of LINE1 and IAP genes in the testes
of 5-month-old mice by northern blot analysis (Fig.3A). LINE1 retrotransposons are
divided based on their sequences into several distinct types, such as type A and type TF,
both of which have similar and unique sequences in their coding and 5’ noncoding
regions, respectively. The expression of type A and TF LINE1 transcripts were
significantly upregulated, while that of IAP remained repressed in the MORC3 mutant
testes. In contrast, the expression of LINE1 transcripts was lower in the MORC3
mutant mice than that of the MILI-null mouse, in which almost no piRNAs were
produced. Such a de-repression of LINE1 transcripts was scarcely observed in the
3-week-old MORC3 mutant mouse.

Next, DNA methylation levels of the LINE1 and IAP retrotransposons in the spermatocytes, before the pachytene stage, of the 12-day-old MORC3 mutant mice were examined by bisulfite sequencing (Fig. 3B and C). A significant reduction in the CpG methylation levels of type A and TF LINE1 genes was observed in the MORC3 mutant male germ cells, compared to the high DNA methylation of the control germ cells. Meanwhile, DNA methylation of the IAP gene was not impaired by MORC3 mutation and, correspondingly, no significant difference in IAP transcript levels was observed (Fig. 3A). The similarity in the phenotypes, i.e., low DNA methylation at LINE1 regions found in the MORC3 mutant, was observed in not only the postnatal testicular germ cells but also in the sperm (Fig. 3D and E). However, the degree of DNA methylation at LINE1 regions in the MORC3 mutant sperm was slightly higher than that in the male germ cells. These data suggested that some germ cells that have quite low methylation levels at LINE1 regions would have been eliminated during the maturation process of spermatogenesis in the MORC3 mutant mice by mechanisms yet to be elucidated.

Role of MORC3 in piRNA biogenesis

DNA methylation marks on LINE1 repeat elements are linked to the nuclear events of the piRNA-dependent DNA methylation pathway, in which MIWI2 is involved. To assess whether MORC3 plays any role in piRNA biogenesis, we purified the total small RNAs from wild type as well as the MORC3 mutant embryonic testes (E16.5) and subjected them to deep-sequencing. After trimming of miRNAs, rRNAs, and tRNAs from both total small RNA libraries using the CLC Genomics Workbench, the reads were mapped to the mouse genome (mm10, UCSC) and normalized to the read-depth of
each library. Comparison of small RNA size distribution profiles among their libraries revealed that the number of 25–31 nt small RNAs (corresponding to the length of piRNAs) was significantly reduced in the MORC3 mutant testes (Fig.4A and B). The expression of miRNAs, used as an internal control, was almost equal in them (supplementary Fig.2A) and the genomic annotations for 25–31 nt small RNAs using the control and MORC3 mutant libraries indicated that much of reads were originated from retrotransposon-related sequences.

Analysis of piRNAs of the MORC3 mutant testes revealed that a significant decrease was largely contributed to the reduction of small RNAs derived from LINE elements (LINE), LTR elements except for IAP (LTR (non-IAP)), and SINE elements (SINE) (Fig.4C). To evaluate the expression of piRNAs corresponding to LINE1 in detail, we mapped 25–31 nt small RNAs to type A and TF LINE1. While both sense and antisense piRNAs levels were decreased to about half, the proportion of 1st U sense and 10th A antisense piRNAs was not altered in the MORC3 mutant mice compared to the control mice (Fig. 4D and E).

Reduction in MIWI2 and MILI loaded piRNAs in the MORC3 mutants

To analyze the function of MORC3 in more detail, we examined the piRNAs bound to MIWI2 or MILI in embryonic testes. We performed immunoprecipitation of the germ cells of wild type and the MORC3 mutant embryonic testes (E16.5) using the anti-MIWI2 or -MILI antibodies and subjected them to deep sequencing. Western blot analysis revealed that the amount of MIWI2 and MILI proteins immunoprecipitated with the corresponding antibodies were almost the same in the control and the mutant testes (supplementary Fig.2B). However, the levels of MIWI2- and MILI-bound piRNAs
were reduced (34% and 24% reduction, respectively) in the MORC3 mutants, compared to the control wild type mice (Fig.5A, and supplementary Fig.2C). Among the MIWI2- or MILI-loaded piRNAs, both the sense and antisense repeat piRNAs mapping to LINE, LTR (non-IAP), and SINE elements were dramatically reduced in the MORC3 mutants. However, the level of piRNAs mapping to IAP (LTR (IAP)) was unaffected in the MORC3 mutants (Fig.5B and C). The proportion of 1st U sense and 10th A antisense piRNAs was not altered in any category (supplementary Fig.2D).

Since the entry of MIWI2 into the nucleus is dependent on its association with piRNAs, we examined the localization of MIWI2 in the MORC3 mutant embryonic testes by immunostaining. MIWI2 was still present in the nucleus but the amount of cytoplasmic MIWI2 was higher in the MORC3 mutant testes (E16.5) (Fig.5D and supplementary Fig.2E). This phenomenon seems to be corresponding with the reduction of MIWI2-bound piRNAs.

Regulation of transcription of embryonic piRNA precursor by MORC3

piRNA loci are defined as the piRNA clusters that are regions of the genome mapped with high-density piRNA sequences. We chose some major piRNA clusters in Chr 7, 8, and 10 in the genome, and analyzed piRNA precursor transcript abundance using the wild type and MORC3 mutant embryonic testes by RT-qPCR (quantitative reverse transcription polymerase chain reaction) (Fig.6A and B). We carefully designed primer sets with unique sequence to identify the transcripts of retrotransposons in piRNA clusters in the genome. For the Chr 7 (1) cluster, Chr 7 (1)-2 and 3 primer sets were located around IAP sequence, and Chr 7 (1)-1, 4, and 5 primer sets were located at a region other than the retrotransposon sequence. Likewise, in Chr 10 cluster, Chr 10-3,
4, 5, and 6 primer sets were located around the retrotransposon sequence whereas Chr 10-1 and 2 primers were not.

The transcript levels of the regions close to LINE, LTR, and SINE sequences were significantly decreased in the MORC3 mutant testes. However, in regions close to IAP the transcript levels were not different between the control and the MORC3 mutant testes. Essentially, similar patterns were also found in Chr 7 (2) and 8 clusters (supplementary Fig.3). These data show that MORC3 has some retrotransposon class-specific effect to regulate the transcription of piRNA precursor from embryonic piRNA clusters.

**Discussion**

Several reports have shown the importance of the role played by MORC family proteins in retrotransposon expression. Homologues of Morc are required for transposon silencing in *Arabidopsis* and *C. elegans*. Furthermore, Morc1, a Morc mouse homologue, is the epigenetic regulator of piRNA-independent transposon gene silencing 34,35,39,40. Results of the current study identified MORC3 as a component of the MIWI2 protein complex and a regulator of transposon repression in male germ cells. MORC3 is essential for repression of retrotransposon via *de novo* DNA methylation involved in the piRNA pathway. Herein, we show that MORC3 is a new player in piRNA biogenesis and acts as an epigenetic regulator of transposon gene silencing via the piRNA pathway.

Immunoprecipitation of the sample from the ZF-MIWI2 Tg testis with anti-Flag antibody and subsequent mass spectrometry analysis revealed a physical association between ZF-MIWI2 and MORC3 (Fig.1A). This association likely involves direct binding, as the experiments using 293T cells expressing MORC3 and MIWI2 confirmed the
binding phenomena (Fig.1C). Basically, the other MIWI2 binding components expressed in male embryonic germ cells are scarcely expressed under this condition. We found that MORC3 expression in the fetal testis was restricted to germ cells and the protein was predominantly localized to the nucleus and piP-body. This localization pattern is quite similar to that of MIWI2 (Fig.1D).

The data of physical binding of MORC3 with MIWI2 prompted us to determine the physiological role of MORC3. We noticed that the DNA methylation level of some classes of retrotransposons was significantly lower and both MIWI2- and MILI-associated piRNAs were reduced by about half in the MORC3 mutant embryonic testes (Fig.3-5). Thus, MORC3 is at least partially required for piRNA-directed de novo DNA methylation. It is conceivable that the loss of MIWI2 loaded piRNAs would be the reason for the incomplete de novo DNA methylation in the MORC3 mutants (Fig.6C).

MORC3 mutants lead to de-silencing of LINE1 but there was no effect on IAP expression. The other proteins involved in piRNA production, such as TDRD1, TDRD9, TDRKH, and EXD1, also regulate LINE1 expression rather than IAP. In contrast, in MILI and MVH mutants, de-suppression of IAP was observed to be as robust as that of LINE1. The loss of piRNA-dependent DNA methylation, which was detected in MILI- and MVH-null testes, leads to meiotic arrest at the pachytene stage in spermatogenesis. In contrast, the vast majority of the MORC3 mutant male germ cells escape such apoptosis and survive through the meiotic prophase (Fig.2A and B). However, the pregnancy rate of the normal female mice crossed with the MORC3 mutant male mice was significantly lower and the fertility of the MORC3 mutant male mice was slightly impaired (Fig.2C and supplementary Fig.1D).

Comparing to the phenotype of the MORC3 mutant mice, impaired fertility was not
observed in EXD1 mutant mice, despite the reduction of MIWI2-bound piRNAs, especially against LINE1. The reason for the normal fertility in EXD1 mutant mice could be the milder effect of de-repression of retrotransposons. Taken together, the high survival rate of the MORC3 mutant germ cells in spermatogenesis is presumably due to the milder degree of the deregulation of LINE1 and reduced levels of piRNAs, compared to those in MILI- and MVH- null mutants. Although it is difficult to conclude that the fertility defect of the MORC3 mutant male mice is dependent on the impaired retrotransposon expression, some association may be anticipated.

Lastly, we wanted to understand the mechanism by which the impaired retrotransposon expression was effected in the MORC3 mutant testis. Previous studies revealed that piRNA precursors are conventional RNA pol II transcripts bearing 5′ caps and 3′ poly (A) tails. The lengths of these nascent RNAs, which are transcribed from transcriptional start sites in the piRNA clusters, are varied. The coordinated increase of pachytene piRNA precursor transcripts is controlled by common transcriptional factors, such as the A-MYB protein, during spermatogenesis, especially around the pachytene stage. It has been suggested that this regulation would be caused by the compartmentalization and reorganization of TAD (topologically associating domain). However, the mechanism of transcriptional regulation for embryonic piRNA precursors remains unclear.

Our RT-qPCR data indicated that transcription of piRNA precursors originated from retrotransposon regions in piRNA clusters and that the transcripts except those of the IAP regions were significantly reduced in the MORC3 mutant embryonic testis (Fig.6A and B, and supplementary Fig.3). These data support the notion that the regulatory mechanism of transcription including IAP regions on the piRNA clusters is
different from that of transcription from the other retrotransposon regions in gonocytes.

The H3K4me3 (H3 tri-methylated lysine 4) signals, that are histone modification associated with RNA pol II transcription start sites, are highly observed in promoters of retrotransposons including piRNA clusters in E16 testes. In addition, the promoter regions of piRNA clusters show much stronger H3K4me3 signals than coding gene promoters in embryonic testis. MORC3 has the PHD X/ZF CW domain, which recognizes and binds to H3K4me3 marks, and the GHKL ATPase domain, which is involved in ATP binding and hydrolysis. Enrichment of MORC3 is found in H3K4me3 sites of active gene promoters in the genome-wide ES cell analysis. Taken these observations together, our findings suggested that MORC3 might recognize and bind preferentially to H3K4me3 marks at the promoter region of not only retrotransposon genes but also piRNA clusters and regulate the transcription of piRNA precursors via chromatin remodeling by hydrolyzing ATP in embryonic testis (Fig.6C).

Considering that MIWI2 is localized at the promoter regions of retrotransposons, it is speculated that MORC3 proteins would be localized to the similar regions through a protein complex formation and would play a role in the embryonic transposon silencing via piRNA pathway. Therefore, notably, MORC3 would have a direct contribution to the initial stages of piRNA biogenesis rather than subsequent secondary piRNA pathway, in which MIWI2 is involved. Thus, MORC3 affects not only secondary but also primary piRNA biogenesis and regulates the transcription of piRNA precursors.

MIWI2 affects the chromatin state by targeting nascent RNAs transcribed from piRNA-dependent retrotransposon regions using piRNAs as guides. The piRNAs may be generated in part from the nascent RNAs transcribed by the regulation of MORC3.
Thus, MIWI2 plays the role as a recruiter of proteins related to de novo DNA methylation on retrotransposons, while MORC3 controls the transcription of piRNA precursors from retrotransposon related piRNA cluster regions of the entire genome. MORC3 and MIWI2 might play an individual role on piRNA-dependent retrotransposon regions, at the first step of primary piRNA biogenesis and after secondary piRNA biogenesis, respectively. Thus, apart from their direct association as the protein complex, the functional relationship between MORC3 and MIWI2 presumably is indirectly in this transposon silencing system.

In this study, we have identified MORC3 as a nuclear effector of retrotransposon silencing via piRNA-dependent de novo DNA methylation. Our data suggest that MORC3 acts as an active regulator of piRNA precursors in the piRNA pathway and subsequently affects piRNA production. Thus, we provide the first mechanistic insights into effector protein at the first step of piRNA biogenesis in embryonic transposon silencing mechanism.

Methods
Animals
All the animal experiments were performed in accordance with the general guidelines of The Institute of Experimental Animal Science, Osaka University Medical School. The transgenic mice carrying the FLAG·NLS·ZFP and FLAG·NLS·ZFP·MIWI2 transgene were previously reported. Morc3 conditional knock out mice were generated using Morc3 floxed mice (gift from Dr Haruhiko Koseki (RIKEN Center for Integrative Medical Sciences) and Tnap·Cre transgenic mice (gift from Dr Andras Nagy). Founder
mice were mated with C57Bl/6 mice to generate the lines. The transgenic mice and

*Morc3* conditional knock out mice were validated by PCR with specific primers.

**Generation of *Morc3* conditional knockout mice**

We generated conditional knockout mouse lines in which we inserted the targeting

vector (pDTMorc3neo) from the bacterial artificial chromosome clone (RP23-119J20) containing mouse genomic DNA using the double Red recombination method as described previously. This targeting vector was constructed to produce an appropriate conditional allele for Morc3. Parallel LoxP sites were inserted such that they flanked a critical portion of the gene: the LoxP sites flanked exon 2 (containing a part of Histidine kinase/HSP90-like ATPase superfamily domain) of Morc3 (supplementary Fig.4). The DNA sequence of the targeting vector (pDTMorc3neo) is shown in supplementary Table 1. The linearized targeting vector was introduced into mouse ES cells (lab-made ES cells originated from hybrid embryos derived from C57BL/6J and C57BL/6N mice) by electroporation (GenePulser; Bio-Rad). The ES cells were grown on feeder cells in an appropriate medium. Each colony was picked up and expanded, and the genomic DNA of each clone was purified. To identify true homologous recombinant colonies, we performed direct sequencing through PCR amplification using the outward primers for the homologous region and the inward primers for the neomycin resistance gene. Targeted ES cells were used for generating chimeric mice. The resulting chimeric mice were backcrossed to C57BL/6J mice for over 5 generations.

**Isolation of testicular germ cells**
Whole testes (D12 after birth) were removed and treated with 1 mg/mL collagenase type II and DNase1 for 15 min at 37°C. These samples were suspended with 0.25% trypsin for 10 min at 37°C and washed with Hank’s Stock Solutions (HBSS) (Nacalai Tesque Inc., Kyoto, Japan). Anti-EpCAM (PE) antibody (CD326) (G8.8, BioLegend) was added to the cell suspension with 5% BSA/PBS. After stirring for 90 min at 4°C, the immune complex was washed with HBSS. The germ cells were sorted using a BD FACS Aria system (BD Biosciences, Franklin Lakes, NJ, USA) after immunostaining with the anti-EpCAM antibody. The germ cell purity was verified by rerunning the sample after sorting and was determined to be > 90%.

**Isolation of sperm**

Mouse sperm were isolated from the cauda epididymis of more than 10-week-old wild type (C57Bl/6) and the MORC3 mutant mice by dissecting tissue in PBS. Sperm were allowed to swim up for 30 min in 37°C with 5% CO2. To avoid the contamination of somatic cells, the upper fraction was collected for bisulfite sequence analysis.

**Western blotting**

The immunoprecipitates or lysates were separated by SDS-PAGE (7.5% gel) and transferred to a PVDF membrane (Millipore, Bedford, MA, USA). After blocking, the filters were incubated with an anti-MIWI2 monoclonal antibody (25D11), anti-MIWI2 polyclonal antibody (MIWI2-C), anti-MILI antibody (26F) 46, anti-MORC3 antibody (D238-3; MBL, Japan), anti-PA antibody (Wako, Japan), anti-FLAG M2 antibody (Sigma Chemical Co., St. Louis, MO, USA), or β-actin antibody (Sigma 5441). Anti-MIWI2 monoclonal antibody (25D11) was produced with GKGRQDFEELGVC (a 69–80 aa
peptide sequence of MIWI2) as the antigen by Dr Yasuyuki Kurihara (Yokohara
National University). An anti-MIWI2-C polyclonal antibody was generated by
immunization with peptides derived from MIWI2 (amino acids 831–847:
SVHKEPSLELANNLFYL). HRP anti-mouse IgG and HRP anti-rabbit IgG (Pierce,
Rockford, IL, USA) were used as the secondary antibody, and the signal was detected
using ECL Western Blotting detection reagents (GE Healthcare, Chalfont, Bucks, UK).

**Bisulfite sequencing**

Sorted germ cells (D12 after birth) were treated with bisulfite using the EpiTect Fast
DNA Bisulfite Kit (Qiagen, Valencia, CA, USA). The primers that we used for the
bisulfite sequencing of type A LINE-1 and type TF LINE-1 loci in this experiment
recognize more than several thousand loci in the mouse genome. Meanwhile, among
IAP retrotransposon species, only 1Δ1-type IAP, which is only 5–6 % of total IAP
elements, shows piRNA-dependent DNA methylation in 5′-LTR. Unfortunately,
5′-LTR sequences of 1Δ1-type IAP could not be distinguished from those of the other
types of IAP. Therefore, we have used the primers specifically recognizing one locus of
1Δ1-type IAP (in this experiment, a locus in Chromosome 3). The PCR primer sequences
are described in the Supplementary information. The first and second rounds of PCR
amplification of the IAP and the *H19* (GenBank accession no. U19619) were carried out
using Ex Taq (Takara, Japan). The PCR conditions were as follows: an initial round of 2
min at 94°C followed by 30 cycles of 30 s at 94°C, 30 s at 50°C (for H19) or 1 min at 55°C
(for IAP), and 1 min at 68°C, and the second round of 2 min at 95°C followed by 15
cycles of 30 s at 95°C, 30 s at 50°C (for H19) or 1 min at 56°C (for IAP), and 1 min at
72°C. Nested PCR was performed to amplify the *H19* differentially methylated regions
(DMRs). The PCR for the type A LINE-1 (GenBank accession no. M13002) and type TF LINE-1 (GenBank acc. no. D84391) was carried out using EpiTaq HS (for bisulfite-treated DNA) (Takara, Japan) under the following conditions: 2 min at 94°C followed by 30 cycles of 30 s at 94°C, 30 s at 55°C (for type A LINE-1) or 30 s at 50°C (for type TF LINE-1), and 1 min at 68°C. The PCR products were purified using the QIAquick Gel Extraction Kit (Qiagen, Valencia, CA, USA), cloned into the pGEM-T Easy Vector (Promega, Madison, WI, USA), and sequenced using an Applied Biosystems 3730 DNA Analyzer (Thermo Fisher Scientific, CA, USA).

**Northern blot analysis**

Total RNA samples were prepared from testes using ISOGEN (Nippon Gene CO., LTD., Tokyo, Japan) and stained with 0.02% methylene blue. RNA Northern blot analysis was performed at 65°C in a solution containing 0.2 M NaHPO₄ (pH 7.2), 1 mM EDTA, 1% BSA, and 7% SDS. The membranes were washed with a 0.2 × SSC, 0.1% SDS solution at 65°C. The subcloned PCR products were labeled with [α-³²P]-dCTP and used as probes. The sequences used for the PCR primers were as follows: the 3′ noncoding region of IAP (GenBank accession no. X04120), nucleotides 4489–4793, and the 5′ noncoding region of type A LINE-1 (M13002), nucleotides 531–1642 and type TF LINE-1(D84391), nucleotides 874–1156.

**Immunohistochemical staining**

Testes of the *Morc3* homozygous and heterozygous mutant and wild type male mice were dissected and fixed in 4% paraformaldehyde for 2 h at 4°C (for D30 after birth) or 2% paraformaldehyde for 1 h at 4°C (for E14.5, 16.5, and 17.5 testes). After washing in
PBS containing 10% and 20% sucrose, the testes were embedded in OCT compound. The cryosections blocked with 10% normal goat serum and 3% BSA in PBS for 0.5 h at room temperature were immunofluorescence stained, after treatment with HistoVT one (Nacalai Tesque Inc., Kyoto, Japan) for E16.5 and E17.5 testes. Sections were treated with anti-MORC3 antibody (D238-3; MBL, Japan), anti-MVH antibody (ab13840; abcam), anti-MILI antibody (#5940; Cell Signaling technology), anti-MIWI2 antibody (ab21869; abcam), or anti-MIWI polyclonal antibody (#2079; Cell Signaling) overnight at 4°C. Alexa Fluor 488- or 568-conjugated anti-rabbit immunoglobulins (H+L) or Alexa Fluor 488- or 568-conjugated anti-mouse immunoglobulins (H+L) (Molecular Probes, Eugene, OR, USA) were used as the secondary antibody for 1 h at room temperature. Nuclei were counterstained with 1 μg/mL DAPI. Immunostained cryosections were examined under a confocal microscope (LSM5Pascal, Carl Zeiss Co. Ltd).

HE staining

Eight-week-old testes and caudal epididymis of the Morc3 homozygous mutant and wild type male mice were dissected and fixed in 4% paraformaldehyde for 2 h at 4°C. After washing in PBS containing 10% and 20% sucrose, they were embedded in OCT compound. The sections were stained with hematoxylin (Wako, Japan) and eosin. After staining, the sections were treated with ethanol and xylene. The image was obtained using the BZ-X710 microscope (Keyence, UK).

Immunoprecipitation, SDS-PAGE, and silver staining

The E16.5 or 3-week-old testes of wild type and the mutant mice carrying the *FLAG-NLS-ZFP* and *FLAG-NLS-ZFP-MIWI2* transgene were homogenized in a lysis
buffer (20 mM Tris [pH 7.5], 150 mM KCl, 0.5% TritonX, 2.5 mM MgCl₂, 1 mM DTT, Rnasin (Promega), and a protease inhibitor tablet (Roche)). The lysates were freeze-thawed and treated with benzonase. The homogenates were centrifuged at 15000 × rpm for 10 min at 4°C and the supernatant was subjected to immunoprecipitation using anti-FLAG (FLA1; MBL) antibody overnight at 4°C and protein G Dynabeads (Invitrogen, US) for 1.5 h at 4°C. The immune complex was washed with the lysis buffer three times. Immunoprecipitates were subjected to SDS-PAGE on a 7.5% gel. The gels were subjected to silver staining using the Silver Stain MS Kit (Wako, Japan) or western blotting.

The transfected 293T cells were treated with lysis buffer (20 mM HEPES [pH 7.5], 0.1% NP-40, 150 mM NaCl, 2.5 mM MgCl₂, 1 mM DTT, and protease inhibitor tablet (Roche)). The lysates were precleared with Protein G Dynabeads (Invitrogen, US) for 1 h at 4°C and incubated with anti-PA antibody (Wako, Japan) or anti-FLAG (FLA1; MBL) antibody overnight at 4°C. The immune complex was washed with the lysis buffer three times. The immunoprecipitates were subjected to western blotting.

**Mass spectrometry**

Mass spectrometry was performed with the UltiMate 3000 Nano LC system, Q-Exactive (Thermo Fisher Scientific) and analyzed with MASCOT software (Matrix Science, London, UK) by CoMIT Omics Center in Osaka University.

**RT-qPCR**

Total RNAs prepared from E16.5 testes were treated with Turbo DNase (Life Technologies) and subjected to RT-PCR using SuperScript III Reverse Transcriptase
(Invitrogen) and Oligo (dT), and to Q-PCR using THUNDERBIRD SYBR qPCR MIX (TOYOBO, Osaka, Japan). The qPCR was analyzed using the CFX384 Real-Time PCR system (BIO-RAD), and the specific primers used are described in the Supplementary information.

piRNA preparation and Deep sequence analysis

Total RNA samples were prepared from E16.5 testes of the Morc3 homozygous mutant and wild type (control) mice using ISOGEN (Nippon Gene CO., LTD., Tokyo, Japan). Small RNA Library was generated with TruSeq small RNA Library Prep Kit and analyzed with Illumina HiSeq (Macrogen Inc). For immunoprecipitation of MILI or MIWI2 piRNA complex, 38 testes from the Morc3 homozygous mutant and wild type (control) mice at E16.5 were collected, respectively. The collected testes were homogenized in lysis buffer (20 mM HEPES (pH 7.3), 150 mM NaCl, 2.5 mM MgCl2, 0.1% NP40, 1 mM DTT) containing protease inhibitor tablet (Roche) and RNasin (Promega). The piRNA complexes were immunoprecipitated using 14 testes by anti-MILI (PM044, MBL) and using 24 testes by anti-MIWI2 (Miwi2-N1) 46 antibodies, respectively. Then, the samples were subjected to RNA purification using ISOGEN-LS (Nippon Gene CO., LTD., Tokyo, Japan) and to western blotting using anti-MILI (26F) and anti-MIWI2 (25D11) antibodies. The small RNA Libraries were generated with TruSeq small RNA Library Prep Kit and analyzed with Illumina NovaSeq. This work was supported by Japan Society for the Promotion of Science (JSPS). The raw sequence data in all samples were processed by CLC Genomics Workbench software (Filgen Inc.) to trim of miRNA (miRbase), rRNA (5S rRNA Database), and tRNA (GtRNAdb) allowing up to 2 mismatches after removing adaptor sequence, low-quality reads, and reads.
below length (15 nt) and above length (45nt). The trimmed 25–31 nt length small RNA reads (corresponding to the length of piRNAs) were mapped to repetitive DNA consensus sequence using Dfam database allowing up to 3 mismatches after mapping the mouse genome allowing 0 mismatches (UCSC / mm10) and mapped to L1A sequence (M13002), L1TF sequence (D84391), and IAP sequence (M17551) allowing up to 2 mismatches. The number of mapped read counts was normalized by the read-depth of each library.

Plasmid, Cell culturing and transient transfection

The 293T cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FCS. The 293T cells were transfected with PA-tagged MIWI2 plasmid and Flag-tagged MORC3 plasmid (gift from Dr Norimitsu Inoue (Department of Tumor Immunology Molecular Genetics, Osaka Medical Center for Cancer and Cardiovascular Disease, Osaka, Japan)) using polyethylenimine (Cosmo Bio, Tokyo, Japan) according to the manufacturer’s instructions. At 27 h post-transfection, the cells were harvested by centrifugation for 10 min at 3000 × rpm at 4°C.

Male fertility test

Sexually mature mutant male mice were caged with 7-week-old wild type (C57Bl/6) females for several months. A check for the vaginal plug was carried out every morning, and the number of pups in the cage was counted.

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**Author Contributions**

K.K.K. performed the experiments and analyzed the data. K.K.K., S.K.M. and T.N. designed the experiments and wrote the paper. H.K. and M.N. supported the generation of conditional knockout mice. S.E.J. discussed about the experiment with K.K.K.
Competing interests

The authors declare no competing interests.

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Figure legends

Figure 1

MORC3 as an interaction partner of MIWI2

(A) Silver staining of the postnatal testis proteins that co-immunoprecipitated with the anti-Flag antibody. Testis lysates of 21-day-old ZF only and ZF-MIWI2 Tg mice were subjected to IP (immunoprecipitation) with the anti-Flag antibody and SDS-PAGE was carried out. The 130 kDa protein that bound specifically to MIWI2 (black arrow) was examined by LC-MS/MS.

(B) Binding of ZF-MIWI2 to MORC3 in embryonic testes. IP of the embryonic testes lysates of E16.5 wild type and ZF-MIWI2 Tg mice was carried out with the anti-Flag antibody. The immunoprecipitated MORC3, ZF-MIWI2, and MIWI2 proteins were detected using the corresponding antibodies in western blotting.

(C) In vitro co-IP assays for MIWI2 and MORC3. HEK 293T cells were transfected with the plasmids expressing PA-tagged MIWI2 and Flag-tagged MORC3. The 293T cells were co-transfected with the tagged protein expression constructs. The lysates were
immunoprecipitated with the anti-PA or anti-Flag antibodies and subsequently subjected to western blotting with these antibodies.

(D) Co-immunostaining of the testes of the E17.5 wild type mice with the anti-MORC3 antibody (red), anti-MIWI2 (green) or anti-MILI (green) antibodies, and DAPI (blue) for DNA are shown. Scale bar, 20 μm.

Figure 2

Subfertility of the MORC3 mutants

(A) Testes and caudal epididymis in the wild type and MORC3-cKO mice (left). Testis weights and sperm count in individual mice (right). Error bars denote SD. Scale bar, 5 mm.

(B) HE staining of the testes and caudal epididymis of 8-week-old wild type and MORC3-cKO mice. Scale bar, 100 μm.

(C) The percentage of delivery by female mice that mated with MORC3-cKO male mice. Error bars denote SD. A significant difference (*p < 0.05 by the t test) between wild type and MORC3-cKO data is shown.

Figure 3

Expression of LINE1 retrotransposons and DNA methylation status of testicular male germ cells and sperm of the MORC3 mutants

(A) Northern blot analysis of LINE1 and IAP-1Δ1 retrotransposons in the testes of 5-month-old control wild type and MORC3-cKO mice, 3-week-old MORC3-cKO and MILI-KO mice. The 5′ UTRs of types A and TF LINE1 and the 3′ UTR of IAP were used as probes.
Representative data of bisulfite sequencing of 12-day-old male germ cells purified by the anti-EpCAM antibody (B) and sperm (D). The 5′ UTR region of LINE1 (types A [GenBank: M13002] and TF [GenBank: D84391]) and the LTR region of the 1Δ1-type IAP in chromosome 3qD were analyzed using the specific primers (sequences provided in the Supplementary information). Dots and circles represent methylated and unmethylated CpGs, respectively. Gaps in the methylation profiles represent mutated or unreadable CpG sites. The percentages of methylated CpGs are shown below each panel.

Statistical analyses of three independent bisulfite sequencing experiments using 12-day-old male germ cells (C) and sperm (E). Error bars denote SD. Significant differences (p < 0.05, “p < 0.01, ***p < 0.0005 by t-test) between the indicated data of the type A and TF LINE1 (top and middle panels) are shown.

Figure 4: piRNA biogenesis and sequence profile in the MORC3 mutant embryonic testes

(A) Length distribution of small RNAs from E16.5 wild type and MORC3-cKO testes. Small RNAs were analyzed after ribosomal RNA (rRNA), micro RNA (miRNA), and transfer RNA (tRNA) mapped reads were removed by CLC Genomics Workbench. Light blue and pink bars show the wild type and MORC3-cKO data, respectively. The accession number of deep sequencing data is DRA011066.

(B) Expression of 25–31 nt small RNAs corresponding to piRNAs in individual samples.

(C) Genomic annotation and classification of repeat piRNAs based on retrotransposon class in the wild type and MORC3-cKO piRNA libraries (upper). Expression of piRNAs corresponding to LINE, LTR (non-IAP), LTR (IAP), and SINE in individual repeat
(D) Expression of piRNAs derived from both strands (upper) and each strand (sense or antisense, bottom) corresponding to LINE1 A (M13002), LINE1 TF (D84391), and IAP (M17551) sequences in the wild type and MORC3-cKO piRNA libraries.

(E) Length distribution of 25–31 nt small RNAs derived from each strand (sense or antisense, left). Nucleotide distribution of the 1st or 10th nucleotide of 25–31 nt small RNAs from each strand (right).

Figure 5

MIWI2- and MILI-associated piRNAs in the MORC3 mutant embryonic testes

(A) Length distribution of small RNAs associated with MIWI2 and MILI from E16.5 wild type and MORC3-cKO testes. The small RNAs were analyzed after ribosomal RNA (rRNA), micro RNA (miRNA), and transfer RNA (tRNA) mapped reads were removed by CLC Genomics Workbench (upper). Reads of 25–31 nt small RNAs corresponding to piRNAs in individual samples (bottom). The accession number of deep sequencing data is DRA009594.

(B) Genomic annotation and classification of repeat piRNAs based on retrotransposon class in MIWI2- and MILI-associated piRNA libraries from wild type and MORC3-cKO testes (upper). Reads of MIWI2- and MILI-associated piRNAs corresponding to LINE, LTR (non-IAP), LTR (IAP), and SINE in individual repeat piRNAs (bottom).

(C) Reads of 25–31 nt small RNAs derived from each strand (sense or antisense) corresponding to LINE1 A (M13002), LINE1 TF (D84391), and IAP (M17551) sequences in the MIWI2- and MILI-associated piRNA libraries from wild type and MORC3-cKO testes.
(D) Co-immunostaining of the testes of the E16.5 MORC3-heterozygous and the mutant mice with anti-MIWI2 antibody (red), anti-MORC3 antibody (green), and DAPI (blue) for DNA are shown. Scale bar, 20 μm.

Figure 6

Expression of piRNA precursors from representative embryonic piRNA clusters

(A and B) Structure of the piRNA clusters (Chr 7 (1) and Chr 10) and positions of individual primer sets. The regions of the IAP element sequence in each cluster are described with red bars (from piRBase in UCSC Genome Browser mm10) (upper).

Quantitative RT-PCR for the expression analysis of piRNA precursors transcribed from embryonic piRNA clusters using E16.5 wild type and MORC3-cKO embryonic testes (bottom). Data is normalized by β-actin and is shown as means and SD (Error bar) from more than triplicate PCR reactions. Significant differences (p < 0.05 by the t test) between wild type and MORC3-cKO data using primer sets for the Chr 7 (1)-1, 4, and 5 positions (A) and Chr 10-1 and 2 positions (B) are shown.

(C) Schematic diagram of de novo DNA methylation pathway mediated piRNAs biogenesis involved with MORC3.
Figure 1

A

B

C

D

E17.5 testis
Figure 2

A

Testis size

WT  MORC3-cKO

Caudal epididymis size

WT  MORC3-cKO

B

Testis

WT  MORC3-cKO

Caudal epididymis

WT  MORC3-cKO

C

The rate of delivery of female mice coupled with MORC3-cKO male mouse

* P<0.05
Figure 3

A

WT     MORC3-cKO    MILI-KO
5M 3W 5M 3W

#115 #122 #127 #203

LINE1(A)
LINE1(TF)
IAP

28S 18S
### Figure 3

#### Postnatal testicular germ cells

|          | WT     | MORC3-cKO |
|----------|--------|-----------|
| **LINE1(A)** | ![Graph](image) | ![Graph](image) |
|           | 91.8%  | 91.0%     |
| **LINE1(TF)** | ![Graph](image) | ![Graph](image) |
|           | 85.9%  | 85.1%     |
| **IAP**   | ![Graph](image) | ![Graph](image) |
|           | 83.3%  | 83.2%     |
| **H19**   | ![Graph](image) | ![Graph](image) |
|           | 94.2%  | 94.1%     |

#### Sperm

|          | WT     | MORC3-cKO |
|----------|--------|-----------|
| **LINE1(A)** | ![Graph](image) | ![Graph](image) |
|           | 93.3%  | 81.9%     |
| **LINE1(TF)** | ![Graph](image) | ![Graph](image) |
|           | 80.2%  | 66.0%     |
| **IAP**   | ![Graph](image) | ![Graph](image) |
|           | 89.1%  | 98.4%     |
| **H19**   | ![Graph](image) | ![Graph](image) |
|           | 91.5%  | 90.8%     |
Figure 4

A

Length distribution and Expression of small RNAs

Length (nt)

Expression of 25-31 nt Small RNAs

WT

MORC3-cKO

B

Expression of 25-31 nt Small RNAs

WT

MORC3-cKO

C

WT

MORC3-cKO

Expression of 25-31 nt Small RNAs

WT

MORC3-cKO
Expression of 25-31 nt Small RNAs (RPM)

Figure 4
Figure 4

E

Ssense

Antisense

WT
MORC3-cKO

Length (nt)

(RPM)

Nucleotide distribution

1st
10th

U
G
C
A

WT
MORC3-cKO

Length (nt)

(RPM)

Nucleotide distribution

1st
10th

U
G
C
A
Figure 5

**A**

**IP MIWI2**

![Graph showing IP MIWI2 distribution with RPM on the y-axis and 20-40 nt on the x-axis for WT and MORC3-cKO.]

**IP MILI**

![Graph showing IP MILI distribution with RPM on the y-axis and 20-40 nt on the x-axis for WT and MORC3-cKO.]

**B**

**IP MIWI2**

![Graph showing pie charts for WT and MORC3-cKO with percentages of LINE, LTR (IAP), LTR (non-IAP), SINE, and Other categories.]

**IP MILI**

![Graph showing pie charts for WT and MORC3-cKO with percentages of LINE, LTR (IAP), LTR (non-IAP), SINE, and Other categories.]

WT vs. MORC3-cKO:
- **LINE**: WT - 33%, MORC3-cKO - 30%
- **LTR (IAP)**: WT - 12%, MORC3-cKO - 10%
- **LTR (non-IAP)**: WT - 14%, MORC3-cKO - 16%
- **SINE**: WT - 6%, MORC3-cKO - 2%
- **Other**: WT - 34%, MORC3-cKO - 61%

**RPM**

![Bar graphs showing RPM for LINE, LTR (IAP), LTR (non-IAP), and SINE for WT and MORC3-cKO.]

- **WT**: LINE - 9000 RPM, LTR (IAP) - 6000 RPM, LTR (non-IAP) - 3000 RPM, SINE - 1000 RPM
- **MORC3-cKO**: LINE - 1000 RPM, LTR (IAP) - 5000 RPM, LTR (non-IAP) - 2000 RPM, SINE - 500 RPM
Figure 5

C  

IP MIWI2

| RPM | sense | antisense |
|-----|-------|-----------|
| WT  |       |           |
| MORC3-cKO |       |           |

IP MILI

| RPM | sense | antisense |
|-----|-------|-----------|
| WT  |       |           |
| MORC3-cKO |       |           |

D  

MORC3-hetero

E16.5 testis
Figure 6

A  Chr 7 (1) piRNA cluster

Relative expression
(normalized by β-actin)

\( N \geq 3 \)

\( *p < 0.05 \)

RT qPCR

Chr 7 (1) (mm10)

Chr 7(1)- 1  Chr 7(1)- 2,3  Chr 7(1)- 4,5

**control**  **MORC3-cKO**

\( *p < 0.05 \)

\( N \geq 3 \)  (E16.5 testis)

B  Chr 10 piRNA cluster

Relative expression
(normalized by β-actin)

\( N \geq 3 \)

RT qPCR

Chr 10 (mm10)

Chr 10- 1,2  Chr 10- 3,4  Chr 10- 5,6

**control**  **MORC3-cKO**

\( *p < 0.05 \)

\( N \geq 3 \)  (E16.5 testis)
Figure 6

C

WT

MORC3

piRNA precursors transcripts

Embryonic piRNA cluster

piRNA biogenesis

MIWI2

piRNA precursors transcripts

Cap

AAA

AAA

AAA

AAA

MORC3

H3K4me2/3

MORC3-cKO

piRNA precursors transcripts

Cap

AAA

AAA

AAA

Embryonic piRNA cluster

piRNA biogenesis

MIWI2

Transposon silencing

Incomplete transposon silencing