

1 Title: Meioc-Piwil1 complexes regulate rRNA transcription for differentiation of
2 spermatogonial stem cells

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18

1 **Abstract**

2 Ribosome biogenesis is vital for sustaining stem cell properties, yet its regulatory mechanisms
3 are obscure. Herein, we show unique properties of zebrafish *meioc* mutants in which
4 spermatogonial stem cells (SSCs) do not differentiate or upregulate rRNAs. Meioc colocalized
5 with Piwil1 in perinuclear germ granules, but Meioc depletion resulted in Piwil1 accumulation
6 in nucleoli. Nucleolar Piwil1 interacted with 45S pre-rRNA. *piwil1^{+/-}* spermatogonia with
7 reduced Piwil1 upregulated rRNAs, and *piwil1^{+/-};meioc^{-/-}* spermatogonia recovered
8 differentiation later than those in *meioc^{-/-}*. Further, Piwil1 interacted with Setdb1 and HP1 α , and
9 *meioc^{-/-}* spermatogonia exhibited high levels of H3K9me3 and methylated CpG in the
10 45S-rDNA region. These results indicate that zebrafish SSCs silence rRNA transcription with
11 repressive marks similar to *Drosophila* piRNA targets of RNA polymerase II, and that Meioc
12 has a unique function on preventing localization of Piwil1 in nucleoli to upregulate rRNA
13 transcripts and to promote SSC differentiation.

14

15

1 **Introduction**

2 The maintenance and control of the differentiation of adult stem cells are indispensable for
3 living animals. In this decade, the relationship between the stem cell system and translation
4 activity control has been revealed. Compared with that of differentiating progenitor cells, the
5 global translation activity is generally inhibited in most adult and embryonic stem (ES) cells,
6 and low translation activity is required for stem cells to maintain undifferentiated status (Ni and
7 Buszczak, 2023; Saba et al., 2021). Despite this common feature of low translational activity,
8 differences in the state of ribosome biogenesis (RiBi) have been observed across stem cell types.
9 ES cells, hair follicle stem cells and *Drosophila* germline stem cells (GSCs) are characterized
10 by high levels of RiBi, whereas quiescent neural stem cells (NSCs) express low levels of
11 ribosomal genes (Llorens-Bobadilla et al., 2015). Hematopoietic stem cells (HSCs) have higher
12 RiBi levels than differentiated bone marrow cells, but they have lower levels than their
13 immediate progenitors, suggesting that cell type-specific mechanisms may regulate RiBi
14 (Jarzebowski et al., 2018). In addition, phenotypes of ribosome biogenesis defects in
15 *Drosophila* GSCs appear to be controversial; attenuation of Pol I activity and the U3 snoRNP
16 complex member *wiched* (*wcd*) mutant show premature differentiation (Fichelson et al., 2009;
17 Zhang et al., 2014), while knockdown of ribosome assembly and loss of DExD/H-box proteins
18 that govern RiBi result in a loss of differentiation (Martin et al., 2022; Sanchez et al., 2016).
19 Therefore, the mechanisms by which RiBi is coupled to proper stem cell differentiation are still
20 obscure.

21 By using an ENU mutagenesis screen to identify gonadogenesis defects in zebrafish
22 (Saito et al., 2011), we isolated a unique mutant, *minamoto* (*moto*), in which germ cells arrest at

1 the early stage of spermatogonia (Kawasaki et al., 2016). By whole genome sequencing, we
2 found that the *moto* phenotype is tightly linked to a mutation within a gene
3 (ENSDARG00000090664) (Bowen et al., 2012), which encodes the coiled-coil
4 domain-containing protein Meioc. Database analysis shows orthologs of MEIOC in vertebrates
5 and most invertebrates, but not in *Drosophila*. The mouse MEIOC has a function to maintain an
6 extended meiotic prophase I with its binding partner YTHDC2 (Abby et al., 2016; Soh et al.,
7 2017). On the other hand, a complex of *Drosophila bam* (*bag of marbles*), a functional homolog
8 of *Meioc*, and *bgn* (*benign gonial cell neoplasm*), a paralog of *Ythdc2* (Jain et al., 2018), plays
9 a pivotal role in promoting stem cell differentiation (Perinthottathil and Kim, 2011). Although
10 zebrafish *meioc* (*moto*) is an orthologous to the mice gene, it appears to be functionally similar
11 to *Drosophila bam*, making it interesting to see how it controls the development of germ cells.

12 The present study revealed that zebrafish had spermatogonial stem cells (SSCs) with low
13 rRNA transcription, a characteristic that differed from many other stem cells, and that Meioc
14 was required for upregulation of rRNA transcription and SSC differentiation. Independently of
15 *Ythdc2*, Meioc regulated intracellular localization of Piwil1 that interacted with transcriptional
16 silencing proteins, *Setdb1* (Eggless) and *HP1 α* (Cbx5). Our results suggested that rRNA
17 silencing maintained the undifferentiated state of zebrafish SSCs, providing a new insight on the
18 relationship between the stem cell system and the RiBi control.

19

20 **Results**

21 **Failure to differentiate spermatogonia in the zebrafish *meioc* mutant**

1 In fish testes, germ cells are surrounded by Sertoli cells within a basement membrane
2 compartment and develop synchronously in cysts. Thus, the developmental stage of
3 spermatogonia can be determined by the number of cells within the cyst (**Figure 1A**). Wild-type
4 spermatogonial cells undergo 8 rounds of cell division before entering meiosis (Leal et al.,
5 2009), however *moto*^{l31533} mutant testes had only up to 3 rounds of division, containing 2-8
6 spermatogonia per cyst (**Figure 1B**). Mutant spermatogonia were positive for the
7 undifferentiated spermatogonia marker, Plzf, which is expressed in single- to 8-cell (1-8-cell)
8 cyst spermatogonia in the wildtype (**Figure 1B**) (Ozaki et al., 2011), but negative for meiotic
9 markers, Sycp1 and Sycp3 (Saito et al., 2014, 2011). Phosphohistone H3 immunostaining
10 indicated that mutant spermatogonia proliferated, but apoptotic cells were increased in the
11 mutant spermatogonia (**Figure S1A**). In zebrafish, mutants that fail to undergo oocyte
12 development become male sex type (Saito et al., 2011; Houwing et al., 2007; Kamminga et al.,
13 2010; Rodríguez-Marí et al., 2010; Shive et al., 2010). Similarly, no females developed in *moto*
14 mutants (**Supplemental Table S1** and **Figure S1B**), as completion of initial stages of meiosis I
15 are a pre-requisite for follicle formation (Elkouby and Mullins, 2017). These results suggested
16 that the *moto*^{l31533} mutant germ cells remained in an undifferentiated state.

17 By whole genome sequencing and genetic analysis, we found that the *moto*^{l31533} mutant
18 phenotype was tightly linked to a mutation within the ortholog of mouse *meioc* gene (**Figure**
19 **S1C**) (Bowen et al., 2012). The *moto*^{l31533} mutant inherited the phenotypes described above for
20 more than 10 generation. Furthermore, fish homozygous for another allele, *moto*^{sa13122}, exhibited
21 the same gonadal phenotype as *moto*^{l31533} homozygotes, and *moto*^{sa13122} failed to complement
22 *moto*^{l31533} confirming that these mutations disrupt the same gene (hereafter denoted *meioc*^{mo})

1 **(Figure S1D)**. We searched for paralogues of *meioc* gene using the conserved coiled-coil
2 domain (PF15189) **(Figure S1C)** but were not able to find another gene in GRCz11 (version
3 111.11). We then generated an antibody against the N-terminus of Meioc **(Figures S1E and F)**
4 and could not detect the expression of truncated Meioc in the mutants **(Figure S1G)**.

5 Zebrafish *meioc* RNA and protein signals were observed in a portion of 1-2-cell cyst
6 germ cells in juvenile gonads at 25 days post-fertilization (dpf), adult ovaries and testes **(Figure**
7 **S2A)**. The cells exhibited a large nucleolus, characteristics of presumed germline stem cells
8 (GSCs) in zebrafish (Kawasaki et al., 2016) and stem-type self-renewing type I germ cells in
9 medaka (Nishimura et al., 2015). Both clear RNA and protein signals were detected in
10 premeiotic germ cell clusters, and the protein was detected as granular structures in the
11 cytoplasm with increasing sizes among cells until meiotic prophase I **(Figures S2A-C)**.

12

13 **Upregulation of translation is required for the differentiation of spermatogonia**

14 Since the global translation activity is generally inhibited in stem cells, we analyzed global
15 translational activities of zebrafish spermatogonia using the O-propargyl-puromycin (OP-Puro)
16 assay (Liu et al., 2012). In wildtype, we observed low levels of de novo protein synthesis in a
17 portion of 1-2-cell cyst spermatogonia and high levels in almost all large cysts with more than
18 32 cells ($32 \leq$ -cell) of differentiated spermatogonia **(Figures 1C and D)**. *meioc*^{mo/mo}
19 spermatogonia showed low levels as that of 1-2-cell cysts **(Figures 1C and D)**. These results
20 suggested that some germ cells within 1-2 cell cysts had low translation activity and that
21 *meioc*^{mo/mo} spermatogonia had low translation activity.

1 In order to know if translational upregulation is required for differentiation of
2 spermatogonia, we examined the effect of cycloheximide on the development of spermatogonia
3 in vitro. Cycloheximide decreased OP-Puro fluorescence intensities in differentiating
4 spermatogonia with a dose-dependent manner in testis organ culture (**Figure S2D**). At 1.0 μ M
5 cycloheximide that reduced OP-Puro fluorescence to approximately 60%, BrdU incorporation
6 decreased in 32 \leq -cell cysts of spermatogonia, while it was not affected in 1-4-cell
7 spermatogonia (**Figure S2E**). We then examined the effect on differentiation of SSCs using a
8 culture system, in which differentiation can be induced on the Sertoli cell line, ZtA6-12
9 (Kawasaki et al., 2016). After propagation of SSCs for 1 month, we transferred the cells onto
10 ZtA6-12 and treated with lower concentration of cycloheximide at 0.2 μ M, which reduced
11 OP-Puro fluorescence to approximately 70%. The treatment maintained SSCs with a large
12 nucleolus and not expressing Sycp3 (**Figures 1E and F**). Bmp2 secretion of ZtA6-12 cells was
13 not affected at 0.2 μ M cycloheximide (**Figure S2F**), suggesting that the effect on Sertoli cell
14 function was presumed to be minimized. These results suggested that a certain level of
15 translational activity was required for the differentiation of zebrafish SSCs.

16

17 ***meioc* mutants do not upregulate rRNAs in 1-2-cell cyst spermatogonia**

18 To estimate the state of ribosome biogenesis, we examined the expression patterns of rRNAs
19 and the ribosomal protein Rpl15 in zebrafish spermatogonia development. Interestingly, the
20 signal intensities of 5.8S, 18S, 28S rRNAs, and Rpl15 were low in a portion of wildtype
21 1-2-cell cyst spermatogonia (**Figure 2A**). We did not observe a low intensity of 5S rRNA.
22 Those with low signals increased in almost all 32 \leq -cell cysts and declined in spermatocytes. To

1 distinguish between cytoplasmic and nucleolar signals, we performed fluorescence *in situ*
2 hybridization of 28S rRNA. We found 1-2 cell cysts with low cytoplasmic signals also had low
3 nucleolar signals (**Figures 2B-D**). Although cytoplasmic 28S rRNA signals increased to
4 32≤-cell cysts, the highest nucleolar signals were detected in portion of 1-2-cell cysts and
5 declined at 32≤-cell cysts. These results suggest that 1-2 cell cysts contain populations with low
6 and high rRNA transcriptional activity.

7 We next tested how rRNA levels were affected in *meioc^{mo/mo}* and found low signals of
8 rRNAs and Rpl15 in spermatogonia (**Figure 2A**). We confirmed these data by performing
9 RT-qPCR from isolated spermatogonia using the *sox17::egfp* SSC marker (Kawasaki et al.,
10 2016), reductions in 5.8S, 18S, 28S rRNAs in *meioc^{mo/mo}* were seen (**Figure 2E**). Furthermore,
11 the homolog of *Drosophila* non-LTR retrotransposable element R2 that transposes exclusively
12 into 28S rDNA(Kojima and Fujiwara, 2003) also decreased in *meioc^{mo/mo}*. We next analyzed
13 unprocessed 45S pre-rRNA and pre-rRNA intermediates by Northern blot to compare levels
14 between *meioc^{mo/mo}* and the wildtype. Each was normalized to 7SL RNA. In *meioc^{mo/mo}* the
15 amount of 45S and intermediates was reduced, however levels of pre-RNA intermediate to 45S
16 pre-rRNA was similar to wildtype: 0.69 (0.49/0.71, mutant/wildtype) of unprocessed 45S
17 pre-rRNA and 0.61 (0.72/1.19) of pre-rRNA intermediates with the 5' external transcribed
18 spacer (ETS) probe, and 0.33 (0.05/0.15) and 0.35 (0.17/0.48) with the internal transcribed
19 spacer 1 (ITS1) probe (**Figure S3A**). These results indicate that 45S pre-rRNA transcripts were
20 reduced in *meioc^{mo/mo}* yet processing into pre-RNA intermediates was not affected.

21

22 **Meioc is required for demethylation of CpG in the IGS of 45S rDNA**

1 Features of silenced rDNA loci, DNA methylation, methylation of histone H3K9, and
2 association of HP1 α , are known (Grummt, 2007). In eukaryotes, the transcription of each rDNA
3 locus is regulated by DNA methylation and histone modifications in promoter and enhancer
4 regions in the intergenic spacer (IGS) region (Santoro, 2005). Therefore, we examined DNA
5 methylation state in the region. Since the regulatory sequence of rDNA was not identified in
6 teleosts, we identified tandem repeats of 90, 120 and 318 bp in the IGS region of 45S-S rDNA
7 (Locati et al., 2017) (**Figure S3B**), similar to carp (Vera et al., 2003). Unmethylated CpG in the
8 tandem repeats was frequently found in isolated wildtype *sox17::egfp* spermatogonia, while was
9 rare in *meioc^{mo/mo}* (**Figure 2F**). These results suggested that Meioc is needed for demethylation
10 of CpG in the tandem repeats of 45S rDNA IGS region.

11

12 ***meioc* and *ythdc2* mutants exhibit different phenotypes in the early stage of spermatogonia**

13 The RNA helicase YTHDC2 is a binding partner of the MEIOIC in mammals (Abby et al., 2016;
14 Soh et al., 2017). Zebrafish spermatogenic cells expressed *Ythdc2* (**Figures S4A and B**), and
15 pull-down analysis showed zebrafish Meioc interacted with *Ythdc2* (**Figure S4C**). It has been
16 recently reported that *ythdc2* KO zebrafish lack germ cells (Li et al., 2022), but staining or
17 expression analysis was not done to confirm that germ cells were completely absent. We
18 separately generated *ythdc2* KO zebrafish by the CRISPR-Cas9 system (**Figure S4A**), and
19 observed that it had up to 8-cell cyst spermatogonia (**Figure 3A**). We observed that the number
20 of 4-8-cell cysts was clearly different between *meioc^{mo/mo}* and *ythdc2^{-/-}*; the ratio of 4-8-cell
21 cysts/1-2-cell cysts in the *ythdc2^{-/-}* was almost the same as that in the wildtype, while that in
22 *meioc^{mo/mo}* significantly decreased (**Figures 3B-D**). Furthermore, *ythdc2^{-/-}* spermatogonia

1 contained both low and high levels of cells with 5.8S, 18S, 28S rRNAs, and Rpl15, similar to
2 the wildtype (**Figure 3E**). The intensity of the 28S rRNA signals in *ythdc2*^{-/-} were almost same
3 as that in the wildtype (**Figure S4D**). These results suggested that Meioc functioned
4 independently of Ythdc2 on the differentiation of 1-2-cell cysts into 4-8-cell cysts.

5

6 **Meioc interacts with Piwil1 and affects its intracellular localization**

7 To explore partners of Meioc on regulation of rRNA transcription, we performed LC/MS/MS
8 for the Meioc-immunoprecipitate (IP) of SSC-enriched hyperplastic testes (Kawasaki et al.,
9 2016). The results showed the enrichment of germ granule (nuage) components, compared with
10 normal testis (Supplemental Tables S2 and S3). By immunostaining, Meioc colocalized with the
11 germ granule components, Tdrd1 (Huang et al., 2011), Tdrd6a (Roovers et al., 2018), Piwil1
12 (Houwing et al., 2007), Piwil2 (Houwing et al., 2008), and Ddx4 (Houwing et al., 2007) in
13 wildtype (**Figures S5A and B**). In *meioc*^{mo/mo}, Piwil1 was strongly detected in the nucleolus,
14 whereas others exhibited the perinuclear localization characteristic of Ddx4-positive granules as
15 in the wildtype (**Figure 4A**). In contrast, we were able to detect low levels of Piwil1 in nucleoli
16 of wildtype 1-4-cell spermatogonia with overexposed detection after the co-staining of the
17 nucleolar marker Fibrillarin (**Figure 4B**). The signals of Piwil1 in Ddx4-positive granules were
18 decreased in *meioc*^{mo/mo} spermatogonia, compared with wildtype (**Figure 4C**). We confirmed
19 that Meioc and Piwil1 directly interacted by Co-IP and revealed that Meioc bound Piwil1
20 through the Coiled coil domain (**Figure 4D; Figure S5C**). These results demonstrated that
21 Piwil1 has a property to localize in nucleoli and that Meioc interacts with Piwil1 in perinuclear

1 germ granules. In addition, we did not detect the accumulation of Piwil1 in nucleoli in *ythdc2*^{-/-}
2 **(Figure S4E)**.

3 Since Piwil1 abnormally accumulated in nucleoli in *meioc*^{mo/mo}, we asked whether
4 Piwil1-dependent piRNA generation (Houwing et al., 2007) was affected. The abundance of
5 piRNA production in *meioc*^{mo/mo} testes was detected at similar levels to those of the wildtype
6 **(Figure S5D)**. piRNAs derived from 28S, 18S rRNAs and R2 apparently decreased in
7 *meioc*^{mo/mo} probably due to the low expression of rRNA **(Figures S5E-G)**. It is unlikely that
8 Meioc critically affected piRNA generation.

9
10 **A reduction of Piwil1 upregulates rRNA transcripts and recovers the *meioc* mutant**
11 **phenotype**

12 From the above analysis, it is hypothesized that accumulation of Piwil1 in the nucleolus
13 suppresses rRNA transcription in the *meioc*^{mo/mo} background. Therefore, we examined the rRNA
14 transcript levels and the phenotype of *meioc*^{mo/mo};*piwil*^{+/-} since the *piwil*^{-/-} depletes germ cells
15 completely before testis differentiation (Houwing et al., 2007). A reduction in Piwil1 was
16 detected in *piwil*^{+/-} testes by Western blot analysis and in the spermatogonia nucleoli by
17 fluorescent immunohistochemistry **(Figures S6A and B)**. The upregulation of 28S rRNA was
18 observed in the cytoplasm and nucleoli of *piwil*^{+/-} 1-2-cell spermatogonia as compared to
19 wildtype and *meioc*^{mo/mo};*piwil*^{+/-} spermatogonia as compared to *meioc*^{mo/mo} **(Figures 5A and B)**.
20 Consistent with the increased expression of 28S rRNA, we observed the differentiation of
21 *meioc*^{mo/mo};*piwil*^{+/-} spermatogonia to the 32-cell stage compared to that of *meioc*^{mo/mo}, in which
22 spermatogonia rarely develop to the 8-cell stage **(Figures 5C and D)**. Furthermore, the number

1 of 4-8-cell cysts increased in *meioc^{mo/mo};piwill^{+/-}* (**Figures 5E and F**). The ratio of 4-8-cell to
2 1-2-cell cysts in *meioc^{mo/mo};piwill^{+/-}* increased compared to that in *meioc^{mo/mo}*, and reached at
3 almost the same level as that in wildtype (**Figure 5G**). These results demonstrated that
4 reduction of Piwill compensated the suppression of rRNA transcription and the SSC
5 differentiation defect of the *meioc^{mo/mo}* phenotypes.

6

7 **Piwill interacts with nascent 45S pre-rRNA in nucleoli of SSCs**

8 To investigate if nucleolar Piwill interact with 45S pre-rRNA, we examined pre-rRNA in the
9 RNA IP (RIP) of Piwill by qRT-PCR. Pre-rRNA was associated with Piwill compared with the
10 control IgG in the wildtype testis, and was more enriched in *meioc^{mo/mo}* (**Figure 6A**).
11 Furthermore, we examined the effect of actinomycin D (RNA polymerase I (Pol I) and Pol II
12 inhibitor), BMH-21 (Pol I inhibitor) (Colis et al., 2014), and α -amanitin (Pol II
13 inhibitor)(Bensaude, 2011) on nucleolar localization of Piwill in the *meioc^{mo/mo}* testis organ
14 culture. The nucleolar signals of Piwill declined upon treatment with actinomycin D and
15 BMH-21 but not α -amanitin, suggesting that the pre-rRNA transcript was involved in Piwill
16 localization to the nucleolus (**Figure 6B**). Together, these results suggested that Piwill
17 interacted with nascent pre-rRNA transcripts in nucleoli of SSCs.

18

19 **Nucleolar Piwill causes accumulation of H3K9me3 and HP1 α**

20 *Drosophila* Piwi requires histone methyltransferase Setdb1 to lead to H3K9me3 deposition,
21 HP1 α accumulation, and heterochromatin formation for silencing of Pol II mediated
22 transcription (Jia et al., 2022). We examined if Setdb1 and HP1 α are localized in nucleoli using

1 anti-Setdb1 (**Figure S6C**) and anti-HP1 α (**Figures S6D** and **E**) antibodies. Setdb1 was detected
2 in nucleoli in spermatogonia of 1-2-cell cysts in both wildtype and *meioc^{mo/mo}*, and co-IP using
3 *meioc^{mo/mo}* testis lysate showed interaction between Piwil1 and Setdb1 (**Figures 6C** and **D**). The
4 higher signal intensity of Setdb1 nucleolar signal was detected in *meioc^{mo/mo}* spermatogonia
5 (**Figure 6E**). To test whether the amount of nucleolar Piwil1 correlates with the silencing state
6 of rDNA loci, we conducted chromatin IP of histone H3K9me3 and Piwil1. Higher interactions
7 of H3K9me3 and Piwil1 with rDNA were detected in *meioc^{mo/mo}* testes, which have increased
8 levels of nucleolar Piwil1, than those in the wildtype (**Figures 6F** and **G**). In addition, lower
9 interactions were detected in *piwil1^{+/-}* testes, which have a low amount of nucleolar Piwil1
10 (**Figures 6F** and **G**). Furthermore, HP1 α was detected in nucleoli in spermatogonia of 1-2-cell
11 cysts in wildtype and *meioc^{mo/mo}*, and co-IP using testis lysate and pull-down assay showed
12 interaction between Piwil1 and HP1 α (**Figures 6H** and **I**; **Figure S6F**). HP1 α was
13 predominantly localized in nucleoli in wildtype and was more accumulated in *meioc^{mo/mo}*
14 (**Figure 6J**). These results indicated that Piwil1 potentially interacted with Setdb1 in nucleoli
15 and led to H3K9me3 formation and HP1 α accumulation in rDNA loci in zebrafish
16 spermatogonia.

17

18 **Meioc expression is correlated with the upregulation of 28S rRNA**

19 To further investigate the relationship with Meioc and upregulation of rRNA in SSCs, we
20 compared the expression patterns of Meioc and 28S rRNA in isolated *sox17::egfp*
21 spermatogonia. Meioc expression ranged from barely detectable to more than 50 granular dots
22 (**Figures 7A**). Nuclear localization of Meioc was observed in some *sox17::egfp* spermatogonia,

1 while in other spermatogonia and spermatocytes Meioc was detected exclusively in the
2 cytoplasm (Figure S2C, S7A). The 28S rRNA intensity correlated with increasing Meioc
3 granules (**Figures 7A**). On the other hand, Piwil1 remained expressed at almost the same across
4 most cells (**Figures 7B**). Since nucleoli were difficult to observe in the isolated cells probably
5 due to the isolation procedure for several hours, it was difficult to verify whether Piwil1 was
6 present in the nucleolus when Meioc expression was low. Therefore, we also examined the
7 expression of Piwil1 in 1-2-cell spermatogonia in wildtype testis sections. Piwil1 was detected
8 at almost same levels in all cells as same as the isolated cells (**Figure S7B**). The large nucleolus
9 was observed in the cells with more than 11-50 Meioc granules, and nucleolar localization of
10 Piwil1 was observed in the cells with 11-50 Meioc granules similar to *meioc^{mo/mo}* spermatogonia,
11 but not in the cells with $51 \leq$ Meioc granules. These results support the idea that Meioc inhibits
12 localization of Piwil1 in nucleoli and that Meioc functioned on upregulation of rRNA
13 transcripts by preventing localization of Piwil1 in nucleoli.

14

15 **Discussion**

16 This study showed that zebrafish had SSCs with low rRNA transcription activity, and that
17 increased activity correlated with differentiation. The state of low rRNA transcription activity is
18 not similar to high RiBi of ES cells or *Drosophila* GSCs, but is closer to the low expression of
19 ribosomal gene of quiescent NSCs (Llorens-Bobadilla et al., 2015). Since zebrafish SSCs has
20 long term BrdU-retaining germ cells in undifferentiated spermatogonia (Nóbrega et al., 2010),
21 low RiBi may be one of the mechanisms in quiescent cells, although HSCs show high RiBi.
22 Zebrafish 1-2-cell cyst spermatogonia had low and high nucleolar 28S rRNA subpopulations,

1 and the high rRNA cells were not found in *meioc* mutants. Therefore, high rRNA is considered
2 to be an activated state for differentiation. The progenitor 32≤-cell cysts had highest
3 translational activity and lower rRNA transcriptional activity than high 1-2-cell cysts.

4 Our results also revealed that *Meioc* was required for the reduction of H3K9me3 levels
5 and demethylation of CpG at rDNA loci. As shown in Figure 7C, *Meioc* prevented nucleolar
6 localization of Piwil1. Piwil1 interacted with Setdb1 and both proteins could be found in the
7 nucleolus. Furthermore, high levels of H3K9me3 and HP1α correlated with higher levels of
8 nucleolar Piwil1. This is similar to *Drosophila* Piwi in piRNA-dependent gene silencing for Pol
9 II transcribed genes (Jia et al., 2022). These data support the model that nucleolar Piwil1
10 interacts with Setdb1 in the nucleolus to silence rDNA transcription via H3K9me3 and
11 recruiting HP1α while *Meioc* promotes differentiation by blocking Piwil1 nucleolar localization
12 and allowing rDNA transcription. The epigenetic state of tandem repeats of the zebrafish rDNA
13 IGS region was similar to human silenced rDNA with CpG hypermethylation, levels of
14 H3K9me3, and HP1α association (Grummt, 2007). The tandem repeats have been reported as
15 enhancers of rRNA in *Xenopus* (Pikaard and Reeder, 1988), and the methylation status of the
16 region is accompanied by rRNA synthesis (Bird et al., 1981). Since Setdb1 associates with Tip5,
17 which is known to interact with DNA methyltransferase as a component of nucleolar
18 remodeling complex (NoRC)(Yuan et al., 2007; Bersaglieri and Santoro, 2019), the Setdb1 that
19 is interacted with nucleolar Piwil1 in this study is presumed to lead to CpG methylation in
20 rDNA IGS region.

21 The phenotypes of the zebrafish *meioc* mutants and *ythdc2* mutants are different from
22 those of mouse *Meioc* KO and *Ythdc2* KO, which arrest at meiotic prophase I or immature

1 follicles (Abby et al., 2016; Soh et al., 2017; Wojtas et al., 2017). In addition to different timing
2 of those expression between mice and zebrafish, the amino acid sequence of Meioc between
3 mice and zebrafish is not well conserved (28.5%), while that of Ythdc2 is well conserved
4 (65.5%). It is likely that zebrafish Meioc have different binding partners and regulate different
5 processes of germ cell development from mouse MEIOC, causing the distinct phenotypes
6 between these mutants. Many unique partners of Meioc, such as RNA binding proteins, myosin
7 proteins, Tudor domain proteins, heat shock proteins and Cnot1 were found in the present study.

8 In this study, we revealed the function of Piwil1 in regulating rRNA transcription. In
9 *Drosophila*, nucleolar localization of Piwi and impairment of its localization by inhibition of
10 RNA pol I in ovarian somatic cells and nurse cells, and accumulation of undifferentiated GSC,
11 delay of GSC proliferation and upregulation of R1, R2 and rRNA transcripts in *piwi* mutants
12 have been reported (Fefelova et al., 2017; Mikhaleva et al., 2019, 2015; Yakushev et al., 2016).
13 In *C. elegans*, the nuclear Argonaute protein NRDE-3 binds to ribosomal siRNAs (risiRNA)
14 and is translocated from the cytoplasm to the nucleolus, in which the risiRNA/NRDE complex
15 associates with pre-RNAs and reduces the level of pre-rRNAs (Zhou et al., 2017b, 2017a; Zhu
16 et al., 2018). Therefore, Argonaute proteins appear to have a conserved function on controlling
17 RiBi. This study showed that zebrafish Piwil1 used the analogous machinery for rRNA
18 transcription as Piwi-dependent silencing Pol II transcription in *Drosophila*. Since Meioc was
19 identified as a component of germ granules, the novel function of Piwil1 will open new insights
20 into the interactions of germ granules and nucleoli to regulate RiBi and SSC differentiation.

21

22 **Materials and Methods**

1 *Zebrafish*

2 The *moto*^{*i31533*} mutant fish were isolated in the Tubingen line by N-ethyl-N-nitrosourea
3 mutagenesis screening as described (Saito et al., 2011). The *moto*^{*sa13122*} mutants were isolated by
4 the Zebrafish Mutation Project (Kettleborough et al., 2013) and provided by the Zebrafish
5 International Resource Center (ZIRC), Eugene Oregon. We used *piwil1*^{*hu2479*} line (Houwing et
6 al., 2007), *India* line, *AB* line, *IM* line (Shinya and Sakai, 2011), *vas::egfp* (Krøvel and Olsen,
7 2002) and *sox17::egfp* (Mizoguchi et al., 2008) transgenic fish. The use of these animals for
8 experimental purposes was approved by the committee on laboratory animal care and use at the
9 National Institute of Genetics (approval identification numbers, 27-12 and 28-13) and the
10 University of Massachusetts Boston Institutional Animal Care and Use Committee (protocol
11 #20120032), and carried out according to the Animal Research Reporting of In Vivo
12 Experiments (ARRIVE) guidelines and to relevant guidelines and regulations.

13

14 *Identification of the meioc gene*

15 We used whole genome sequencing (WGS) to map the *moto*^{*i31533*} mutation to an approximately
16 19 Mb (~14 cM) region on chromosome 3 (Bowen et al., 2012), revealing two nonsynonymous
17 changes within this region. Analysis of the WGS data revealed two nonsynonymous changes
18 within this region. One was a missense mutation affecting the *smg1* gene
19 (ENSDARG00000054570) and the other was a nonsense mutation affecting
20 ENSDARG00000090664, orthologue of human C3H17orf104, now called *Meioc* (Abby et al.
21 2016). By further recombination mapping of the *moto*^{*i31533*} mutation using Simple Sequence
22 Length Polymorphisms (SSLPs), the mutation was found in a region of about 2.1 Mb, between

1 markers z22516 and z8680, which contained the C3H17orf104 locus and excluded the *smg1*
2 locus.

3

4 *The ythdc2 mutant*

5 The *ythdc2*^{-/-} fish were generated by CRISPR-Cas9 mutagenesis based on protocols (Chen et al.,
6 2017; Hwang et al., 2013). A single-guide RNA (Supplemental Table S5) was designed to
7 target exon 5 of *ythdc2* (ENSDART00000166268.2) to delete functional domains predicted by
8 Pfam 35.0 (Figure S2 F; Mistry et al., 2021). *ythdc2* sgRNA (100ng/μl) and 10 pmol/μl Cas9
9 NLS protein (abm) were co-injected into 1-cell stage *India* embryos. Founders were
10 backcrossed with *India* fish, and the F1 siblings were screened by genotyping. Heterozygous
11 *ythdc2* knockout carrying a -14 bp frameshift mutation in exon 5 (*ythdc2*⁻, a -14 bp deletion
12 affecting the codons from G206 that generates 63 amino acid residues from the wrong frame
13 and stop codon after the 269th amino acid) were obtained.

14

15 *Meioc and Ythdc2 antibodies*

16 To produce specific antibodies against Meioc and Ythdc2, *meioc* cDNA encoding 356 amino
17 acid residues from N-terminus and *ythdc2* cDNA encoding amino acid residue Arg743 to
18 Leu1381 were cloned into a pQE-30 vector (QIAGEN) and a pET-21a (+) vector (Novagen),
19 respectively. The 6 x histidine tag fused proteins were expressed and purified as described
20 previously (Ozaki et al., 2011). Rats and rabbits were immunized with the purified Meioc and
21 Ythdc2 recombinant proteins, respectively. Then, anti-Meios rat IgG and anti-Ythdc2 rabbit

1 IgG were purified with CNBr activated sepharose (Cytiva) conjugated with recombinant
2 proteins according to manufacturer's instructions.

3

4 *Western blot analysis*

5 Western blot analyses were performed as described (Ozaki et al., 2011) using antibodies
6 (Supplemental Table S6). Chemiluminescent signals generated with ECL Prime (GE
7 Healthcare) were detected and quantified with Chemidoc XRS Plus (Bio-Rad). For the
8 quantification of Piwil1, eight wildtype testes and nine *piwil1*^{+/-} testes were individually used
9 for the protein extraction and western blotting analysis. The amount of Piwil1, BMP-2, and
10 α -Tubulin were quantified using quantity tools of ImageLab software version 6.0.1 (Bio-Rad),
11 and Piwil1 and BMP-2 signal intensities were normalized with signal intensities of α -Tubulin.

12

13 *Histological observation*

14 Testes and juveniles were fixed in 4% PFA in PBS or Bouin's solution (Sigma) for 2 hours at
15 RT. Paraffin sections were prepared at a 5 μ m thickness and stained with hematoxylin and eosin.
16 For the count of spermatogonial cysts, complete serial sections of 3 testes each of wildtype,
17 *ythdc2*^{-/-}, *meioc*^{mo/mo}; *piwil1*^{+/+}, and *meioc*^{mo/mo}; *piwil1*^{+/-} were stained with PAS (periodic acid
18 Schiff) to stain the Sertoli cells (Saito et al., 2014). The number of spermatogonia in cysts were
19 identified by observation of adjacent sections. For wildtype testes, we counted them in
20 randomly selected 10 sections. Undifferentiated spermatogonia (1- to 8-cell spermatogonia) in
21 *ythdc2*^{-/-}, *meioc*^{mo/mo}; *piwil1*^{+/+}, and *meioc*^{mo/mo}; *piwil1*^{+/-} testes were counted in randomly
22 selected 10 fields (23547.2 μ m²/ field) of sections, and estimated average number of each stage

1 of spermatogonia. The area of the sections of wildtype testes and the fields used for the
2 counting were calculated by using ImageJ/Fiji software (Schindelin et al., 2012).

3

4 *Immunohistochemistry and in situ hybridization in testis sections*

5 Immunohistochemistry was performed with slight modifications as described (Kawasaki et al.,
6 2016). Rehydrated sections were antigen retrieved using Immunosaver reagent (Nisshin EM) as
7 manufacturer's instructions, and blocked with EzBlockChemi (Atto) containing 5% BSA
8 (Sigma). Used antibodies and reagents were listed in supplemental Table S6. To analyze Ddx4,
9 Piwil1, Piwil2, Tdrd1 and Tdrd6 localization in *meioc^{mo/mo}* spermatogonia, we used anti-Ddx4
10 IgG labeled with fluorescein using Fluorescein labeling kit – NH2 (Dojindo) and performed
11 double staining with other antibodies because all antibodies were generated in rabbits.

12 *In situ* hybridization of rRNAs was performed with slight modifications as described
13 (Ozaki et al., 2011). To synthesize digoxigenin (DIG) labeled cRNA probes, cDNA of rRNAs
14 were amplified from RT-PCR of testicular RNA using primer sets (Supplemental Table S5), and
15 cloned into the pGEM-T Easy vector (Promega). Twelve loci of precursor 45S rRNA were
16 identified in zebrafish genome (Locati et al., 2017), we designed the specific primer sets that are
17 able to detect rRNAs derived from more than 6 loci. The DIG labeled cRNA probes were
18 synthesized using DIG RNA labeling kit (Roche). The reagents and antibodies were listed in
19 supplemental Table S6.

20 For the fluorescence detection, images were obtained under a FV1000 confocal
21 microscope (Olympus). Overexposed Piwil1 images were acquired under conditions of high
22 detector sensitivity ignoring halation of Piwil1 signals in cytoplasm. The signal strength were

1 quantified using ImageJ/Fiji software (Schindelin et al., 2012). Three testes of wildtype,
2 *meioc^{mo/mo}*, *piwill^{+/-}* and *piwill^{+/-}* were used for the quantification. The signal intensities were
3 normalized to the intensities of neighboring sperm or the intensities of myoid cells of basement
4 membrane of lobules.

5

6 *TdT-mediated nick-end labeling (TUNEL)*

7 TUNEL assays were performed using in situ cell death detection kit (AP; Roche, Germany) as
8 described by the manufacturers. The experiments were repeated three times.

9

10 *Whole-mount in situ hybridization and immunohistochemistry*

11 Whole-mount in situ hybridization and immunohistochemistry were performed as described
12 (Nakamura et al., 2006). A cDNA clone for *meioc* was isolated from RT-PCR of testicular RNA
13 using the primer set (Supplemental Table S5). The digoxigenin (DIG)-labeled RNA probes
14 were synthesized using the *meioc* cDNA and DIG RNA-labeling kit (Roche), and the reagents
15 and antibodies were listed in supplemental Table S6.

16

17 *Protein synthesis assay*

18 Measurement of protein synthesis of germ cells was performed by the Click-iT Plus OPP Alexa
19 Fluor 594 protein synthesis assay kit (Molecular Probes) with slight modifications as described
20 (Sanchez et al., 2016). The fragments of the *meioc^{mo/mo}* and wildtype *sox17:egfp* testes were
21 treated with Leibovitz's L-15 medium (Sigma) containing a 1:400 dilution of Click-iT OPP
22 reagent at 28 °C for 30 minutes. Fluorescence images were acquired with confocal microscope

1 (FV-1000, Olympus). Quantification of OP-Puro fluorescence intensity was performed using
2 ImageJ as described (Sanchez et al., 2016). The signal intensities were normalized to the
3 intensities of neighboring sperm and the myoid cells of basement membrane of lobules. The
4 experiments were repeated three times.

5

6 *Inhibition of ribosome translation in culture*

7 Testis fragments were cultured on the floating membrane in the spermatogonia proliferation
8 medium without growth factors (Kawasaki et al., 2012), with cycloheximide at 0.1 μ M, 1.0 μ M
9 and 10 μ M or the same amount of DMSO as a control for 2 days. Then, BrdU incorporation was
10 analyzed using Cell proliferation kit (GE Healthcare). At least n=3 tissues were examined.

11 Spermatogonia of a *sox17::egfp* hyperplasia testis were cultured as described (Kawasaki
12 et al., 2016). After 1 month of SSC propagation, SSCs were transferred onto the Sertoli cell line
13 ZtA6-12 to induce differentiation, and 0.2 μ M cycloheximide was added. After 9 days, the
14 SSCs (large cells with a few large nucleolus) and the differentiated spermatogonia (small cells
15 with several small nucleolus) were counted according to their morphology and eGFP expression.
16 n=3 dishes were examined.

17

18 *Co-IP*

19 Co-IP was performed with slight modifications as described (Houwing et al., 2007). Testes were
20 homogenized with cell lysis buffer M (Wako) containing cOmplete,Mini Protease Inhibitor
21 cocktail (Roche). One IP generally contained 20 μ l protein G beads (Protein G HP SpinTrap,

1 Cytiva), three testes, and 20µg of antibodies against Meioc or Piwill (Supplemental Table S6)
2 in a total volume of 500µl. The experiments were repeated three times.

3 For RIP, testes were homogenized with 133µl of the lysis buffer containing 100U/ml of
4 SUPERase-In RNase inhibitor (Thermo Fisher Scientific) for 1mg of testis. One IP contains
5 30µl Dynabeads Protein G (Thermo Fisher Scientific), 7.2 µg of anti-Piwill antibody, and
6 250µl of the lysates. Beads and 1% of the lysates were used for Trizol (Thermo Fisher
7 Scientific) RNA isolation. RNA was reverse transcribed using the PrimeScript RT Reagent Kit
8 with gDNA Eraser (Takara), and used for RT-qPCR with TB Green Premix Ex Taq II (Takara)
9 using LightCycler480 (Roche). The thermal cycling were as follows: initial hold for 2 min at
10 95 °C followed by 60 cycles of 30 s at 95 °C, 30 s at 58 °C for primer set for unprocessed
11 pre-rRNA (5'ETS-18S rRNA in supplemental Table S5; (Heyn et al., 2017), and 20 s at 72 °C.
12 Fold enrichment was calculated with -ddCt by normalization with input using Sigma
13 RIP-qRT-PCR Data Analysis Calculation Shell, associated with the Sigma Imprint RIP kit
14 (<http://www.sigmaaldrich.com/life-science/epigenetics/imprint-rna.html>). The experiments were
15 repeated three times.

16

17 *Mass spectrometry*

18 For mass spectrometry, co-IP using anti-Meios IgG and Normal rat IgG were performed using
19 normal testes and hypertrophied testes as described above. We used Dynabeads Protein G
20 (Thermo Fisher Scientific) and the anti-Meios antibody was cross-linked with the beads as
21 described by the manufacturers. The IP was used for Mass spectrometry as described (Ishiguro
22 et al., 2020). The raw LC-MS/MS data was analyzed against the UniProt Knowledgebase

1 restricted to *Danio rerio* using Proteome Discoverer version 1.4 (Thermo Fisher Scientific) with
2 the Mascot search engine version 2.5 (Matrix Science).

3

4 *Construction of expression vectors*

5 For transfection of *Meioc*, *Ythdc2*, *Piwill1*, and *HP1 α* , cDNA fragments encoding zebrafish
6 full-length *meioc*, *ythdc2*, *piwill1*, and *hp1 α* were amplified by RT-PCR using primer sets
7 (Supplemental Table S5). The amplified fragments of *meioc*, *ythdc2* and *piwill1* were subcloned
8 into a pFLAG-CMV-5a expression vector (Sigma-Aldrich) by using *EcoRI* and *SallI*, to create a
9 FLAG tag at the C-terminus of the expressed protein. The amplified fragment was assembled
10 via overlap sequence using NEBuilder HiFi DNA Assembly Master Mix (New England
11 BioLabs). Each 500ng of plasmids were used for transfection to the cells cultured in 35mm
12 dish.

13 For transfection of *Escherichia coli* (Rosetta-gami2 (Novagen)), cDNA fragments
14 encoding zebrafish *dnmt3Aa*, *dnmt3Ab*, *dnmt3Ba*, *dnmt3Bb1*, *dnmt3Bb2*, *dnmt3Bb3*, and *hp1 α*
15 were amplified by RT-PCR using primer sets (Supplemental Table S5), and cloned into the
16 pET-21a (+) vector (Novagen) using NEBuilder HiFi DNA Assembly Mix (New England
17 BioLabs). Expression of cloned cDNAs were induced with 1 μ M Isopropyl- β
18 -D-thiogalactopyranoside (IPTG, Wako).

19

20 *Pull-down assay*

21 The expression vectors were transfected into HEK-293 cells using Lipofectamine 3000
22 Transfection Reagent (Thermo Fisher Scientific). After 48 h, cells were harvested and used for

1 immunoprecipitation as described above. We used anti-Flag M2 agarose resin (Sigma). The
2 experiments were repeated three times.

3

4 *Small RNA-seq library preparation and sequencing*

5 Total RNA from 6 *meioc*^{+/*mo*} and 5 *meioc*^{*mo/mo*} testis samples at 10 months old was isolated using
6 Trizol RNA isolation according to manufacturer's instructions (Thermo Fisher Scientific).
7 RNAs smaller than 40nt were isolated using a 15% TBE-Urea polyacrylamide gel (BioRad),
8 and purified with sodium chloride/isopropanol precipitation. NGS library preparation was
9 performed using the NEBNext Multiplex Small RNA Library Prep Set for Illumina (New
10 England BioLabs) as recommended by the manufacturer (protocol version v2.0 8/13), with 14
11 PCR cycles for library amplification. The PCR-amplified DNA was purified using AMPure XP
12 beads (Beckman Coulter). Size selection of the small RNA library was done on LabChip XT
13 instrument (Perkin Elmer) using a DNA 300 assay kit. The library fractions in the range
14 120-161 bp were pooled in equal molar ratio. The resulting 2nM pool was denatured and diluted
15 to 10 pM with 5% PhiX spike-in DNA and sequenced (single read, 51 cycles, high output
16 mode) on 2 lanes of HiSeq 2000 system (Illumina).

17

18 *Small RNA-seq data analysis*

19 The raw NGS reads in FastQ format were cleaned from partial 3' adapter sequences using
20 Flexbar v.2.4 (Dodt et al., 2012) using parameters: *-m 18 -ao 10 -as*
21 *AGATCGGAAGAGCACACGTCTGAACTCCAGTCAC*. Read mapping to the *Danio rerio*
22 reference genome (Zv9/danRer7 build from Illumina iGenomes) was carried out using Bowtie

1 v.0.12.8(Langmead et al., 2009) with parameters: *-n 0 -e 80 -l 18 -y --best --nomaqground*. Reads
2 were assigned to the predicted zebrafish 28S and 18S rRNAs, defined as 4110 nt and 1887 nt
3 sequences from GenBank records CT956064 and BX537263 with highest Megablast homology
4 to the respective rRNAs from *Cyprinus carpio* (GenBank JN628435), as well as to the zebrafish
5 R2 transposon (GenBank AB097126), using Bowtie with perfect match parameters (*-v 0 -m 1*).
6 Quality assessment of the raw data and length profiling of the mapped reads was performed
7 with FastQC (<http://www.bioinformatics.babraham.ac.uk/projects/fastqc>). The sequence data
8 have been deposited in the NCBI GEO repository (<http://www.ncbi.nlm.nih.gov/geo>) under
9 accession number GSE84060. [The temporal token (password) for reviewers access to the GEO
10 record before publication is: wjunoeqmxidpep]

11

12 *Inhibition of RNA polymerases*

13 Fragments of *meioc^{mo/mo}* testes were cultured in the spermatogonia proliferation medium without
14 growth factors (Kawasaki et al., 2012). Actinomycin D (Wako) at 1 µg/ml, α-amanitin
15 (Bensaude, 2011) at 10 µg/ml and BMH-21 (Sigma) (Colis et al., 2014) at 1 µM were added.
16 After 1 hr incubation at 28° in 5% CO₂/20% O₂, testicular samples were fixed and analyzed by
17 immunohistochemistry for Piwill1. The experiments were repeated three times.

18

19 *In situ hybridization and immunocytochemistry of isolated spermatogonia*

20 After sorting spermatogonia of *sox17::egfp* wild-type that expresses EGFP at the equivalent
21 stage to *meioc^{mo/mo}* spermatogonia (Kawasaki et al., 2012) using JSAN cell sorter (Bay
22 Bioscience), cells were plated on the CREST-coated glass slides (Matsunami) for 10 minutes,

1 and fixed with 4% paraformaldehyde for 10 minutes. The cells were treated with 0.5% Triton
2 X100 in PBS for 5 minutes. To detect 28S rRNA, the cells were acetylated with 0.025% acetic
3 anhydrid in triethanolamine 10mM for 5 minutes and followed as described above. After
4 hybridization, antibodies and reagents (Supplemental Table S6) were used. Images were
5 obtained under a FV1000 confocal microscope (Olympus). The signal strength of 28S rRNA,
6 Meioc and Piwil1 was quantified using ImageJ/Fiji software (Schindelin et al., 2012). The
7 signal intensities of in situ hybridization (28S rRNA) and immunocytochemistry (Meioc and
8 Piwil1) were normalized to the intensities of S probe and control IgG, respectively.

9

10 *DNA methylation analysis*

11 Because there was no information of the structure of IGS region in zebrafish, we analyzed IGS
12 sequence of known active locus of 45S-S rDNA (Ch5: 831,755-826,807 in GRCz11) (Locati et
13 al., 2017) using Tandem repeats finder software (Benson, 1999). The result was summarized in
14 the schema of Supplemental Figure S3B. The undifferentiated spermatogonia of
15 *meioc^{mo/mo};sox17::egfp* and *sox17::egfp* wildtype were sorted using JSAN cell sorter (Bay
16 Bioscience) and used for bisulfite conversion using MethylEasy Xceed Rapid DNA Bisulfite
17 modification kit (Takara). The tandem repeat region of 45S-S rDNA IGS was amplified by PCR
18 with EpiTaq HS (Takara) using the primer sets (Supplemental Table S5). The primers were
19 designed using Meth primer software (Li and Dahiya, 2002). All PCR products were subcloned
20 into the pCRII vector (Thermo Fisher Scientific) and used for sequencing analysis using online
21 QUMA software (Kumaki et al., 2008).

22

1 *Comparison of RNA expression levels*

2 After sorting spermatogonia of *vas::egfp; meioc^{mo/mo}* and *sox17::egfp* wildtype that expresses
3 EGFP at the equivalent stage to *meioc^{mo/mo}* spermatogonia using JSAN cell sorter (Bay
4 Bioscience), total RNA were extracted and used for RT-qPCR analysis as described in
5 “Immunoprecipitation” section using primer sets (Supplemental Table S5). Relative gene
6 expression levels were calculated using the comparative Ct method (Schmittgen and Livak,
7 2008) and normalized to the expression of *gapdh*. The experiments were repeated three times.

8

9 *Northern blot analysis*

10 Northern blot analysis was performed using DIG northern starter kit (Roche). cDNA fragments
11 of 5'ETS and ITS1 region of 45S-S rDNA(Locati et al., 2017) and 7SL were amplified from
12 RT-PCR of testicular RNA using the primers containing T7 and T3 promoter sequence
13 (Supplemental Table S5) and used for cRNA probe synthesis with T3 RNA polymerase using
14 DIG RNA labeling kit (Roche). Signals of 45S pre-rRNA, pre-rRNA intermediates and 7SL
15 were detected with a Chemidoc XRS Plus (Bio-Rad), and quantified using quantity tools of
16 ImageLab software version 6.0.1 (Bio-Rad). For the quantification, three wildtype and
17 *meioc^{mo/mo}* testes were individually used. Signal intensities of 45S pre-rRNA, pre-rRNA
18 intermediates were normalized with signal intensities of 7SL.

19

20 *Chromatin IP (ChIP)-qPCR analysis*

21 Chip assay was performed with slight modifications as described (Imai et al., 2017). Ten testes
22 were used for 6 ml of chromatin suspension. Sonication was carried out with a Bioruptor®

1 Standard apparatus (Diagenode) at high power for four series of 7 cycles (30 s on, 30 s off). For
2 IP, 1 ml of chromatin suspension was incubated with 20 μ l of Dynabeads® Protein A (Thermo
3 Fisher Scientific) pre-incubated with 3 μ g of rabbit anti-histone H3 tri-methyl K9 (Abcam,
4 ab8898) and Piv11 antibodies. Quantitative PCR (qPCR) was performed with SYBR Premix
5 Ex Taq II (Takara) using LightCycler480 (Roche) using primer sets (Supplemental Table S5).
6 Fold enrichment was calculated with -ddCt by normalization with 10% input sample using
7 Sigma RIP-qRT-PCR Data Analysis Calculation Shell, associated with the Sigma Imprint RIP
8 kit (<http://www.sigmaaldrich.com/life-science/epigenetics/imprint-rna.html>). The experiments
9 were repeated six times.

10

11 **Quantification and Statistical analysis**

12 Data were presented as the mean \pm standard deviation of at least three independent experiments
13 as indicated in each method and figure legend. Statistical difference between two groups was
14 determined using unpaired Student's *t*-test when the variance was heterogeneous between the
15 groups, and Welch's *t*-test was used when the variance was heterogeneous. $P < 0.05$ was
16 considered statistically significant. Graphical presentations were made with the R package
17 ggplot2 (Wickham, 2009).

18

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20

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10

11 **Author Contributions**

12 Toshihiro Kawasaki, Kellee R. Siegfried and Noriyoshi Sakai conceived and designed the work.
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15 R. Siegfried and Noriyoshi Sakai performed the experiments and analyzed the data. Toshihiro
16 Kawasaki, Kellee R. Siegfried and Noriyoshi Sakai wrote the manuscript.

17

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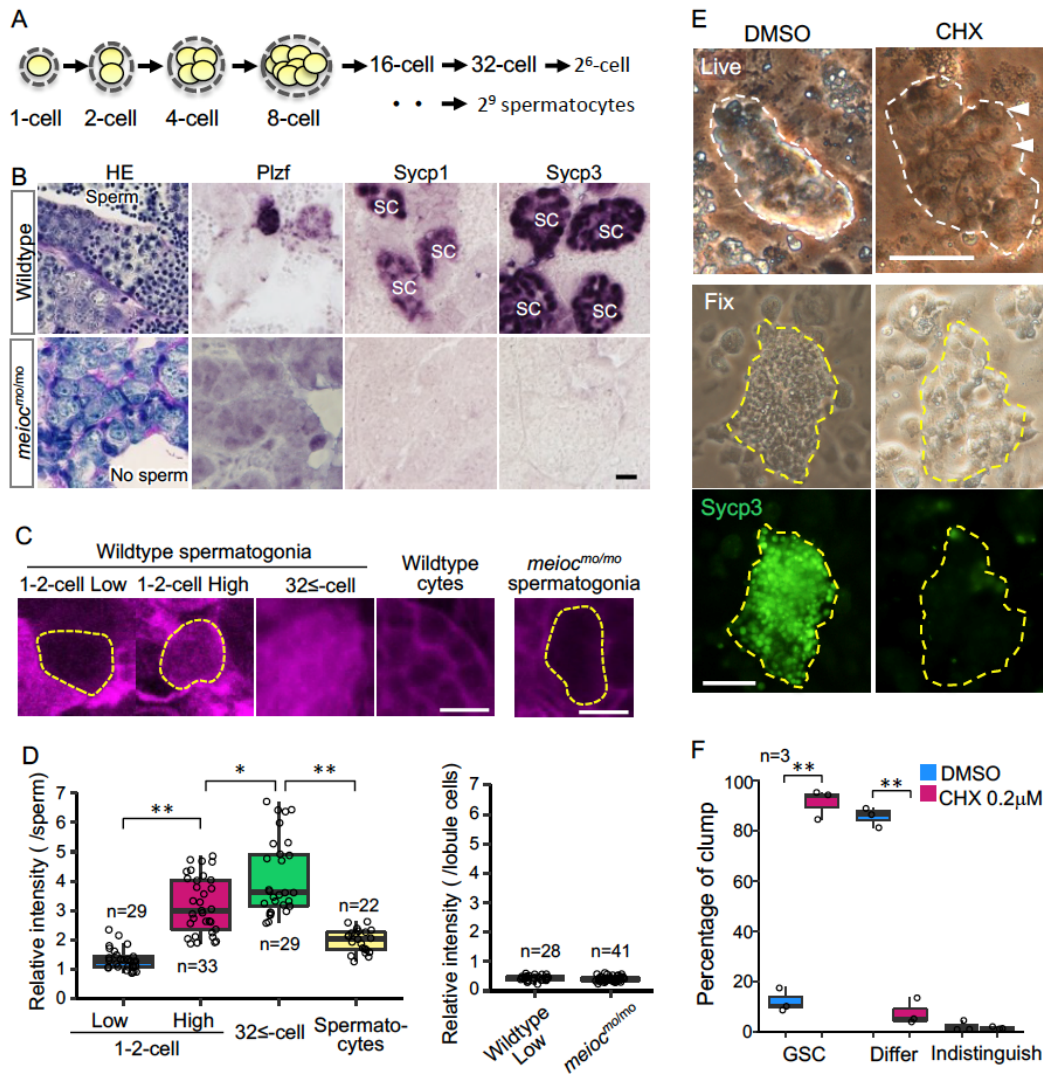
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1 **Figure legends**

2



3

4 **Figure 1. Low translational activity of *meioc* mutant spermatogonia**

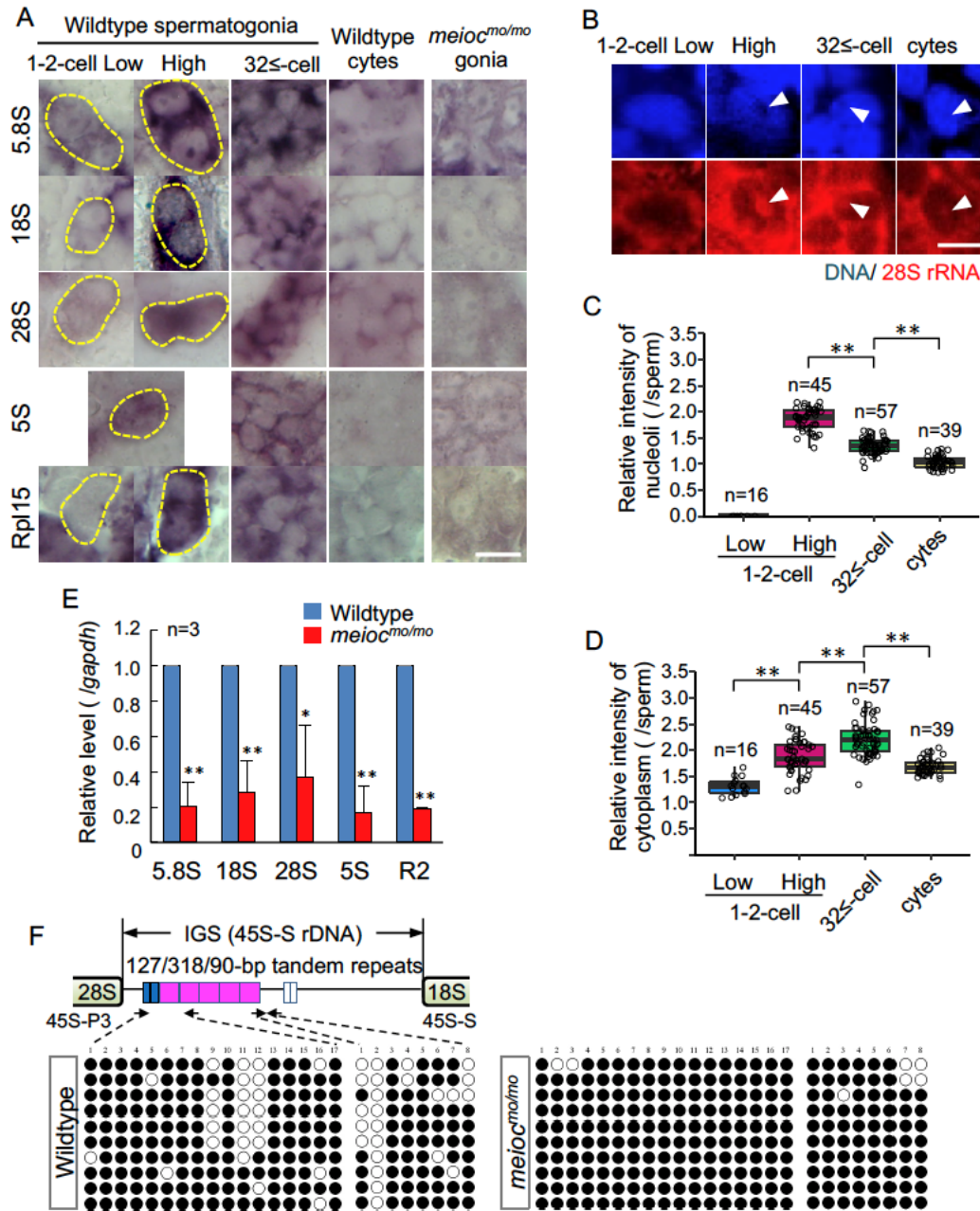
5 (A) Schema of the development of spermatogonial cysts surrounded by Sertoli cells in zebrafish.

6 (B) Histology (HE) and immunostaining against Plzf and spermatocyte markers (Sycp1, 3) in

7 the wildtype and the *meioc*^{mo/mo} testes. Scale bar: 10 μm. (C, D) OP-Puro fluorescence analysis (C)

8 and quantification of the signal intensities (D) in wildtype and *meioc*^{mo/mo} spermatogenic cells.

1 Scale bars: 10 μm . (E-F) Effect of cycloheximide (CHX, 0.2 μM) on differentiation of SSCs in
2 culture. Yellow dotted lines; germ cell clumps. Sycp3; Immunostaining of Sycp3. Arrowheads;
3 examples of a cell with a large nucleolus. The graph (F) presents the percentage of clumps of
4 SSCs and differentiated cells (Differ) shown in Panel E. Scale bar; 50 μm . Data are represented
5 as mean \pm SD.
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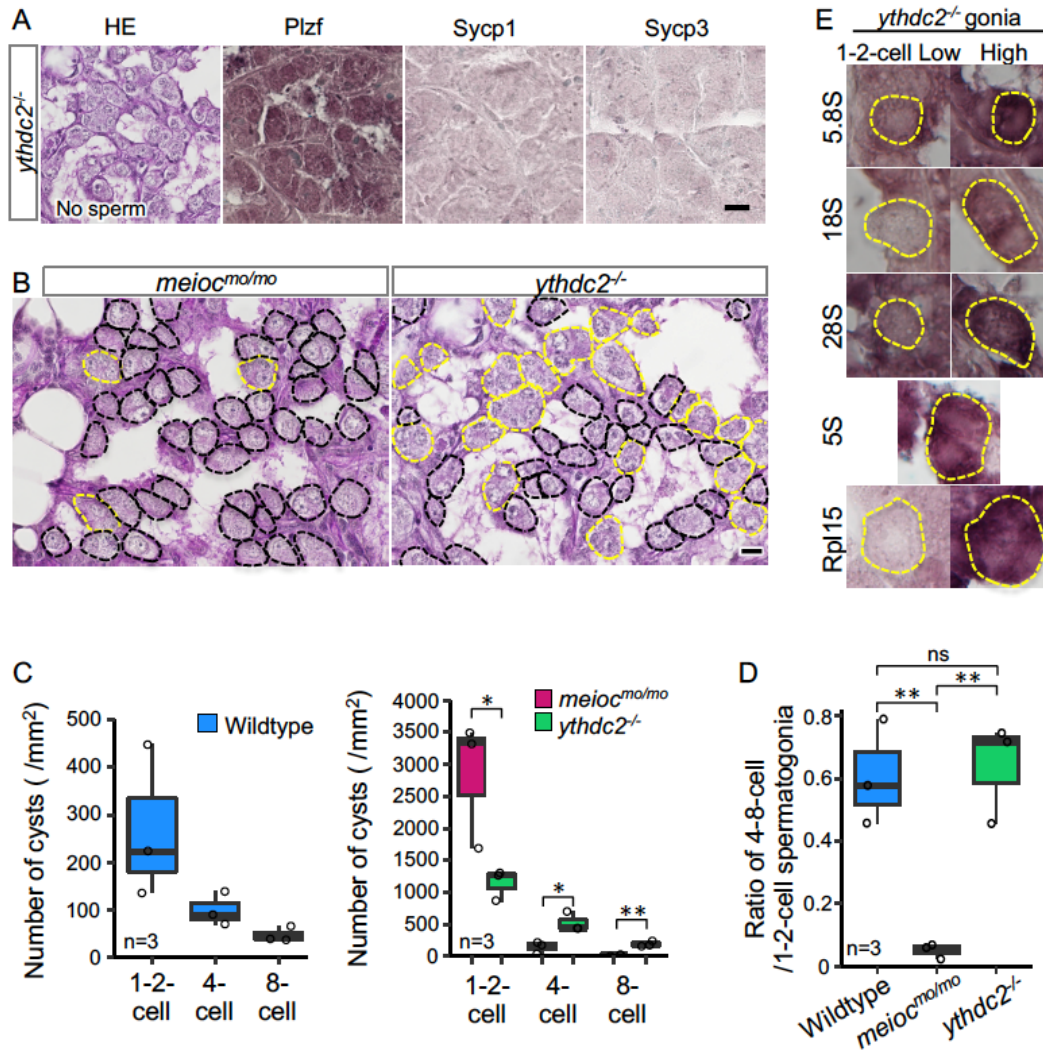


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2 **Figure 2. Defect on upregulation of rRNA transcription in *meioc^{mo/mo}* spermatogonia**

3 (A) In situ hybridization of 5S, 5.8S, 18S, and 28S rRNA and immunohistochemistry with
4 anti-Rpl15 antibody in spermatogonia (gonia) and spermatocytes (cyte) in wildtype and
5 *meioc^{mo/mo}*. Yellow dotted lines indicate 1-2-cell spermatogonia. (B-D) Fluorescent in situ

1 hybridization of 28S rRNA in wildtype. The graphs present quantification of signal intensities
2 of 28S rRNA in nucleoli (C) and in cytoplasm (D) in spermatogenic cells. Arrowheads; nucleoli.
3 (E) qRT-PCR analysis of rRNAs and R2 between wildtype and *meioc^{mo/mo}* purified
4 undifferentiated spermatogonia. (F) Bisulfite-sequencing analysis of the tandem repeat region in
5 the IGS region of the 45S-S rDNA locus in purified undifferentiated spermatogonia of wildtype
6 and *meioc^{mo/mo}*. Arrows; position of bisulfite primers in the tandem repeat elements (blue,
7 magenta and white boxes), black dots; methylated CpG sites, white dots; unmethylated sites. *p
8 < 0.05, **p < 0.01. Scale bars: 10 μ m.
9



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2 **Figure 3. *ythdc2* mutant spermatogonia have different defects from *meioc^{mo/mo}*.**

3 (A) Histology (HE) and immunostaining against Plzf and spermatocyte markers (Sycp1, 3) in

4 the *ythdc2^{-/-}* testes. (B) Representative image of *meioc^{mo/mo}* and *ythdc2^{-/-}* testes sections stained

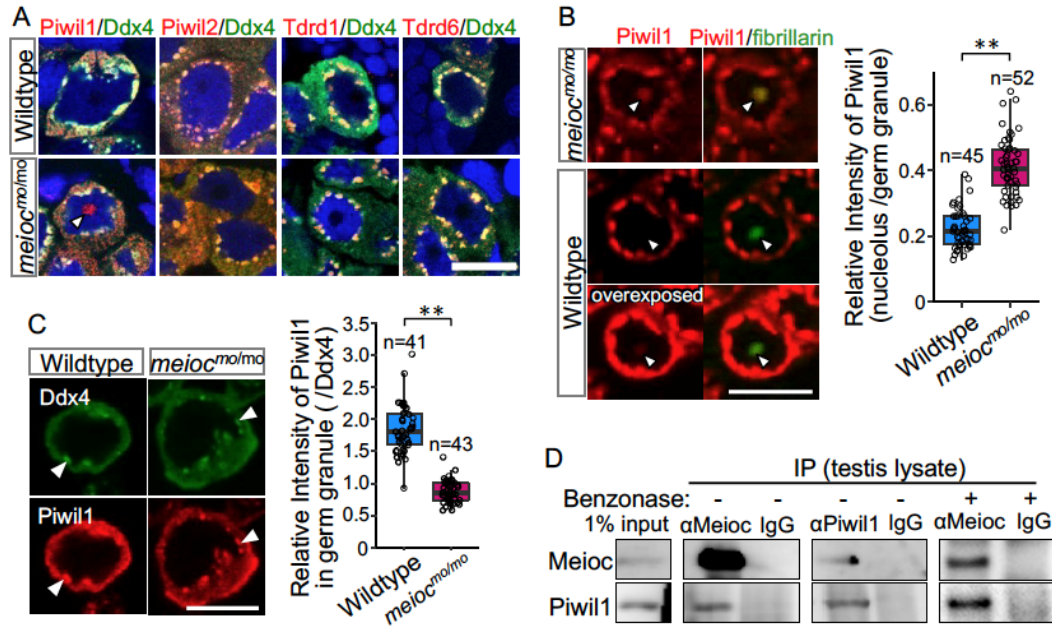
5 with PAS (periodic acid Schiff) and hematoxylin. Dotted lines; 1-2-cell cyst spermatogonia

6 (black) and 4≤-cell cysts (yellow). (C) The number of 1-2-, 4-, and 8-cell cyst spermatogonia

7 per mm² of sections in wildtype, *meioc^{mo/mo}* and *ythdc2^{-/-}* testes. (D) Ratio of the number of

8 4-8-cell cyst spermatogonia to 1-2-cell cysts in wildtype, *meioc^{mo/mo}* and *ythdc2^{-/-}* testes. (E) In

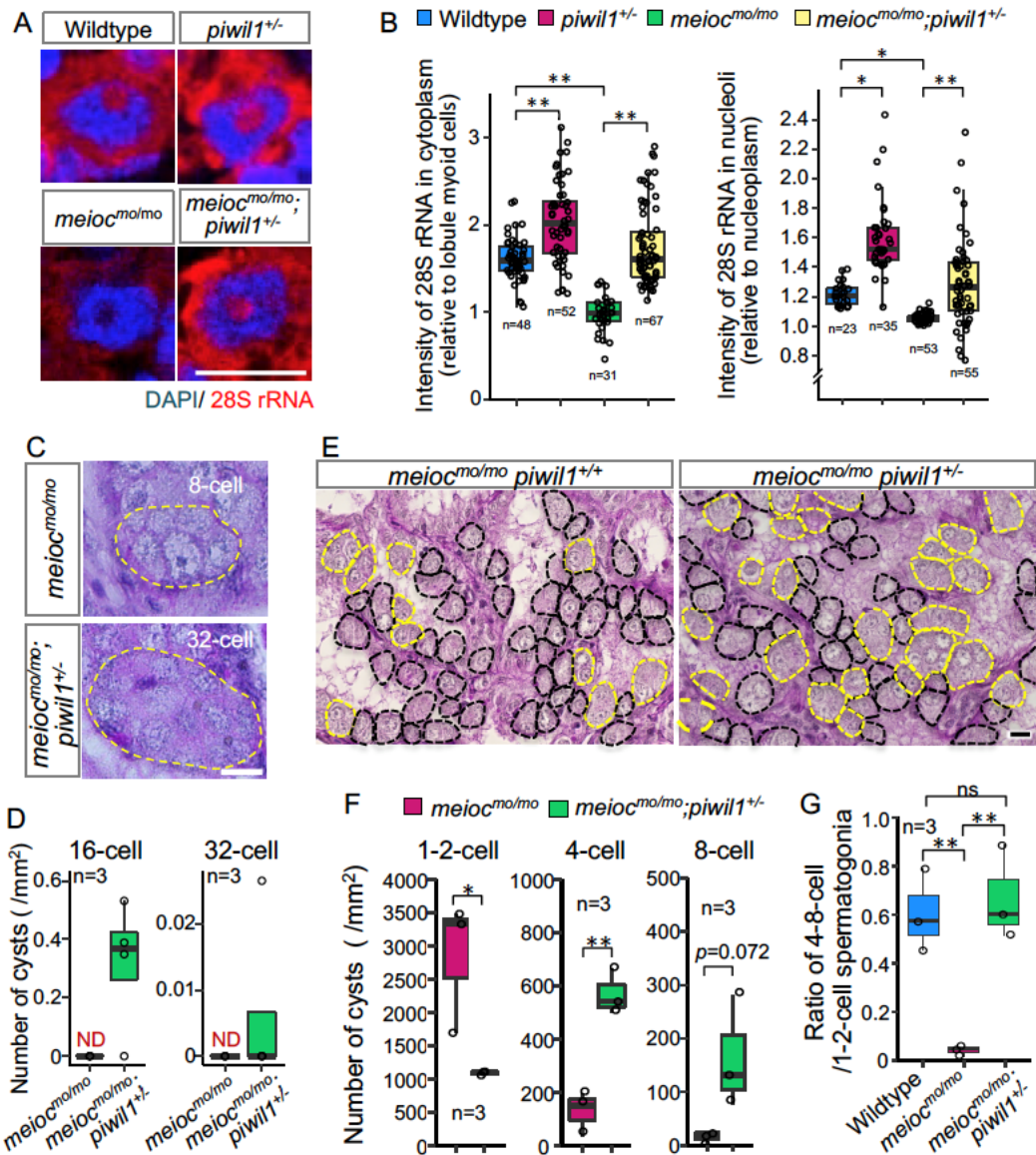
- 1 situ hybridization of 5S, 5.8S, 18S, and 28S rRNA and immunohistochemistry with anti-Rpl15
- 2 antibody in the *ythdc2*^{-/-} testes. Yellow dotted lines; 1-2-cell spermatogonia. *p < 0.05, **p <
- 3 0.01. Scale bars: 10 μm.
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Figure 4. Meioc directly binds with Piwil1 and affects the localization of Piwil1.

(A) Immunostaining of Ddx4 and Piwil1, Piwil2, Tdrd1 and Tdrd6 in wildtype and *meioc^{mo/mo}* spermatogonia. The arrowhead; the Piwil1 signal in the nucleolus. (B) Immunostaining against Piwil1 and fibrillarin (left panels) and quantification of nucleolar Piwil1 (right panel) in wildtype and *meioc^{mo/mo}* spermatogonia. Arrowheads; Fibrillarin positive nucleolus. (C) Immunostaining of Piwil1 and Ddx4 (left panels) and quantification of Piwil1 in germ granules (right panel) in wildtype and *meioc^{mo/mo}* spermatogonia. Arrowheads; Ddx4 positive germ granules. (D) Coimmunoprecipitation of Meioc and Piwil1 using testis lysate. Meioc signals were detected in Piwil1 immunoprecipitate and vice versa. Benzoylase: addition of benzoylase nuclease. ***p* < 0.01. Scale bars: 10 μ m.



1

2 **Figure 5. Reduction of Piwil1 compensated phenotypes of *meioc*^{mo/mo}.**

3 (A, B) In situ hybridization of 28S rRNA in wildtype and *meioc*^{mo/mo}; *piwil1*^{+/-} spermatogonia

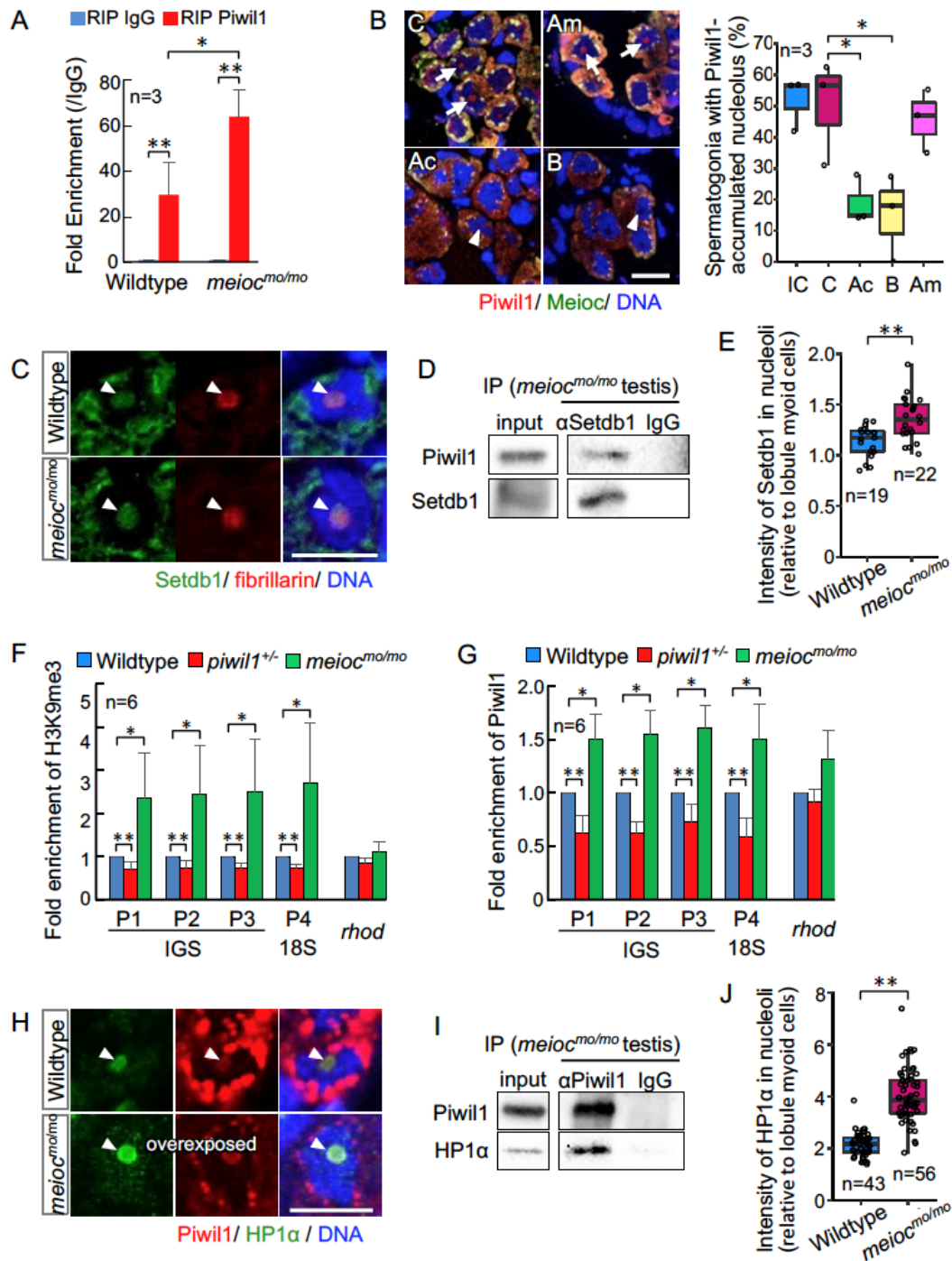
4 (1-2-cell cysts). Graphs (B) show the relative signal intensity in the cytoplasm normalized to the

5 intensity of lobule myoid cells (left) and nucleoli normalized to the intensity of the nucleoplasm

6 (right). (C, D) Differentiated spermatogonia in *meioc*^{mo/mo} and *meioc*^{mo/mo}; *piwil1*^{+/-} testes. Yellow

7 dotted lines; differentiated spermatogonia. Graphs (D) show the number of 16-cell and 32-cell

1 cyst spermatogonia per mm² of sections. ND: not detected. (E-G) *meioc^{mo/mo}* and *meioc^{mo/mo}*
2 *piwill^{+/-}* testis sections stained with PAS and hematoxylin. Cysts of 1-2-cell spermatogonia
3 (black) and 4≤-cell cysts (yellow) are indicated by dotted lines. Graphs show numbers of 1-, 2-,
4 4-, and 8-cell cysts per mm² in sections of *meioc^{mo/mo}* and *meioc^{mo/mo};piwill^{+/-}* testes (F), and
5 ratio of the number of 4-8-cell cysts to 1-2-cell cysts in wildtype, *meioc^{mo/mo}* and
6 *meioc^{mo/mo};piwill^{+/-}* (G). *p < 0.05, **p < 0.01. Scale bars: 10 μm.
7



1

2 **Figure 6. Nucleolar Piwil1 interacted with Setdb1 and causes silenced epigenetic state of**

3 **rDNA loci**

1 (A) Fold enrichment of pre-rRNA (5'ETS-18S rRNA) in Piwil1 immunoprecipitated RNA
2 relative to the control IgG in wildtype and *meioc^{mo/mo}* testes. (B) Immunostaining of Piwil1 (left
3 panels) and the percentage of spermatogonia with detectable nucleolar Piwil1 (right panel) in
4 the *meioc^{mo/mo}* testes treated with α -amanitin (Am), actinomycin D (Ac) and BMH-21 (B).
5 Arrows; Piwil1 detectable nucleoli, arrowheads; Piwil1 undetectable nucleoli, C; control
6 without inhibitors, IC; initial control. (C) Immunostaining of Setdb1 and fibrillarin in wildtype
7 and *meioc^{mo/mo}* spermatogonia. Arrowheads; nucleoli. (D) Co-IP of Piwil1 and Setdb1 using
8 *meioc^{mo/mo}* testes lysate. Piwil1 was detected in Setdb1 IP. (E) Intensities of Setdb1 in nucleoli
9 in wildtype and *meioc^{mo/mo}* spermatogonia. (F, G) ChIP-qPCR analysis of H3K9me3 (F) and
10 Piwil1 (G) levels in 45S-rDNA region in wildtype, *piwil1^{+/-}*, and *meioc^{mo/mo}* testes. Position of
11 primers were indicated in Supplemental Figure S3. Mean \pm s.d. are indicated. (H)
12 Immunostaining of HP1 α and Piwil1 in wildtype and *meioc^{mo/mo}* spermatogonia. Arrowheads;
13 nucleolus. (I) Co-IP of Piwil1 and HP1 α using *meioc^{mo/mo}* testis lysate. HP1 α was detected in
14 Piwil1 IP. (J) Intensities of HP1 α in nucleoli in wildtype and *meioc^{mo/mo}* spermatogonia. *p <
15 0.05, **p < 0.01. Scale bars: 10 μ m.

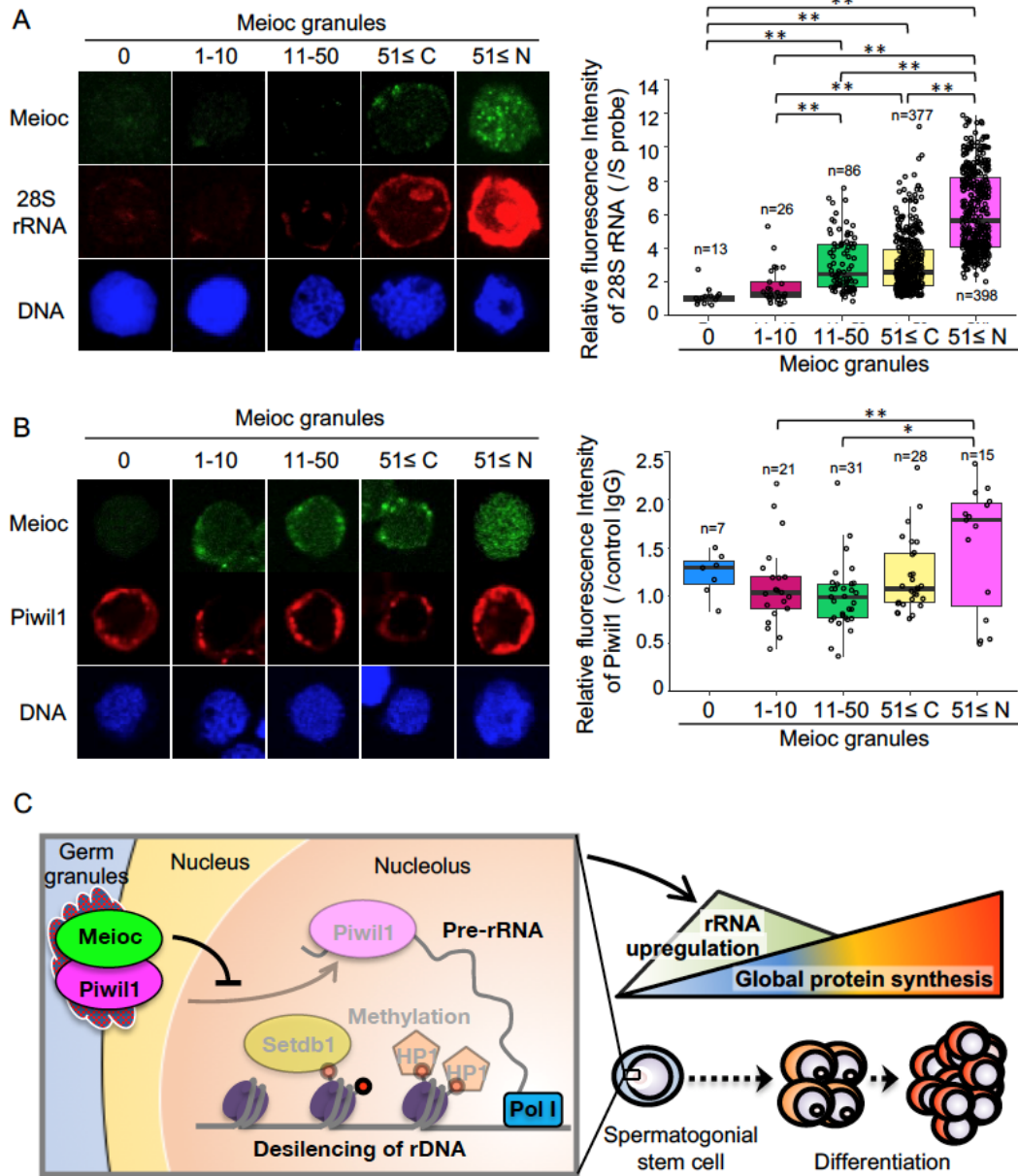


Figure 7. Meioc was required for upregulation of 28S rRNA

(A, B) Expression patterns of 28 rRNA (A) and Piwil1 (B) in isolated *sox17::egfp* spermatogonia, based on the amount of Meioc granules and the localization. Right panels are intensities of 28 rRNA (A) and Piwil1 (B) for each class of the purified wildtype 1-2-cell spermatogonia. 51≤ C and N; 51≤ cytoplasmic and nuclear Meioc granules, respectively. * $p <$

- 1 0.05, **p < 0.01, n: number of analyzed spermatogonia. Scale bars: 10 μ m. (C) Graphical
- 2 abstract. Meioc prevents the nucleolar localization of Piwil1 and its associated Setdb1 and
- 3 HP1 α to upregulate rRNA transcripts that are required for zebrafish SSCs to differentiate.