Discrete Small RNA-Generating Loci as Master Regulators of Transposon Activity in Drosophila

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SUMMARY

Drosophila Piwi-family proteins have been implicated in transposon control. Here, we examine piwi-interacting RNAs (piRNAs) associated with each Drosophila Piwi protein and find that Piwi and Aubergine bind RNAs that are predominantly antisense to transposons, whereas Ago3 complexes contain predominantly sense piRNAs. As in mammals, the majority of Drosophila piRNAs are derived from discrete genomic loci. These loci comprise mainly defective transposon sequences, and some have previously been identified as master regulators of transposon activity. Our data suggest that heterochromatic piRNA loci interact with potentially active, euchromatic transposons to form an adaptive system for transposon control. Complementary relationships between sense and antisense piRNA populations suggest an amplification loop wherein each piRNAdirected cleavage event generates the 5' end of a new piRNA. Thus, sense piRNAs, formed following cleavage of transposon mRNAs may enhance production of antisense piRNAs, complementary to active elements, by directing cleavage of transcripts from master control loci.

INTRODUCTION

Mobile genetic elements, or their remnants, populate the genomes of nearly every living organism. Potential negative effects of mobile elements on the fitness of their hosts necessitate the development of strategies for transposon control. This is critical in the germline, where transposon activity can create a substantial mutational burden that would accumulate with each passing generation.

Hybrid dysgenesis exemplifies the deleterious effects of colonization of a host by an uncontrolled mobile element. The progeny of intercrosses between certain Drosophila strains reproducibly show high germline mutation rates with elevated frequencies of chromosomal abnormalities and partial or complete sterility (Bucheton, 1990; Castro and Carareto, 2004; Kidwell et al., 1977). Studies of the molecular basis of this phenomenon linked the phenotype to transposon mobilization (Pelisson, 1981; Rubin et al., 1982).

Hybrid dysgenesis occurs when a transposon, carried by a male that has established control over that element, is introduced into a naïve female that does not carry the element. The transposon becomes active in the progeny of the naïve female, causing a variety of abnormalities in reproductive tissues that ultimately result in sterility (Engels and Preston, 1979). Since the dysgenic phenotype is often not completely penetrant, a fraction of the progeny from affected females may survive to adulthood. Such animals can develop resistance to the mobilized element, although in many cases, several generations are required for resistance to become fully established (Pelisson and Bregliano, 1987). Immunity to transposons can only be passed through the female germline, indicating that there are both cytoplasmic and genetic components to inherited resistance (Bregliano et al., 1980).

Studies of hybrid dysgenesis have served a critical role in revealing mechanisms of transposon control. In general, two seemingly contradictory models have emerged. The first model correlates resistance with an increasing copy number of the mobile element. A second model suggests that discrete genomic loci encode transposon resistance.

The first model is supported by studies of the I element. Crossing a male carrying full-length copies of the I element to a naïve female leads to I mobilization and hybrid dysgenesis (Bregliano et al., 1980; Bucheton et al., 1984). The number of I copies builds during subsequent crosses of surviving female progeny until it reaches an average of 10-15 per genome (Pelisson and Bregliano, 1987). At this point, I mobility is suppressed, as the initially naïve strain

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gains control over this element. Thus, a gradual increase in I element copy number over multiple generations was implicated in the development of transposon resistance.

The second model, which attributes transposon resistance to specific genetic loci, is illustrated by studies of gypsy transposon control (Bucheton, 1995). Genetic mapping of gypsy resistance determinants led to a discrete locus in the pericentric β-heterochromatin of the X chromosome that was named flamenco (Pelisson et al., 1994). Females carrying a permissive flamenco allele (one that allows gypsy activity) showed a dysgenic phenotype when crossed to males carrying functional gypsy elements. Permissive flamenco alleles exist in natural Drosophila populations but can also be produced by insertional mutagenesis of animals carrying a restrictive flamenco allele (Robert et al., 2001). Despite extensive deletion mapping over the flamenco locus, no transposon repressor from flamenco has been identified. For P elements, a repressor of transposition has been identified as a 66 kDa version of the P element transposase. Expression of the repressor was proposed to correlate with increasing P element copy number, leading to a self-imposed limitation on P element mobility (Misra and Rio, 1990). However, studies of resistance determinants indicated that control over P elements could also be established by insertion of P elements into specific genomic loci, arguing for an alternative, copy number-independent control pathway (Biemont et al., 1990). Studies of inbred lines or of wild isolates with natural P element resistance indicated that P insertions near the telomere of X (cytological position 1A) were sufficient to confer resistance if maternally inherited (Biemont et al., 1990; Ronsseray et al., 1991). Additionally, several groups isolated insertions of incomplete P elements in this same cytological location that acted as dominant transposition suppressors (Marin et al., 2000; Stuart et al., 2002). Importantly, these defective P elements lacked sequences encoding the repressor fragment of transposase.

Both models of transposon resistance, those determined by specific genomic loci and those caused by copy number-dependent responses might be linked to small RNA-based regulatory pathways. Copy numberdependent silencing of mobile elements is reminiscent of copy number-dependent transgene silencing in plants (cosuppression) (Smyth, 1997) and Drosophila (Pal-Bhadra et al., 1997). In both cases, silencing occurs through an RNAi-like response where high-copy transgenes provoke the generation of small RNAs, presumably through a double-stranded RNA intermediate (Hamilton and Baulcombe, 1999; Pal-Bhadra et al., 2002). Moreover, mutations in RNAi pathway genes impact transposon mobility in flies (Kalmykova et al., 2005; Sarot et al., 2004; Savitsky et al., 2006) and C.elegans (Ketting et al., 1999; Tabara et al., 1999). Finally, small RNAs (rasiRNAs) corresponding to transposons and repeats have been isolated from flies and zebrafish (Aravin et al., 2001, 2003; Chen et al., 2005).

At the core of the RNAi machinery are the Argonaute proteins, which directly bind to small RNAs and use these as guides for the identification and cleavage of their targets (Liu et al., 2004). In animals, Argonautes can be divided into two clades (Carmell et al., 2002). One contains the Argonautes, which act with microRNAs and siRNAs to mediate gene silencing. The second contains the Piwi proteins. Genetic studies have implicated Piwi proteins in germline integrity (Cox et al., 1998; Harris and Macdonald, 2001). For example, *piwi* mutations cause sterility and loss of germline stem cells (Cox et al., 1998; Lin and Spradling, 1997). aubergine is a spindle-class gene that is required in the germline for the production of functional oocytes (Harris and Macdonald, 2001). The third Drosophila Piwi gene, Ago3, has yet to be studied. Mutation of Piwi-family genes also affects mobile elements. For example, piwi mutations mobilize gypsy (Sarot et al., 2004), and aubergine mutations impact TART (Savitsky et al., 2006) and P elements (Reiss et al., 2004). Finally, both Piwi and Aubergine bind rasiRNAs (Saito et al., 2006; Vagin et al., 2006) targeting a number of mobile and repetitive elements. These complexes are enriched for antisense small RNAs, as might be expected if they were actively involved in silencing transposons by recognition of their RNA products.

Recently, a new class of small RNAs, the piRNAs, was identified through association with Piwi proteins in mammalian testes (Aravin et al., 2006; Girard et al., 2006; Grivna et al., 2006; Lau et al., 2006). These 26-30 nt RNAs are produced from discrete loci, generally spanning 50-100 kb. Interestingly, mammalian piRNAs are relatively depleted of transposon sequences. Despite apparent differences in the content of Piwi-associated RNA populations in mammals and Drosophila, Piwi-family proteins share essential roles in gametogenesis, with all three murine family members, Miwi2 (M.A. Carmell et al., submitted), Mili (Kuramochi-Miyagawa et al., 2004), and Miwi (Deng and Lin, 2002), being required for male fertility.

In order to probe mechanisms of transposon control in Drosophila and to understand the relationship between Piwi protein function in flies and mammals, we undertook a detailed analysis of small RNAs associated with Piwi proteins in the Drosophila female germline. Our studies indicate that Drosophila Piwi-family members function in a transposon surveillance pathway that not only preserves a genetic memory of transposon exposure but also has the potential to adapt its response upon contact with active transposons.

RESULTS

Piwi-Family Members Have Distinct Expression Patterns

In Drosophila, Piwi-clade consists of three members: Piwi, Aubergine (Aub), and Ago3. As a prerequisite to further studies of this family, we experimentally determined the sequence of Ago3 and raised antisera specific to each Drosophila Piwi-family protein (Figures S1 and S2).

Previous studies have used myc-tagged Piwi and green fluorescent protein (GFP)-tagged Aub transgenes to investigate their spatial and temporal expression patterns

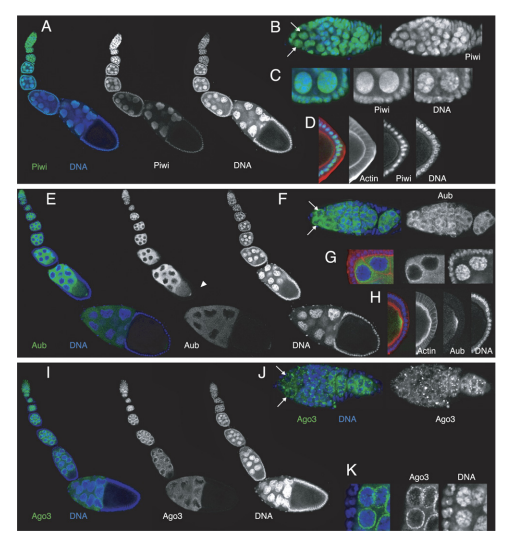


Figure 1. Expression and Localization of Piwi-Family Members in Ovarioles

In all panels, signal from the indicated Piwi-family member is in green; DNA is in blue and actin in red.

- (A) Overview of Piwi localization in the ovariole.
- (B) Detailed view of the germarium containing the two stem cells (arrows).
- (C) Nuclear localization of Piwi in nurse cells and surrounding follicle cells.
- (D) Indicates a weak accumulation of maternally deposited Piwi at the posterior pole of stage 10 oocytes.
- (E) Overview of Aubergine localization in the ovariole with enrichment of Aub at the posterior oocyte pole in late egg chambers (arrow head).
- (F) Detailed view of Aub localization in the germarium containing the two stem cells (arrows).
- (G) Enrichment of Aub in the cytoplasm and perinuclear nuage of germline cells; staining is absent from surrounding somatic follicle cells.
- (H) Accumulation of Aub at the posterior pole of a stage 10 oocyte.
- (I) Overview of Ago3 localization in the ovariole.
- (J) Detailed view of Ago3 staining in the germarium showing strong enrichment around the stem cell nuclei (arrows) and in discrete foci.
- (K) Detailed view of Ago3 localization to nuage in nurse cells.

during oogenesis (Cox et al., 2000; Harris and Macdonald, 2001). We used our specific antibodies to examine the expression patterns of the endogenous proteins and to extend analyses to the third family member, Ago3.

As previously reported (Cox et al., 2000), Piwi is predominantly nuclear and is present not only in germline cells but also in the somatic cells of the ovary (Figures 1A-1D). Strong Piwi staining is seen in the cap cells that surround the germline stem cells and in the follicle cells that envelop the developing egg chamber. In later stage egg chambers, Piwi is detectable in the cytoplasm of the developing oocyte with a slight enrichment at the posterior where the germline primordia of the embryo will form.

Aubergine is expressed at very low or undetectable levels outside the germline (Figures 1E-1H). Aub is primarily cytoplasmic. As reported previously for GFP-Aub, we detect endogenous protein in germline stem cells, developing cystoblasts, and the nurse cells of developing egg



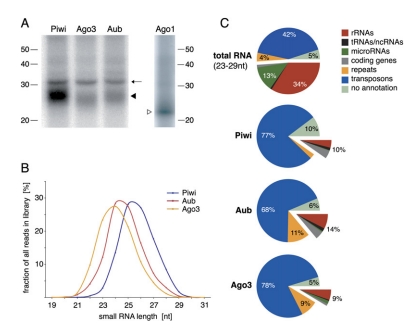


Figure 2. Characteristics of *Drosophila* piRNAs

- (A) Radioactively labeled RNA isolated under identical conditions from specific Piwi-family RNPs and Ago1 was analyzed on a denaturing polyacrylamide gel. The positions of RNA size markers, electrophoresed in parallel, are shown to the left. Indicated are piRNAs (solid arrowhead), miRNAs (open arrow head), and 2S rRNA (arrow), which is also present in purifications using control antibodies.
- (B) Size distributions of sequenced piRNAs specifically bound by the three Piwi-family members.
- (C) Pie chart summarizing the annotation of piRNA populations in total RNA and those bound by Piwi, Aub, and Ago3.

chambers. Aubergine is enriched in nuage, a perinuclear, electron dense structure, displaying a localization pattern very similar to Vasa. Like Vasa, Aubergine is deposited into the developing oocyte from early stage 10 onward and is localized to the pole plasm.

Ago3 protein is also predominantly cytoplasmic (Figure 1I–1K). It is present in the germline but not detectable in somatic cells of the egg chamber, although we do find Ago3 in the somatic cap cells of the germarium. Ago3 shows a more striking accumulation in nuage than does Aub, and it is also found in prominent but discrete foci of unknown character in the germarium. Despite its localization to nuage, Ago3 does not accumulate at the posterior pole of the developing oocyte, and Ago3 is not detected in the pole plasm of early embryos.

Piwi-Family Members Bind Distinct Populations of Small RNAs

To investigate the small RNA populations bound by each Drosophila Piwi protein, we purified RNP complexes from ovary lysates. All three proteins associate with small RNAs ranging in length from 23 to 29 nt (Figure 2A). We prepared cDNA libraries from each complex and from 23-29 nt RNAs purified from total ovary RNA. 454 sequencing yielded 60,691 reads (17,709 for Piwi; 23,376 for Ago3; 14,872 for Aub; and 4,734 for ovary total RNA) that match perfectly to Release 5 of the Drosophila genome or to nonassembled Drosophila sequences from GenBank. These were used for subsequent analysis. It should be noted that we isolated small RNAs from the Oregon R strain, rather than the sequenced strain $(y^1; cn^1 bw^1 sp^1)$. A detailed discussion of potential differences between these genomes and any impacts on the data that we present can be found in the Supplemental Data.

Based both upon gel mobility (Figure 2A) and size distributions (Figure 2B), each Piwi protein bound a specific class of piwi-interacting RNA (piRNA). Piwi-associated RNAs were the largest (25.7 nt), followed by Aub (24.7) and Ago3 bound (24.1) RNAs. As with mammalian piRNAs, Piwi and Aub bound RNAs have a strong preference for a 5' terminal uridine (83% and 72%, respectively), a trend that is essentially absent from the Ago3 bound population (37% terminal U). *Drosophila* piRNA populations are quite complex, with most RNAs being cloned only once (87% for Piwi, 81% for Aub, and 73% for Ago3).

Despite their size differences, the small RNAs obtained from each complex were remarkably similar in the types of genomic elements to which they correspond. Overall, roughly three quarters of all sequences from each complex could be assigned to annotated transposons or transposon remnants, with nearly all known transposons being represented (Figures 2C and S3). An additional 2%–11% of small RNAs were derived from regions of local repeat structure, such as the subtelomeric TAS repeats or pericentromeric satellite repeats. Thus, nearly 80% of *Drosophila* piRNAs can also be characterized as rasiRNAs. An additional group of sequences (10% for Piwi, 6% for Aub, and 5% for Ago3) map to unannotated regions of the genome, most often to heterochromatic, transposon-rich loci.

Drosophila piRNAs Are Derived from Discrete Genomic Loci

In *Drosophila*, intact and potentially active transposons populate the euchromatin as well as pericentromeric and telomeric heterochromatin. There are also numerous transposon remnants that have mutated sufficiently to negate their potential for transposition. These are strongly enriched in the β -heterochromatin that borders

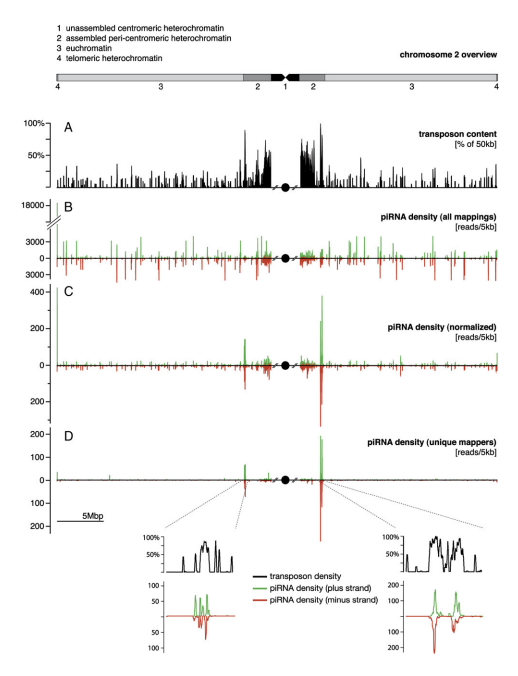


Figure 3. Drosophila piRNAs Map to Discrete Genomic Loci For reference, a sketch of chromosome 2 with the major chromatin domains is provided.

(A) Density of annotated transposons.

(B-D) Density of cloned piRNAs (green: sense; red: antisense) along chromosome 2; the y axis indicates relative units, and the centromere is shown as a circle. In (B) each genomic position corresponding to a cloned piRNA is given equal weight; in (C) each piRNA-genomic match is normalized for the number of times it maps to the genome, resulting in proportionally less signal for piRNAs that map many times. In (D) only those piRNAs that map uniquely to the genome are plotted. The most prominent piRNA clusters (enlarged below and shown together with transposon density in black) are typically found at the border of the pericentromeric heterochromatin.

centromeres and are generally absent from euchromatic chromosome arms (Hoskins et al., 2002).

A depiction of the chromosomal locations matched by Drosophila piRNAs closely resembles a plot of transposon density (Figure 3). However, most piRNAs match multiple chromosomal sites. Therefore, to address the genomic origin of piRNAs, it was necessary to restrict our analysis to the \sim 20% that match the genome at a unique position (Figure S4). A density plot of this small RNA subset shows a striking clustering of piRNAs at discrete loci (e.g.,

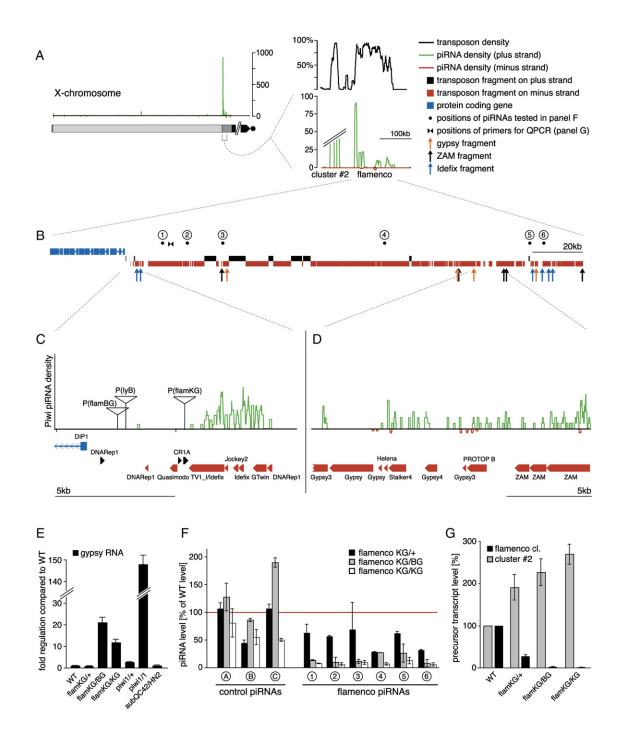


Figure 4. flamenco Is a piRNA Cluster

(A) piRNA density on the X chromosome showing two large clusters (numbers as in Table S1) in the pericentromeric heterochromatin (chromatin domains are shaded as in Figure 3). Densities of uniquely mapping sense (green) and antisense (red) piRNAs in two clusters is shown enlarged alongside with corresponding transposon density (black).

- (B) Shown is a more detailed map of the flamenco locus, indicating protein-coding genes (blue), sense (black), and antisense (red) transposon fragments. The flamenco cluster ends ~180 kb proximal to DIP1 in a gap of unknown size in the genome assembly. Arrows highlight the retroelements gypsy (orange), Idefix (blue), and ZAM (black), which are known to be regulated by the locus.
- (C) Shown are the first 10 kb of the flamenco locus with the flanking DIP1 gene (blue), annotated transposon fragments, the P element insertions that result in inactive flamenco alleles (triangles), and the density of Piwi-associated piRNAs that map to this region (note that over 99% of the uniquely mapping piRNAs in the flamenco cluster derive from the sense strand and that more than 95% associate with Piwi).
- (D) Fifteen kilobases of the flamenco locus showing Piwi bound piRNAs and a detailed view of transposon fragments from the region, including gyspy and ZAM.

Figure 3D). A similar plot can be obtained for all piRNAs if each piRNA-genomic match is divided by the number of genomic hits for that sequence (normalized) (Figure 3C). We identified 142 genomic locations as sites of abundant piRNA generation (Tables S1 and S2). These clusters produce 81% of all piRNAs that match the genome at a single site. Although these sites comprise only 3.5% of the assembled genome, more than 92% of all sequenced piRNAs could potentially be derived from these loci.

Only seven piRNA clusters occur in potentially euchromatic regions, with the remainder being present in pericentromeric and telomeric heterochromatin. Telomeric clusters most often consist of satellite sequences and correspond to the subtelomeric terminal associated sequence (TAS) repeats (Karpen and Spradling, 1992). These flank the telomeric arrays of HetA and TART transposons, for which we also find corresponding piRNAs. Telomeric clusters, supported by the presence of uniquely mapping piRNAs, are found on most chromosome arms (X, 2R, 2L, and 3R). Clusters in the pericentromeric β-heterochromatin display a high content of annotated transoposons (typically from 70% to 90%), with the majority being partial or defective nested elements.

The size of Drosophila piRNA clusters varies substantially with the largest being a 240 kB locus in the pericentromeric heterochomatin of chromosome 2R (cytological position 42AB). This cluster produces 20.8% of all uniquely mapping piRNA sequences and could potentially give rise to 30.1% of all the piRNAs, which we identified (Table S1). Overall, the largest 15 clusters account for 57% of the unique piRNAs and up to 70% of the total.

piRNA Clusters Are Master Regulators of Transposon Activity

Numerous genetic studies have pointed to discrete genomic loci that suppress the activity of specific transposons. The best understood is the recessive flamenco/COM locus (Prud'homme et al., 1995). flamenco was originally identified as a locus controlling the activity of the retroviral gypsy element (Pelisson et al., 1994). This locus has subsequently been shown to regulate two additional retroelements, Idefix and ZAM (Desset et al., 2003).

Through the use of numerous deficiencies, flamenco was mapped proximally to the DIP-1 gene and proposed to span a region of at least 130 kb. This locus corresponds precisely to a piRNA cluster (cluster 8; Table S1, Figures 4A and 4B). Eighty-seven percent of the sequence in the locus consists of nested transposable elements spanning a total length of 179 kb. The locus includes numerous fragments of all three transposable elements that were shown to be controlled by flamenco/COM (gypsy, Idefix, and ZAM; Figurse 4B-4D) in addition to many other transposon families.

A second piRNA cluster that has been genetically linked to transposon control corresponds to the subtelomeric TAS repeat on the X chromosome (X-TAS) (Table S1; cluster 4). Numerous studies indicate that insertions of one or two P elements into X-TAS are sufficient to suppress P-M hybrid dysgenesis (Marin et al., 2000; Ronsseray et al., 1991; Stuart et al., 2002). Transposon silencing by these insertions has been linked to the Piwi family, as it is relieved by mutations in aubergine (Reiss et al., 2004). The precise insertion sites of three suppressive P elements in X-TAS have been mapped, and they correspond to areas, which give rise to multiple small RNAs (not shown). In accord with a trans-acting model for silencing, lacZcontaining P elements inserted into X-TAS can suppress euchromatic lacZ transgenes in the female germline (Roche and Rio, 1998; Ronsseray et al., 1998).

A Functional Pathway Links flamenco-Derived piRNAs to gypsy Suppression

To probe the relevance of the piRNA cluster mapped to flamenco, we made use of mutations that negate the ability of this locus to silence *gypsy*. The only molecularly defined flamenco allele corresponds to a P element insertion ~2 kb proximal to DIP1 and 550 bp upstream of the first piRNA uniquely mapped to this cluster (P(lyB); Figure 4C) (Robert et al., 2001). We obtained two additional lines that harbor P element insertions near P(lyB) and tested their effects on gypsy expression. gypsy RNA levels increased by ~20-fold in strains carrying homozygous trans-heterozygous insertions, indicating that P(KG00476) (flamKG) and P(BG02658) (flamBG) (Figure 4C) are indeed *flamenco* mutant alleles (Figure 4E).

We next examined the levels of mature piRNAs from flamenco in wild-type animals and flamenco mutants. Using quantitative RT-PCR, we found substantial reductions in piRNAs that uniquely map to the flamenco piRNA cluster in mutant animals (2, 3, and 5; Figures 4B and 4F). In contrast, piRNAs definitively derived from other clusters were unaffected. We also probed levels of piRNAs that did not map uniquely to flamenco (1,4, and 6; Figures 4B and 4F). These were also substantially reduced in flamenco mutants, indicating that they arise mainly from this cluster despite not being uniquely assignable based upon sequence information alone.

The flamenco piRNA cluster preferentially loads the Piwi protein, with 94% of its uniquely mapping RNAs being Piwi partners. An examination of gypsy RNA levels reveals a 150-fold increase in piwi mutants (Figure 4E) (Sarot et al., 2004). In contrast, aubergine mutations show no

⁽E) Quantitative RT-PCR analysis on gypsy transcript levels in RNA isolated from wild-type, flamenco mutant, piwi mutant, and aub mutant ovaries. Shown are average levels (n = 3), and error bars indicate standard deviation (SD).

⁽F) Quantitative RT-PCR analysis on several individual piRNAs from small RNA libraries prepared from wild-type or flamenco mutant ovaries. Positions of tested piRNAs in *flamenco* are indicated in (B). Error bars indicate SD.

⁽G) Quantitative RT-PCR analysis on precursor transcripts from two different piRNA clusters in ovaries from wild-type and flamenco mutant flies. Position of primers used for the flamenco primary transcript are indicated in (B). Shown are average levels (n = 3), and error bars indicate SD.

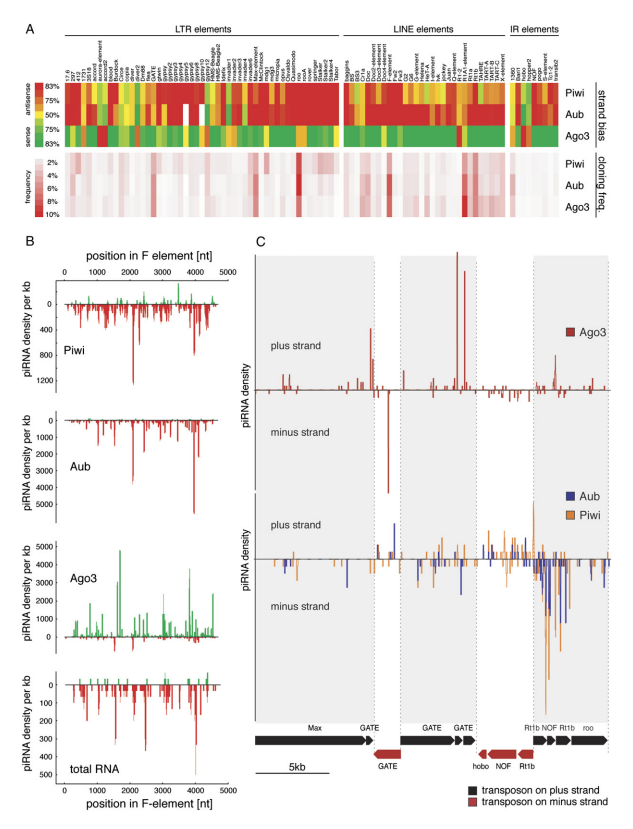


Figure 5. Piwi Complexes Show Profound Strand Biases

(A) Shown in the upper panel is a heat map indicating the strand bias of cloned piRNAs with respect to canonical transposon sequences (indicated at the top). Transposons are grouped into long terminal repeat (LTR) elements, long interspersed nuclear elements (LINE) elements, and inverted repeat elevation of gypsy levels, consistent with a minority of flamenco piRNAs entering this complex and with a lack of Aub expression in follicle cells where gypsy is expressed. The greater effect of piwi compared to flamenco mutations is consistent with flamenco locus contributing a substantial fraction of but not all gypsy repressive piRNAs. In this regard, the flamenco cluster has the potential to produce 79% of all piRNAs that target ZAM, 30% of those matching Idefix, and 33% of piRNAs complementary to gypsy.

The P element insertions that we analyzed strongly reduced the abundance of piRNAs generated from sequences up to 168 kb away (Figures 4B and 4F). Considering sequences that map uniquely, flamenco produces piRNAs with a marked strand asymmetry that correlates with a strongly biased orientation of transposon fragments in the locus. These observations can be accommodated by a model in which piRNAs are produced from long, unidirectional, precursor transcripts that traverse flamenco. Indeed transcripts containing multiple transposons, and several kb of the flamenco locus can be easily detected by RT-PCR, and these are lost in flamenco mutants (Figure 4G and not shown).

Argonaute3 Binds Sense-Strand piRNAs

Drosophila rasiRNAs show a strong bias for sequences that are antisense to transposable elements (Vagin et al., 2006). We asked whether this observation held for our sequenced piRNAs by examining the strand biases of those derived from Piwi, Aub, and Ago3 complexes. We aligned all piRNA sequences to a comprehensive set of consensus sequences for D. melanogaster transposons (canonical set v9.41, Flybase). Piwi and Aub preferentially incorporate piRNAs matching the antisense strand of transposons (76% and 83%, respectively). In contrast, Ago3 complexes contain piRNAs that are strongly biased for sense transposon strands (75%). As piRNAs derived from total RNA retain an antisense bias, Ago3 complexes must be less abundant overall.

The pattern of asymmetry among the three RNPs was preserved when we evaluated each transposable element separately (Figure 5A). As an example, a plot of piRNAs along the F element reveals numerous antisense piRNAs that are loaded into Piwi and Aub and numerous sense piRNAs that enter Ago3 (Figure 5B). There are a few notable exceptions where asymmetry remains marked but is reversed for Piwi/Aub and Ago3 complexes (see for example accord2, gypsy12, diver2, and hopper2; Figure 5A).

Unlike *flamenco*, transposons within most piRNA clusters lack an orientation bias. For example, the largest piRNA cluster, at 42AB, contains a high density of randomly oriented, nested transposons and produces uniquely mapping piRNAs from both strands (Figure 5C). Even within this cluster, the strand asymmetry of Piwi complexes is preserved. An interesting illustration is two adjacent GATE fragments that are in opposite orientations. Uniquely mapping RNAs in the Ago3 complex correspond to the sense strand of each copy, while Aub, and to some extent Piwi, show the opposite trend (Figure 5C).

Mechanisms of piRNA Biogenesis

To investigate mechanisms of piRNA biogenesis, we examined the relationship between sense and antisense piRNAs. A processing mechanism resembling siRNAs or miRNAs would predict the detection of sense-antisense piRNA pairs that reflect the 2 nt 3' overhangs produced by RNase III enzymes. According to this scenario, complementary sense and antisense piRNAs should have 5' ends separated by 23 nt (assuming an average piRNA size of 25 nt) and correspondingly show 23 nt of complementary sequence.

We plotted the distance between each piRNA 5' end and the 5' end of its neighbors on the opposite strand. Instead of the expected peak at 23 nucleotides, we found that 5' ends of complementary piRNAs are most frequently separated by exactly 10 nucleotides (Figure 6A). On the whole, 20% of all piRNAs have a partner whose 5' end can be mapped ten nucleotides away on the complementary strand. We found the strongest complementary relationships between piRNAs in Ago3 and Aub complexes (Figure 6B). Even though our sequencing efforts are not saturating, more than 48% of small RNAs in the Ago3 library had complementary partners in the Aub library. Statistically significant, although less pronounced, interactions are indicated between Piwi and Ago3, and self complementarity is seen in Aub and Ago3 comparisons.

The 10 nt overlap between sense and antisense piRNAs provoked the hypothesis that the Piwi proteins have a role in piRNA biogenesis. In such a model, an antisense piRNA, derived from a piRNA locus and complexed with Aub or Piwi, would recognize and cleave a sense transposon transcript. This cleavage event would occur opposite nucleotides 10 and 11 of the antisense piRNA, generating a 5' end 10 nt distant, and on the opposite strand, from the end of the original piRNA (see Figures 6A and 7). The cleaved product would be loaded into a second Piwi family protein, likely Ago3 based upon observed strand biases, ultimately becoming new piRNA after processing at the 3' end by an unknown mechanism (see Figure 7). piRNAs generated by Piwi-mediated cleavage events are designated as secondary piRNAs.

elements (IR). Color intensities indicate the degree of the strand bias (green: sense; red: antisense; yellow: unbiased). In the lower panel, the cloning frequency for individual transposons in all three complexes is indicated as a heat map. The key for relative cloning frequency is shown at the left. (B) Shown is the density (sense in green, antisense in red) of all cloned piRNAs assigned to the canonical F element sequence with up to three mismatches. Frequencies in each Piwi-family RNP and in total ovarian RNA are shown individually, as indicated. The relative nucleotide position within the consensus sequence is indicated along the x axes.

(C) Shown is a fragment of the largest piRNA cluster (position 42AB) with only those piRNAs that map uniquely to this region (Ago3 piRNAs in red, Piwi piRNAs in orange, and Aub piRNAs in blue). Top strand piRNAs are shown above the x axes, while bottom strand piRNAs are shown below. Annotated transposon fragments are indicated at the bottom of the panel. Shaded areas mark the boundaries where transposon orientation reverses.

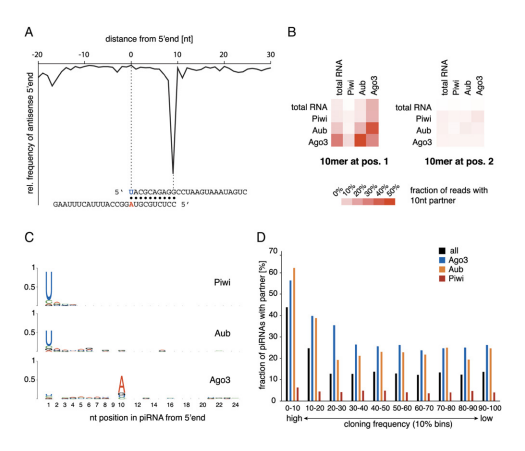


Figure 6. A Slicer-Mediated Amplification Loop for piRNAs

(A) Shown is a frequency map of the separation of piRNAs mapping to opposite genomic strands. The spike at position 9 (the graph starts at 0)

indicates the position of maximal probability of finding the 5' end of a complementary piRNA. One example of this 10 nt offset is shown below. (B) Shown are heat maps that indicate the degree to which complementary 5' 10-mers are found in pairwise library comparisons, with a key to the intensity of the signal shown below. The panel to the right represents a control analysis performed with the 10-mer from position 2–11. (C) The relative nucleotide bias of each position in all piRNAs obtained from Piwi, Aub, and Ago3 complexes, as indicated.
(D) Ten bins were constructed for each Piwi complex (as indicated) and for all sequences combined (all) by sorting piRNAs according to their cloning frequency (e.g., the bin labeled 0–10 contains the 10% of sequences that were most frequently cloned). The fraction of piRNAs within each bin that

(D) Ten bins were constructed for each Piwi complex (as indicated) and for all sequences combined (all) by sorting piRNAs according to their cloning frequency (e.g., the bin labeled 0–10 contains the 10% of sequences that were most frequently cloned). The fraction of piRNAs within each bin that has a complementary partner was graphed on the y axis.

This model is consistent with the known biochemical properties of Piwi and Argonaute proteins (Lau et al., 2006; Liu et al., 2004; Saito et al., 2006). Moreover, it explains the observed lack of a U bias at the 5' end of sense-strand piRNAs in Ago3 complexes. If 5' U-biased antisense piRNAs produce sense piRNAs following cleavage, these should have an unbiased 5' end. However, sense piRNAs should show an enrichment for A, the complement of the 5' U, at position 10 (Figure 7). Strikingly, 73% of all Ago3 bound RNAs conform to this prediction, suggesting that the majority of these arise from piRNA-directed cleavage events (Figure 6C).

DISCUSSION

An Amplification Loop that Reinforces Transposon Silencing

In *C. elegans*, effective RNAi depends upon an amplification mechanism (Sijen et al., 2001). Small RNAs from the

primary dsRNA trigger are largely dedicated to promoting the use of complementary targets as templates for RNA-dependent RNA polymerases (RdRPs) in the generation of secondary siRNAs. In *Drosophila*, no RdRPs have been identified. However, the ability of Piwi-mediated cleavage to prompt the production of new piRNAs could create an amplification cycle that serves the same purpose as the RdRP-driven secondary siRNA generation systems in worms (Figure 7).

The cycle is initiated by generating primary piRNAs, sampled from the piRNA clusters that we have identified. As these are composed mainly of defective transposon copies, they serve as a genetic memory of transposons to which the population has been exposed. piRNAs that are antisense to expressed, dispersed transposons would identify and cleave their targets, resulting in the genesis of a new, sense piRNA in an Ago3 complex. The Ago3 bound sense piRNA would then seek a target, probably a precursor transcript from a master control locus that contains

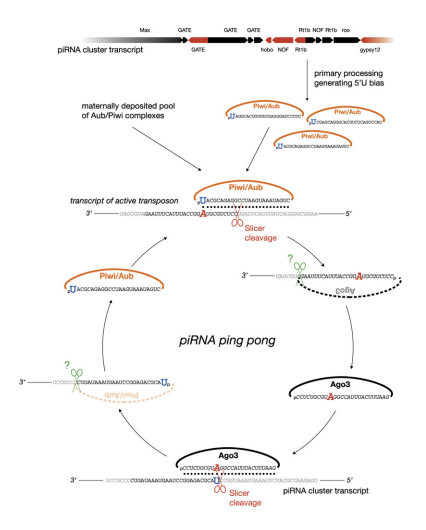


Figure 7. The piRNA Ping-Pong Model Illustrated is the amplification loop consisting of Piwi/Aub complexes, Ago3 complexes, piRNA cluster transcripts, and transcripts of active transposons. Nucleotide cleavage events are shown as scissors. Potential sources of primary piRNAs are piRNA cluster transcripts and maternally inherited piRNA complexes.

antisense transposon sequences. Ago3-directed cleavage would then generate additional antisense piRNAs capable both of actively silencing their target element and reinforcing the cycle through the creation of additional sense piRNAs.

The existence of such an amplification cycle essentially permits sense and antisense piRNAs act in concert to increase production of silencing-competent RNAs in response to the activity of individual transposons. Since Argonautes act catalytically, a significant amplification of the response could be achieved by even a relatively low level of sense piRNAs in Ago3 complexes. This model predicts that piRNAs participating in this process, namely those with complementary partners, should be more abundant than piRNAs without detectable partners. In accord with this hypothesis, the most frequently cloned Aub and Ago3-associated piRNAs show an increased probability of having partners within the data set (Figure 6D). Interestingly, Piwi-associated RNAs do not follow this pattern. Since the amplification cycle consumes target transposon transcripts as part of its mechanism, posttranscriptional gene silencing may be sufficient to explain transposon repression. However, we cannot rule out the

possibility that transcriptional silencing may also be triggered by Piwi-family RNPs.

The amplification cycle may not be absolutely essential for silencing of all elements, as loci such as flamenco may operate in somatic follicle cells where the absence of Aub and Ago3 forces it to act in a stoichiometric fashion. In this regard, flamenco is unusual in that the vast majority of transposon fragments within this locus exist in a common orientation, which can lead to the production of antisense primary piRNAs given a long, unidirectional, precursor transcript.

Origins of Strand Bias in piRNA Populations

In contrast to flamenco, most piRNA loci appear to be both bidirectionally transcribed and contain transposon sequences in random orientation. Nevertheless, the marked asymmetry of Piwi/Aub and Ago3 complexes is conserved in piRNAs that uniquely map to clusters (e.g., Figure 5C). Among piRNAs that match transposons, 77% and 79% of unambiguously cluster-derived Piwiand Aub-associated RNAs are antisense, while 73% of those in Ago3 are sense. These observations strongly suggest that piRNA clusters themselves participate in



an amplification cycle in a manner that informs strand choice. According to our model, the remarkable strand asymmetry in piRNA populations hinges on informative interactions between master control loci and active transposons, which by their nature produce sense RNAs. Our observations identify Ago3 as the principal recipient of piRNAs derived from transposon mRNAs. Thus, as long as there is an input to the system from active transposon transcripts via Ago3 and a preferential relationship between Ago3 and Aub for generating secondary piRNAs in their reciprocal complexes, a strand bias can be maintained even if primary and secondary piRNAs can both be derived from master control loci.

Biogenesis of Primary piRNAs

The amplification cycle must be initiated by primary piRNAs. Presently, the biogenesis pathway that generates primary piRNAs from piRNA clusters remains obscure. Our data suggest that the piRNA precursor is a long, single-stranded transcript that is cleaved, preferentially at U residues. We detect transcripts from piRNA loci by RT-PCR that encompass multiple transposon fragments (not shown) and find numerous small RNAs that cross junctions between adjacent transposons. In the case of flamenco, P element insertions near the proximal end of the locus have a polar effect both on these long RNA transcripts and on flamenco piRNAs.

Equally mysterious is the generation of piRNA 3' ends. Mature piRNAs could arise by two cleavage events and subsequent loading into Piwi complexes. Alternatively, piRNAs could be created following 5' end formation and incorporation of a long RNA into Piwi by resection of their 3' ends. The latter model is attractive, since it could provide an explanation for observed size differences between RNAs bound to individual Piwi proteins, as piRNA size would simply reflect the footprint of each Piwi protein.

Although de novo biogenesis mechanisms must exist, maternally inherited piRNA complexes could also serve to initiate the amplification cycle. All three Piwi proteins are loaded into the developing oocyte (Figure S2) (Harris and Macdonald, 2001; Megosh et al., 2006), and Piwi and Aub are concentrated in the pole plasm, which will give rise to the germline of the next generation. Inherited piRNAs could enhance resistance to transposons that are an ongoing challenge to the organism, augmenting zygotic production of primary piRNAs. Indeed, maternally loaded rasiRNAs were detected in early embryos (Aravin et al., 2003), and their presence was correlated with suppression of hybrid dysgenesis in *D. virilis* (Blumenstiel and Hartl, 2005).

A Model for Transposon Silencing in Drosophila

Our data point to a comprehensive strategy for transposon repression in *Drosophila* that incorporates both a long-term genetic memory and an acute response to the presence of potentially active elements in the genome. The model that emerges from our studies shows many parallels to adaptive immune systems. The piRNA loci

themselves encode a diversity of small RNA fragments that have the potential to recognize invading parasitic genetic elements. Throughout the evolution of Drosophila species, a record of transposon exposure may have been preserved by selection for transposition events into these master control loci, as this is one key mechanism through which control over a specific element can be achieved. Evidence has already emerged that X-TAS can act as a transposition hotspot for P elements (Karpen and Spradling, 1992), raising the possibility the piRNA clusters in general may attract transposons. Once an element enters a piRNA locus, it can act, in trans, to silencing remaining elements in the genome, either directly through primary piRNAs or by engaging in the amplification model described above. A comparison of D. melanogaster piRNAs to transposons present in related Drosophilids shows a lack of complementarity when comparisons are made at high stringency. However, when even a few mismatches are permitted, it is clear that piRNA loci might have some limited potential to protect against horizontal transmission of these heterologous elements (Figure S3). The existence of a feed-forward amplification loop can be compared to clonal expansion of immune cells with the appropriate specificity following antigen stimulation, leading to a robust and adaptable response.

EXPERIMENTAL PROCEDURES

Antibodies

Rabbit polyclonal antisera directed against the N-terminal 14–16 AA of Piwi, Aub, and Ago3 (see Figure S1) were raised and affinity purified as described (Denli et al., 2004). Primary antibody dilutions were 1:2000 for western analysis and 1:500 for immunohistochemistry. Actin staining was with Rhodamine-coupled Phalloidin (1:100). Ovaries were dissected on ice (phosphate-buffered saline) and fixed for 20 min in 4% Formaldehyde/PBS/0.1%Triton X-100.

Immunoprecipitation of Piwi RNP Complexes

Ovary extract was prepared in Lysis buffer (20 mM HEPES-NaOH at pH 7.0, 150 mM NaCl, 2.5 mM MgCl2, 250 mM Sucrose, 0.05% NP40, 0.5% Triton X-100, 1x Roche-Complete). Cleared extracts were incubated with primary antibodies (1:50) for 4 hr at 4°C. Antibody complexes were isolated and analyzed for RNA content as described (Aravin et al., 2006).

Small RNA Cloning and Sequencing

Small RNA cloning was performed as described in Pfeffer et al. (2005), and a detailed protocol is available upon request. Sequencing of small RNA libraries was at 454 Life Sciences.

Quantitative RT-PCR

See Supplemental Data for primer sequences and for the amplification of primary cluster transcripts, *gypsy* transcript, and piRNAs.

Bioinformatic Analysis of piRNAs

Sequence extraction and genomic mapping was as described in Girard et al. (2006). We used the Release 5 assembly of the *Drosophila melanogaster* genome and GenBank to identify all piRNAs matching 100% to genomic sequences. The only GenBank entry (L03284) that recovered hits not present in Release 5 corresponds to the tip of the X chromosome, which differs significantly between the sequenced strain and Oregon R, the strain used for our analysis (Abad et al., 2004). Annotation of small RNAs was done using the following



databases: Repbase (http://www.girinst.org/) on the Release 5 assembly, Transposable element canonical sequences (http://www.fruitfly. org/p disrupt/TE.html). Flybase annotations for coding and noncoding genes (extracted from http://genome.ucsc.edu), and microRNA annotations from Rfam (http://microrna.sanger.ac.uk/sequences). Density analysis of transposons along Release 5 chromosome arms was by counting all nucleotides in a 50 kb window (10 kb offset) annotated as transposons.

piRNA Cluster Analysis

All piRNAs except the 10% of reads corresponding to miRNAs, rRNAs. tRNAs, other ncRNAs, and the sense strand of annotated genes were mapped to Release 5 and the telomeric X-TAS repeat L03284. Nucleotides corresponding to the 5' end of each piRNA were weighted according to N/M with N = cloning frequency and M = number of genomic mappings. We used a 5 kb sliding window to identify all regions with densities greater than 1 piRNA/kb. Windows within 20 kb of each other were collapsed into clusters. Clusters with at least 5 piRNAs that uniquely matched to the cluster were retained.

Analysis of piRNAs Mapping to Transposable Elements

All piRNAs were matched to the canonical sequences of Drosophila transposable elements with high (0 mismatches), medium (3 mm), or low (5 mm) stringency, and the strand relative to the transposon sense strand was determined. We calculated the ratio of all piRNAs per library that match exclusively to the plus or minus strand and excluded those that matched to both (e.g., in inverted repeat [IR] elements). The relative density of piRNAs for each transposon is the number of piRNAs matching a specific element as fraction of all transposonmatching piRNAs.

10 nt Offset Analysis and Nucleotide Bias

The relative relationship of each piRNA to its nearest neighbor was calculated as described in the text, except that all genomic matches were weighted according to the normalized model to avoid skewing by piRNAs that match the genome multiple times. For all subsequent analyses, partners were defined as piRNAs, whose 5' 10 nt are reverse complements. For Figure 6D we sorted all piRNAs in each library according to their cloning frequency and determined the fraction of piRNAs with partners in bins, each of which contained 10% of all sequences. Position-dependent nt biases for each library were determined by their log-odds score relative to library-specific nt compositions.

Supplemental Data

Supplemental Data include four figures, two table, Supplemental Experimental Procedures, and Supplemental References and can be found with this article online at http://www.cell.com/cgi/content/full/ 128/6/■ ■ ■/DC1/.

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Accession Numbers

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