Zucchini-dependent piRNA processing is triggered by recruitment to the cytoplasmic processing machinery

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The piRNA pathway represses transposable elements in the gonads and thereby plays a vital role in protecting the integrity of germline genomes of animals. Mature piRNAs are processed from longer transcripts, piRNA precursors (pre-piRNAs). In Drosophila, processing of pre-piRNAs is initiated by piRNA-guided Slicer cleavage or the endonuclease Zucchini (Zuc). As Zuc does not have any sequence or structure preferences in vitro, it is not known how piRNA precursors are selected and channeled into the Zuc-dependent processing pathway. We show that a heterologous RNA that lacks complementary piRNAs is processed into piRNAs upon recruitment of several piRNA pathway factors. This processing requires Zuc and the helicase Armitage (Armi), Aubergine (Aub), Argonaute 3 (Ago3), and components of the nuclear RDC complex, which are required for normal piRNA biogenesis in germ cells, are dispensable. Our approach allows discrimination of proteins involved in the transcription and export of piRNA precursors from components required for the cytoplasmic processing steps. piRNA processing correlates with localization of the substrate RNA to nuage, a distinct membraneless cytoplasmic compartment, which surrounds the nucleus of germ cells, suggesting that sequestration of RNA to this subcellular compartment is both necessary and sufficient for selecting piRNA biogenesis substrates.

Keywords: piRNA biogenesis; nuage; Zucchini; tethering; Piwi; RNA sequestration

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How piRNA precursors are identified by the biogenesis machinery remains unresolved. Other RNA processing events, such as microRNA (miRNA) or CRISPR RNA biogenesis, splicing, or polyadenylation, require specific sequence and/or structure motifs in precursor RNA that are recognized by the processing machinery [Zeng et al. 2005; Park et al. 2011; Barrangou and Marraffini 2014; Li and Patel 2016; Tsai and Joung 2016]. Common sequence or structural motifs that are shared by all piRNA precursors were not identified [Muerdter et al. 2012]. In the absence of sequence motifs, two mutually nonexclusive models have been proposed to explain precursor selection in the germline, which we call the “persistent nuclear mark model” and the “selection by pre-existing piRNA model.”

The “persistent nuclear mark model” proposes that specific proteins associate with piRNA precursors in the nucleus and remain associated with the transcripts in the cytoplasm, where they interact with the processing machinery. In the germline, genomic regions that give rise to the majority of piRNAs, called piRNA clusters, are marked by the histone 3 Lys9 trimethylation [H3K9me3] mark and the RDC complex. RDC consists of the HP1 homolog Rhino and two additional proteins, Cutoff (Cuff) and Deadlock, and is required for transcription and early processing of cluster transcripts [Klattenhoff et al. 2009; Pane et al. 2011; Le Thomas et al. 2014; Mohn et al. 2014; Zhang et al. 2014; Chen et al. 2016]. The well-conserved transcription export (TREX) complex is also enriched at cluster loci and binds piRNA precursors cotranscriptionally in an RDC-dependent fashion [Zhang et al. 2012; Hur and Chung 2016; Hur et al. 2016]. Components of either the RDC or the TREX complex were proposed to constitute the mark that triggers processing in the cytoplasm. However, the nature of the mark and the mechanism by which it engages the processing machinery remain unclear. Evidence for RDC or TREX remaining associated with piRNA precursors after nuclear export is also lacking. Finally, tethering of Rhino to a single-stranded transgene {without its concomitant tethering to a complementary antisense transgene} does not trigger piRNA biogenesis [Zhang et al. 2014], arguing against the idea that binding of RDC by itself is sufficient to specify piRNA precursors.

The “selection by pre-existing piRNA model” suggests that precursors are specified in the cytoplasm by complementary piRNAs associated with the cytoplasmic piwi proteins. Existing piRNAs can target pre-piRNAs and induce their processing via the ping-pong cycle [Brennecke et al. 2007; Gunawardane et al. 2007]. piRNA-dependent cleavage not only generates complementary piRNAs (ping-pong partners) but also leads to phased processing of the precursor downstream from the initial cleavage [Han et al. 2015; Homolka et al. 2015; Mohn et al. 2015; Senti et al. 2015; Wang et al. 2015]. This process generates Piwi-loaded piRNAs. Mutation of the two piwi proteins involved in the ping-pong cycle, Aub and Ago3, greatly reduces Piwi-bound piRNAs in the germline, leading to the suggestion that Zuc-mediated piRNA biogenesis in germ cells is triggered by the ping-pong cycle [Han et al. 2015; Homolka et al. 2015; Mohn et al. 2015; Senti et al. 2015; Wang et al. 2015].

Follicular cells of the fly ovary lack ping-pong biogenesis factors as well as the RDC complex yet produce piRNAs, suggesting an alternative precursor selection mechanism. Some piRNA precursors specific to follicular cells, such as the mRNA of traffic-jam [tj] and transcripts from the flamenco piRNA cluster, harbor sequences that are bound by the RNA-binding protein Yb [Ishizu et al. 2015] and can trigger processing into piRNAs [Homolka et al. 2015; Ishizu et al. 2015]. Whether other somatic piRNA precursors also harbor sequence motifs that are bound by Yb and whether binding of Yb leads to piRNA processing remain to be elucidated.

Genetically, many factors have been identified to be essential for piRNA biogenesis; however, a mechanistic dissection of which steps these factors are involved in {i.e., transcription, export, recognition, or cytoplasmic processing} is largely missing. Assigning many of the identified factors to specific steps of the biogenesis pathway is hindered by the absence of an in vitro system that allows independent analysis of different stages of the pathway. In vivo analysis is impaired because disrupting proteins acting in early steps of the pathway, such as transcription of pre-piRNAs, eliminates piRNA precursors and thereby the possibility of testing whether these factors also act in downstream steps of processing.

The core of the cytoplasmic processing machinery, including the endonuclease Zuc, is shared by nurse and follicular cells, raising the question of how the same processing machinery has evolved to recognize its targets by two distinct {sequence-specific in follicular cells and sequence-independent in nurse cells} mechanisms. We show that direct recruitment of a heterologous sequence to the cytoplasmic piRNA processing machinery can bypass the necessity for sequence-specific recognition, as well as the need for both complementary piRNAs and a unique chromatin architecture. Our work reveals that sequestration of RNA to the cytoplasmic processing machinery is both necessary and sufficient for selecting substrates for piRNA biogenesis, leading to a unified model of piRNA specification in the two cell types. The described experimental approach bypasses nuclear biogenesis steps, enabling identification and functional dissection of factors that act in Zuc-mediated processing downstream from nuclear export.

Results

Recruitment of Yb and Piwi to RNA in follicular cells leads to piRNA processing

Two transcripts that are processed into piRNAs in follicular cells, the mRNA of tj and transcripts from the flamenco piRNA cluster, were shown to harbor sequences that associate with the Yb protein and trigger piRNA processing when transferred to heterologous transcripts [Homolka et al. 2015; Ishizu et al. 2015]. To determine whether the interaction of a transcript with Yb is
sufficient to induce processing, we tested whether we could trigger piRNA production in follicular cells by tethering a heterologous sequence that does not have sequence homology with natural piRNA substrates to Yb (Fig. 1A). The sequence of the reporter mRNA contains four BoxB sequences in its 3’ untranslated region (UTR) that are bound by the λN domain of anti-terminator protein N fused to Yb (Fig. 1A). We coexpressed λN-tagged Yb with the reporter in follicular cells of the fly ovary and sequenced ovarian small RNAs. Tethering of the transcript to Yb resulted in production of reporter-derived small RNAs, while small RNAs were absent upon recruitment of the control GFP protein (Fig. 1B, C). The majority of small RNAs produced from the reporter upon Yb tethering was 25–28 nt long and had a U bias at the first nucleotide (82.2% of all reporter-derived reads had a 5’U), consistent with these sequences being piRNAs [Fig. 1D, E]. Thus, recruitment of Yb to a novel transcript is sufficient to trigger piRNA processing in follicular cells in the absence of any sequence or structure motifs in the RNA.

In follicular cells, piRNAs are loaded into Piwi, the only member of the piwi clade expressed in these cells. Therefore, we asked whether recruitment of Piwi itself to RNA is sufficient to trigger processing. Tethering of Piwi also induced generation of piRNAs from the reporter [Fig. 1B–E]. We note that the piRNA profiles along the reporter differ depending on the factor recruited. Piwi recruitment leads to processing of the entire transcript, while Yb recruitment leads to piRNA biogenesis only downstream from the BoxB sites, to which the Yb fusion protein binds [Fig. 1C]. This is similar to the piRNAs observed in the tj mRNA and when using the recognition sequence from flamenco, both of which lead to processing only downstream from the recognition sequence [Homolka et al. 2015; Ishizu et al. 2015]. Our data indicate that Yb binding not only identifies piRNA precursors but also specifies the entry site for processing. Thus, in follicular cells, recruitment of either Yb or Piwi is sufficient to trigger piRNA processing from a novel substrate, and Yb might be responsible for specifying where processing initiates within the RNA.

Recruitment of the nuclear RDC and TREX complexes or Zuc is not sufficient to trigger piRNA generation in germ cells

In the germline, piRNA biogenesis is more complex than in somatic follicular cells and depends on numerous nuclear and cytoplasmic factors. Two nuclear complexes—RDC and TREX, which localize to chromatin of piRNA loci and nascent pre-piRNA, respectively—were proposed to play a role in selecting piRNA precursors and guiding them to the processing machinery in germ cells. To test whether the recruitment of RDC or TREX to the heterologous transcript is sufficient to trigger piRNA processing in germ cells, we tethered the RDC component Cuff and the TREX component Thoc5 to the reporter in the female germline and profiled ovarian small RNAs. Tethering of these factors did not trigger piRNA generation [Fig. 2A, B], suggesting that recruitment of RDC or TREX is not sufficient to trigger biogenesis from novel RNA substrates in germ cells.

**Figure 1.** Yb and Piwi recruitment to RNA in follicular cells triggers piRNA production from reporter mRNA. [A] Schematic diagram of the reporter construct used to study the effect of piRNA pathway component recruitment to RNA. [PAS] PolyA signal. [B] Recruitment of Yb and Piwi, but not GFP, to a reporter results in small RNA reads mapping to the reporter sequence. Shown are small RNA sequencing reads from 19- to 30-nt total ovarian RNA. RPM (reads per million mapped reads) was calculated for reads mapping exclusively to the reporter, normalized to total reads mapping to the DM3 genome. Error bars indicate standard deviation between replicates. n = 2. [C] The small RNA reads produced upon tethering Yb or Piwi map to the reporter. Shown are profiles for all reads [black] and reporter-specific reads [red] mapping to the reporter, normalized to all reads mapping to the DM3. [D] Tethering of Yb or Piwi results in production of piRNA-length reads from the reporter. Shown are size profiles of all reads mapping to the reporter. Error bars indicate standard deviation between replicates. n = 2. [E] Reporter-derived sequences upon Yb or Piwi tethering exhibit a 1U bias. The Weblogos were generated from all reads mapping to the reporter.
piRNA biogenesis induced by RNA tetherno. However, upon NaIO4 treatment, reporter-derived small RNAs generated upon Zuc tethering revealed the presence of abundant small RNAs arising from the reporter transcript (Fig. 2A,B). However, small RNA sequences were predominantly 22–23 nt in length (mostly from the 5′ region of the reporter) (Supplemental Fig. S1) and were depleted of 5′U reads (13.9% of all reporter-mapped reads had a 5′U; standard deviation = 1.3; note that the reporter is depleted of uracils, with an overall U content of 21.3%) (Fig. 2C–F). Small RNAs produced upon Zuc tethering were sensitive to NaIO4 treatment (Fig. 2A,F), further confirming that the majority of reporter-derived reads upon tethering Zuc is not bona fide piRNAs. Thus, Zuc tethering induces processing of the transcript but not the efficient production of piRNAs. A closer look at the size profile revealed a smaller peak (27.4% of all reporter-derived reads; standard deviation = 0.01) around the 25- to 28-nt piRNA size range (Fig. 2C). This small RNA population had slightly more 5′U RNAs (28.4% had 5′U; standard deviation = 0.03) and was derived from the entire length of the reporter (Fig. 2E; Supplemental Fig. S1).

To test whether any genuine piRNAs are formed upon recruitment of Zuc, we immunoprecipitated Piwi/piRNA complexes and cloned the isolated piRNAs. Analysis of Piwi-associated RNAs yielded reporter-derived small RNAs with a size profile and 5′U bias characteristic of piRNAs (41.8% had 5′U) (Fig. 2A,G,H). Thus, tethering of Zuc leads to reporter processing into small RNAs; however, only a small fraction of the formed small RNAs is genuine piRNAs.

**Recruitment of Piwi and cytoplasmic piRNA pathway components triggers piRNA biogenesis in germ cells**

Recent studies suggested that in germ cells, both slicer-dependent processing and Zuc-mediated processing are triggered by initial Aub- or Ago3-mediated cleavage guided by complementary piRNAs (Han et al. 2015; Mohn et al. 2015). Thus, tethering of Zuc leads to reporter processing into small RNAs; however, only a small fraction of the formed small RNAs is genuine piRNAs.

Next we tested whether recruitment of Zuc—the endonuclease responsible for generating the 5′ end and, to some extent, the 3′ end of piRNAs (Malone et al. 2009; Ipsaro et al. 2012; Nishimasu et al. 2012; Han et al. 2015; Mohn et al. 2015)—is sufficient to trigger piRNA biogenesis. Small RNA sequencing (RNA-seq) upon Zuc tethering revealed the presence of abundant small RNAs arising from the reporter transcript (Fig. 2A,B). However, small RNA sequences were predominantly 22–23 nt in length (mostly from the 5′ region of the reporter) (Supplemental Fig. S1) and were depleted of 5′U reads (13.9% of all reporter-mapped reads had a 5′U; standard deviation = 1.3; note that the reporter is depleted of uracils, with an overall U content of 21.3%) (Fig. 2C–F). Small RNAs produced upon Zuc tethering were sensitive to NaIO4 treatment (Fig. 2A,F), further confirming that the majority of reporter-derived reads upon tethering Zuc is not bona fide piRNAs. Thus, Zuc tethering induces processing of the transcript but not the efficient production of piRNAs. A closer look at the size profile revealed a smaller peak (27.4% of all reporter-derived reads; standard deviation = 0.01) around the 25- to 28-nt piRNA size range (Fig. 2C). This small RNA population had slightly more 5′U RNAs (28.4% had 5′U; standard deviation = 0.03) and was derived from the entire length of the reporter (Fig. 2E; Supplemental Fig. S1).

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neither the genomic location of the target nor its sequence was crucial for processing.

We tested whether other components of the piRNA pathway can trigger processing in germ cells upon their recruitment to the reporter. Recruitment of nuclear proteins that work together with or downstream from Piwi to repress Piwi targets such as Asterix (Arx), Panoramix (Panx), and the histone methyltransferase SetDB1/Egg (Donertas et al. 2013; Muerdter et al. 2013; Sienski et al. 2015; Yu et al. 2015) did not trigger piRNA biogenesis (Fig. 3A,B). Recruitment of the cytoplasmic components Aub and Armi, two nuage components involved in pong-pong and Zuc-dependent processing, respectively, resulted in small RNA biogenesis (Fig. 3A,B). Small RNAs generated upon Piwi, Aub, or Armi tethering were resistant to NaIO4 treatment, confirming that they are generated by the canonical piRNA biogenesis machinery (Supplemental Fig. S4A).

The bulk of the reporter-mapped small RNAs upon Piwi, Aub, and Armi tethering was 25–28 nt in length [Fig. 3C], and the majority (60.4% [standard deviation = 1.8], 55.9%, and 78.4% [standard deviation = 0.06], respectively) had a strong bias for U at the first position [Fig. 3D; Supplemental Fig. S4B], both features of piRNAs. Cloning of small RNAs isolated from immunoprecipitated Piwi complex upon Piwi or Aub tethering confirmed that they are indeed bona fide piRNAs (Fig. 3E–G). Reporter piRNAs are derived only from the sense strand of the reporter [99.4% [standard deviation = 0.001], 99.4% and 99.8% [standard deviation = 0.001], respectively] and lack a bias for A at position 10 (a feature of ping-pong piRNAs) [Fig. 3D,G]. Together with the lack of complementary piRNAs, this suggests that these piRNAs are formed through a ping-pong-independent mechanism. We note that neither of our tethering experiments resulted in phased small RNAs from the reporter, although we did detect phasing in these libraries when we analyzed cluster 42AB (Supplemental Fig. S4C). We conclude that tethering of several different factors, such as Piwi, Aub, and Armi, to the reporter in germ cells
triggers piRNA biogenesis. The mechanism by which reporter piRNAs are generated does not require complementary piRNAs and ping-pong processing and thus seems similar, if not identical, to the mechanism observed in follicular cells.

Identification of factors required for piRNA processing in germ cells

Having shown that tethering of Piwi, Aub, or Armi is sufficient for piRNA biogenesis, we set out to identify factors that are required for piRNA processing of the reporter mRNA. We combined tethering of Piwi to the reporter with germline knockdown [Supplemental Fig. S5] of the respective gene followed by small RNA cloning.

Factors that work with Piwi to induce transcriptional repression [Panx and Egg] [Donertas et al. 2013; Muerdter et al. 2013; Sienski et al. 2015; Yu et al. 2015] are not required for piRNA biogenesis upon Piwi tethering (Fig. 4A,B). Similarly, Cuff and Thoc5, components of the RDC and TREX complexes, respectively, were dispensable for generation of reporter piRNAs [Fig. 4A,B]. These results indicate that nuclear Piwi partners as well as RDC and TREX are not required for piRNA biogenesis triggered by Piwi recruitment.

In contrast, reporter piRNAs were strongly [sevenfold] reduced upon knockdown of Zuc [Fig. 4A,B]. Similarly, strong [14-fold] reduction of piRNA biogenesis was observed upon knockdown of Armi [a putative RNA helicase]. These data confirm that reporter-derived small RNAs are genuine piRNAs formed through the canonical Zuc-dependent biogenesis pathway. A somewhat weaker effect on piRNA biogenesis was seen upon knockdown of other cytoplasmic piRNA processing proteins, such as the nuage component Vasa, an RNA helicase, and the Tudor domain protein Krimper, which reduced reporter piRNA levels approximately twofold [Fig. 4A,B]. The cytoplasmic piwi proteins Aub and Ago3 were dispensable for piRNA generation from the reporter upon Piwi tethering. This result further confirms that reporter piRNAs are generated independently of the ping-pong cycle [Fig. 4A, B]. Knockdown of Cuff and Ago3 resulted in an apparent increase in reporter-derived small RNAs when depicted as RPM of all mapping reads [Fig. 4A,B], likely due to the massive loss of endogenous piRNAs mapping to the DM3 genome. Overall, our results indicate that reporter RNA is processed by the canonical piRNA processing machinery, which includes Zuc and Armi, and that tethering bypasses the requirement of nuclear proteins.

Tethering to Piwi leads to transcript accumulation in nuage

We used RNA-FISH to test whether piRNA biogenesis triggered by tethering of Piwi leads to a change in the subcellular localization of the reporter RNA. Reporter RNA was uniformly distributed in the cytoplasm in control cells upon GFP tethering. Accumulation of the transcript in distinct perinuclear foci, which also show vasa enrichment, was observed upon Piwi tethering, indicating that Piwi recruits the bound RNA to nuage (Fig. 5A). Thus, induction of piRNA biogenesis from a heterologous reporter correlates with sequestration of the reporter mRNA to nuage.

Discussion

Compared with the processing of other classes of small RNAs, piRNA biogenesis is very complex and requires numerous proteins with, to date, unspecified roles in piRNA processing. How the precursor RNAs are discriminated from other cellular RNA and directed to the processing machinery is also not clear. As standard genetic and biochemical approaches failed to provide clear answers to...
these questions, we used an alternative approach, which is based on tethering of several piRNA pathway components to a heterologous reporter, to gain a better understanding of piRNA biogenesis. We found that tethering of several proteins (Piwi, Aub, and Armi) to an RNA that lacks homology with other piRNA precursors triggers its processing into piRNAs (Fig. 3A,B). The piRNAs produced by this approach have all of the features of canonical piRNAs: They have the correct size and a bias for U at the first nucleotide position and are loaded into Piwi (Fig. 3C–G). Biogenesis of these piRNAs is mediated by the canonical processing machinery (Fig. 4).

**Experimental approach to dissect the piRNA biogenesis pathway**

Most of the numerous proteins required for piRNA biogenesis were identified through genetic screens [Handler et al. 2011, 2013; Muerdter et al. 2013]. Mutations in these genes diminish or eliminate mature piRNAs; however, it was difficult to assign specific molecular functions to these proteins, largely due to the fact that we do not know intermediate steps of piRNA processing that could be analyzed in these mutants. For example, the nuclear RDC and TREX complexes are required for piRNA biogenesis and were proposed to direct pre-piRNAs to the processing machinery [Klattenhoff et al. 2009; Pane et al. 2011; Le Thomas et al. 2014; Mohn et al. 2014; Zhang et al. 2014; Chen et al. 2016; Hur and Chung 2016; Hur et al. 2016], however, the molecular mechanism by which these complexes might bring precursors to the cytoplasmic machinery remains unresolved.

The approach used in this study, tethering combined with knockdown of piRNA biogenesis factors, provides a valuable tool to dissect steps of the piRNA biogenesis pathway and discriminate proteins involved in processing and precursor localization per se from factors involved in upstream steps of biogenesis, such as transcription and nuclear export of pre-piRNAs. As expected, processing absolutely requires Zuc, the endonuclease that mediates the formation of both the 5′ and 3′ ends of piRNAs [Fig. 4; Malone et al. 2009; Ipsaro et al. 2012; Nishimasu et al. 2012; Han et al. 2015; Mohn et al. 2015]. While recruitment of Zuc to RNA leads to processing of the transcript, the majority of the obtained small RNAs is not piRNAs, indicating that Zuc by itself is not sufficient for efficient piRNA processing (Fig. 2). We show that another factor that is required for piRNA biogenesis upon tethering is the putative RNA helicase Armi (Fig. 4). Thus, efficient piRNA processing requires the interaction between Zuc and other proteins such as Armi.

We found that components of the nuclear RDC and TREX complexes, which associate with chromatin of piRNA clusters and nascent pre-piRNAs, respectively, are not required for piRNA processing upon tethering (Fig. 4), suggesting that their requirement for piRNA biogenesis can be bypassed. Previous studies have shown that both RDC and TREX are required for transcription of pre-piRNAs [Klattenhoff et al. 2009; Pane et al. 2011; Le Thomas et al. 2014; Mohn et al. 2014; Zhang et al. 2014; Chen et al. 2016; Hur and Chung 2016; Hur et al. 2016]. TREX is likely also involved in nuclear export of pre-piRNAs. It is possible that RDC and TREX are only necessary in the nucleus for transcription and export of pre-piRNAs and do not interact with the cytoplasmic processing machinery. Alternatively, these complexes might be required to channel natural precursors to the processing machinery, but this function can be effectively bypassed by recruitment of several different proteins to the RNA. Either way, RDC and TREX are not part of the core processing machinery. The same approach can be used to assign roles to other [including yet unidentified] piRNA biogenesis factors.

**Zuc-dependent, ping-pong-independent piRNA processing exists in germ cells**

Tethering of several proteins to an RNA that has no complementarity to mature piRNAs triggers its processing to piRNAs. This processing is observed in both follicular and germ cells [Figs. 1,3] and is independent of pre-existing piRNAs [Supplemental Fig. S2] and ping-pong. Historically, two types of piRNA biogenesis were proposed: the ping-pong cycle, which relies on the catalytic (slicer)
activity of Aub and Ago3 to generate the 5’ end of new piRNAs [Brennecke et al. 2007; Gunawardane et al. 2007], and the so-called primary processing, which was thought to be independent of ping-pong [Malone et al. 2009]. Recent studies suggested that in germ cells, “primary” biogenesis is triggered by initial ping-pong processing [Han et al. 2015; Mohn et al. 2015; Senti et al. 2015; Wang et al. 2015]. These results suggested that “primary” piRNA biogenesis is actually downstream from ping-pong (or “secondary” biogenesis), making these titles obsolete. Our data show that piRNA biogenesis that is independent of the ping-pong machinery can be achieved—at least by the artificial tethering approach—in germ cells. Therefore, both germ cells and somatic follicular cells (which lack the ping-pong machinery) [Malone et al. 2009] possess the same machinery, which allows them to process piRNAs in a ping-pong-independent fashion. We propose that Aub and Ago3 might promote Zuc-dependent processing of piRNA precursors through sequestration of these RNA to a specific cellular compartment rather than through its slicing activity [see below].

The sequestration to nuage might explain the selection of substrates for piRNA processing

How piRNA precursors are recognized by the piRNA processing machinery is still unresolved. The core piRNA processing machinery, composed of Zuc and Armi, is similar in both cell types, suggesting that a common principle for precursor selection should exist. piRNAs could be generated from artificial sequences inserted into piRNA clusters [Muerdter et al. 2012]. piRNA processing from genomic and tRNA transcripts was also observed [correlating with transcript abundance] in Kc167 cells. Both of these observations argue against the need for specific sequence/structure motifs in precursor transcripts [Vrettos et al. 2017]. We propose that sequestration of RNA into a distinct cellular compartment might be the central principle for precursor selection that is shared by germline and follicular cells. Based on our proposed model, any RNA that is localized to the processing compartment will be processed to piRNAs in a sequence-independent fashion [Fig. 5B].

Several lines of evidence support this hypothesis. First, efficient piRNA biogenesis can be triggered by recruitment of several different proteins to RNA, which have little in common except for their localization to nuage. Some of these factors, such as Aub, localize to nuage in steady state, while others localize transiently, such as Piwi on its way to the nucleus [Lim and Kai 2007; Le Thomas et al. 2013; Huang et al. 2014; Webster et al. 2015]. The only compartment where both Aub and Piwi are present together is nuage [Malone et al. 2009]. Recruitment of Zuc—the central player in processing, according to both previous results and our data—does not lead to efficient processing into piRNAs [Fig. 2]. Zuc associates with mitochondria and therefore is localized throughout the cytoplasm and not exclusively to nuage [Handler et al. 2013]. Second, we found that even though Zuc-dependent piRNA biogenesis does not require ping-pong, biogenesis is reduced upon knockdown of Vasa and Krimp, proteins that localize to nuage and were implicated previously in ping-pong [Fig. 4; Xiol et al. 2014; Webster et al. 2015]. Knockdown of Vasa disrupts the overall structure of nuage [Liang et al. 1994]; this might be true for knockdown of Krimp as well, making biogenesis that relies on physical integrity of this compartment less efficient. Finally, we observed that reporter RNA normally distributed throughout the cytoplasm is localized to nuage upon tethering [Fig. 5A]. A study published during the revision of this report showed that artificial reporter tethering to Armi in follicular cells, where Armi localizes to Yb bodies, leads to reporter processing into piRNAs. Mutating either the helicase or the ATP-binding domain of Armi delocalizes it from Yb bodies, and, consistent with our results, reporter tethering to these mutant (and thus delocalized) versions of Armi greatly reduced reporter processing into piRNAs [Pandey et al. 2017]. Thus, it seems that processing directly correlates with localization of the substrate to nuage.

How are pre-piRNAs recruited to nuage in germ cells and to Yb bodies in follicular cells? In our experiments, artificial tethering of the heterologous RNA to Piwi was sufficient to sequester it to nuage. Natural piRNA substrates might be recruited to Yb bodies through RNA-binding proteins that recognize specific motifs in RNA sequences, such as the sequences identified in tij or Flamenco [Hommola et al. 2015; Ishizu et al. 2015]. In germ cells, pre-piRNAs might be recruited to nuage through interaction with Aub and Ago3 loaded with complementary piRNAs. Piwi-loaded piRNAs are substantially reduced in the Aub/Ago3 double mutant, which was interpreted as a requirement for slicer cleavage to trigger Zuc-dependent processing [Han et al. 2015; Mohn et al. 2015; Senti et al. 2015; Wang et al. 2015]. We propose that these results can be explained by failure to sequester pre-piRNAs to nuage in the Aub/Ago3 double mutant. Expression of catalytically impaired Ago3 and Aub, which cannot cleave precursors but are likely able to bind and recruit them to nuage, can partially rescue Zuc-dependent processing of Piwi-associated piRNAs [Wang et al. 2015].

Compartmentalization is important for many RNA processing pathways, such as splicing, rRNA maturation, etc. [Zhang et al. 1994; Henras et al. 2015]. However, our hypothesized model of piRNA precursor selection goes beyond stating that compartmentalization is important for piRNA processing: While RNA sequence motifs are still required for processing in all other pathways [Zeng et al. 2005; Park et al. 2011; Barrangou and Marraffini 2014; Li and Patel 2016; Tsai and Joung 2016], indicating that localization of substrates to the processing compartments is necessary but not sufficient to trigger processing, we propose that for the piRNA pathway, localization of RNA into nuage/Yb granules is both necessary and sufficient to initiate piRNA processing.

Our model of piRNA biogenesis allows unification of two seemingly different mechanisms for selection of piRNA precursors operating in germ and follicular cells. Processing induced by artificial recruitment enables separation of proteins involved in upstream steps of piRNA biogenesis, while Zuc-based biogenesis is triggered by initial processing of piRNA precursors, leading to increased biogenesis. This unification is consistent with previous results and our data.
biogenesis from factors involved in Zuc-mediated processing. Finally, selection of piRNA precursors through recruitment to the processing machinery enables the design of artificial sequences that can be processed efficiently into piRNAs and thus opens possibilities of using the piRNA pathway as a tool for both transcriptional and post-transcriptional gene regulation.

**Materials and methods**

**Fly stocks**

The sh-White, sh-AGO3, sh-Argi, sh-Aub, sh-Cuff, sh-Krimper, sh-Vasa, and sh-Zuc lines were obtained from the Bloomington *Drosophila* Stock Center [BDSC, nos. 33523, 33522, 34789, 33728, 35182, 35230, 32434, and 35227]. The GFP-Piwi line was a generous gift from the Hannon laboratory. The sh-SetDB1, sh-Phalan, sh-Flag-Piwi, AN-Flag, and Tubulin-GFP-BoxB lines were generous gifts from the Brennecke laboratory. The TJ-Gal4 line was obtained from the Kyoto Stock Center (DGGR, no. 104055). To obtain the sh-Thoc5 line, the short hairpin sequences were ligated into the pValium20 vector [Ni et al. 2011] using T4 DNA ligase from New England Biolabs (M0202), according to the manual, and then integrated into the attP2 landing site (BDSC, 25710). Hairpin sequences are listed in Supplemental Table S2. UASp-AN-eGFP-Piwi, UASp-AN-eGFP-Aub, UASp-ArmiHA-AN, UASp-Zuc-HA-AN, UASp-AN-eGFP-GFP, UASp-AN-HA-Yb, UASp-AN-eGFP-Thoc5, UASp-AN-eGFP-Cuff, UASp-AN-eGFP-Asterix, UASp-AN-eGFP-Panoramix, UASp-AN-eGFP-Egg, and UASp-mKate2-4xBoxB-K10polyA were generated in the laboratory as described previously [Chen et al. 2016]. These constructs were generated by P-element integration and driven by maternal a-tubulin67C-Gal4 [BDS, 7063].

Flies carrying the AN transgenes and the α-tubulin67C-Gal4 driver were crossed to flies carrying the UASp-mKate2-4xBoxB-K10polyA reporter and, where applicable, the respective hairpin to obtain flies that express all components of the system in the female germline. For expression in follicular cells, flies carrying the AN transgenes and the TJ-gal4 driver were crossed to flies carrying the UASp-mKate2-4xBoxB-K10polyA reporter. Flies were put on yeast for 3–4 d prior to dissection and were 5–8 d old at the time of dissection.

**RT-qPCR**

Total RNA was isolated from 20 ovaries with Ribozol [Amresco, N580] followed by DNase treatment with amplification-grade DNase I [Invitrogen, 18068-015] according to the manual. Reverse transcription was carried out using SuperScript III [Invitrogen] with oligo dT15. qPCR was performed on a Mastercycler ep Realplex PCR thermal cycler machine (Eppendorf). RT-qPCR data target expression was normalized to the rp49 locus expression. PCR primers are listed in Supplemental Table S2.

**Small RNA-seq**

Ovaries were dissected, and total RNA was isolated with Ribozol [Amresco, N580]. Small RNAs within a 19- to 29-nt window were isolated from 15% polyacrylamide gels from 4 µg of ovarian total RNA. For samples that were NaO4-treated, 5x borate buffer (pH 8.6, 150 mM borax, 150 mM boric acid) and 200 mM sodium periodate were added to the size-selected small RNA, and the samples were incubated for 30 min at 25°C. The NaO4-treated small RNA was then ethanol-precipitated before proceeding to library construction. The small RNA libraries were constructed using the NEBNext small RNA library preparation set for Illumina [no. E7330S], according to the protocol, using NEBNext multiplex oligos for Illumina [no. E7335S]. Libraries were sequenced on the Illumina HiSeq 2500 [SE 50-bp reads] platform. The resulting reads were mapped against the reporter sequence and the DM3 genome using Bowtie 0.12.7 [Langmead et al. 2009] with the settings “-v 0 -m 10000 –best-strata,” retaining reads that map with zero mismatches. The reads were also mapped to RepBase using Bowtie 0.12.7 [Langmead et al. 2009] with the settings “-v 3 –best-strata.” Reads that specifically mapped to the reporter sequence and not to the DM3 genome with zero mismatches were extracted from the libraries. Where applicable, 22- to 23-nt reads or 25- to 28-nt reads were extracted from the reads mapping exclusively to the reporter. All four sets of reads (all reads, all reads-specific, 22- to 23-nt reads-specific, and 25- to 28-nt reads-specific) were plotted along the reporter sequence normalized to the number of total reads or 25- to 28-nt reads mapping to the DM3 genome, respectively, using Integrative Genomics Viewer 2.3.68 [Robinson et al. 2011]. Reads that mapped specifically to the 42AB locus [chromosome 2R: 2,144,349–2,386,719] and not to the rest of the DM3 genome were extracted from the libraries.

Weblogos were generated using an R script that uses the seqLogo library [https://rdr.io/bioc/seqLogo] to display the nucleotide frequency occurring at each position of the input reads calculated within a homemade bash script. Sequencing data are summarized in Supplemental Table S1.

**Immunoprecipitation small RNA-seq**

Ovaries (~100 per immunoprecipitation) from flies expressing UASp-AN-eGFP-Piwi, UASp-AN-eGFP-Aub, or UASp-Zuc-HA-AN with UASp-GFP-Piwi, and the UASp-mKate2-4xBoxB-K10polyA reporter line driven by maternal α-tubulin67C-Gal4 were dissected and lysed on ice in 250 µL of lysis buffer [30 mM HEPES-KOH at pH 7.4, 100 mM KOAc, 2 mM Mg(OAc)2, 5 mM DTT, 0.5% (v/v) NP40, proteinase inhibitor [Roche, 11836170011], RNasin Plus [Promega, N2611]. Lysate was dounced and clarified by centrifugation at maximum speed at 4°C. The supernatant was incubated with protein A/G agarose beads [Thermo Fisher, 20421] preconjugated with rabbit polyclonal anti-GFP [Covance, affinity-purified in our laboratory] or anti-Piwi [Brennecke et al. 2007] for 2 h at 4°C. The immunoprecipitation and RNA isolation were carried out as described previously [Vagin et al. 2006]. A fifth of the RNA was CIP-treated [New England Biolabs, M0290S] in NEB buffer #3 [New England Biolabs, B700S] for 30 min at 37°C and then ethanol-precipitated after phenol:chloroform and chloroform extraction. The CIP-treated RNA was then PNK-treated with 1 µL of 10x T4 polynucleotide kinase buffer [New England Biolabs, B0210S], 2 µL of γ-<sup>P</sup>32ATP [PerkinElmer, BLU502A250UC], and 1 µL of T4 polynucleotide kinase [New England Biolabs, M0201S] for 45 min at 37°C. The CIP- and PNK-treated RNA was added back to the remainder of the RNA isolated from the immunoprecipitation. Size selection, library preparation, and analysis were performed as described above, except that fragments were gel-extracted based on labeled immunoprecipitation material.

**RNA-FISH and immunofluorescence**

Probes specific to the mKate2 transcript were designed and ordered from Stellaris [Biosearch Technologies] and resuspended in 200 µL of DEPC-treated TE buffer (10 mM Tris-HCl, 1 mM EDTA at pH 8.0) and stored at −20°C as stock. Fish were performed on 30–50 pairs of ovaries that were dissected and washed

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with ice-cold DEPC-treated PBS. Prior to fixation, the ovaries were exposed to blue light for 15 min to quench GFP fluorescence from the tethering construct [UA5pAN-eGFP-Piwi and UA5pAN-eGFP-GFP]. The ovaries were fixed with 300 µL of fixation solution (4% paraformaldehyde, 0.15% Triton X-100 in DEPC-treated PBS) with shaking at 300 rpm at room temperature and washed three times with PBX (DEPC-treated PBS, 0.3% Triton X-100) for 10 min each at room temperature. Samples were dehydrated in 500 µL of 70% ethanol (in DEPC-treated water) and permeabilized overnight at 4°C on a nutator. Stock probes were diluted 100 times in hybridization buffer (DEPC-treated 2× SSC, 10% [w/v] dextran sulfate, 10% [v/v] formamide). Following rehybridization of ovaries in 500 µL of wash buffer (DEPC-treated 2× SSC, 10% [v/v] formamide) for 5 min, the diluted probe was added to the samples for overnight incubation at 37°C in a hybridization chamber. Samples were washed twice with wash buffer and incubated in wash buffer twice for 30 min at 37°C. Samples were blocked with SBX buffer (DEPC-treated 2× SSC, 1% [w/v] BSA, 0.3% Triton X-100) for 2 h at room temperature and incubated with 1:100 diluted anti-Vasa [Developmental Studies Hybridoma Band, A.C. Spradling and D. Williams] overnight at 4°C followed by five washes with SBX buffer for 10 min at room temperature and incubation with 1:400 diluted chicken anti-rat IgG secondary antibody and Alexa fluor 647 (Thermo Fisher, A21472) in SBX buffer for 4 h at room temperature in the dark without agitation. Ovaries were washed five times for 10 min with SSCT buffer (DEPC-treated 2× SSC, 0.1% Triton X-100) at room temperature and incubated with 1:5000 diluted DAPI in 2× SSC buffer for 10 min at room temperature on a nutator. Ovaries were washed twice with 2× SSC buffer and then mounted on a glass slide using Prolong Gold anti-fade reagent (Thermo Fisher, P36934).

Accession numbers

High-throughput sequencing data for small RNA-seq experiments are available through Gene Expression Omnibus (GSE102961).

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