

Ectopic assembly of heterochromatin in *Drosophila melanogaster* triggered by transposable elements

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A persistent question in biology is how *cis*-acting sequence elements influence *trans*-acting factors and the local chromatin environment to modulate gene expression. We reported previously that the DNA transposon *1360* can enhance silencing of a reporter in a heterochromatic domain of *Drosophila melanogaster*. We have now generated a collection of variegating phiC31 landing-pad insertion lines containing *1360* and a heat-shock protein 70 (*hsp70*)-driven *white* reporter to explore the mechanism of *1360*-sensitive silencing. Many *1360*-sensitive sites were identified, some in apparently euchromatic domains, although all are close to heterochromatic masses. One such site (line 1198; insertion near the base of chromosome arm 2L) has been investigated in detail. ChIP analysis shows *1360*-dependent Heterochromatin Protein 1a (HP1a) accumulation at this otherwise euchromatic site. The phiC31 landing pad system allows different *1360* constructs to be swapped with the full-length element at the same genomic site to identify the sequences that mediate *1360*-sensitive silencing. Short deletions over sites with homology to PIWI-interacting RNAs (piRNAs) are sufficient to compromise *1360*-sensitive silencing. Similar results were obtained on replacing *1360* with *Invader4* (a retrotransposon), suggesting that this phenomenon likely applies to a broader set of transposable elements. Our results suggest a model in which piRNA sequence elements behave as *cis*-acting targets for heterochromatin assembly, likely in the early embryo, where piRNA pathway components are abundant, with the heterochromatic state subsequently propagated by chromatin modifiers present in somatic tissue.

epigenetics | position-effect variegation

Transposable elements (TEs) are major structural features of nearly all eukaryotic genomes, and can play a role as *cis*-acting regulatory features with profound influence over the gene regulatory networks of their host (1). However, if left unchecked, the deleterious effects of TE mobilization can become insurmountable for the system, damaging genome integrity. Thus, silencing mechanisms that prevent TE mobilization are fundamental to the faithful propagation of genetic information. Position-effect variegation (PEV), a mosaic expression pattern resulting from silencing in some cells that normally express a gene, has frequently been used as an indicator of heterochromatic environments (2, 3). We reported previously that the DNA transposon *1360* can enhance reporter silencing in repeat-rich heterochromatic regions (4). To gain insight into the mechanisms involved, we sought to identify sequence elements present in *1360* that contribute to *1360*-sensitive PEV, using an adjacent heat-shock protein 70 (*hsp70*)-driven *white* gene as the reporter.

We used the phiC31 landing pad system, where phiC31 integrase mediates recombination between a landing pad (inserted in the genome) containing phage attachment (attP) sites and a donor construct containing bacterial attachment (attB) sites (5). This system allows us to maintain a constant genomic context for multiple transgenic *1360* sequence constructs. A genomic test site identified at the base of chromosome arm 2L (site 1198) indeed exhibits *1360*-dependent silencing, which we find reflects accumulation of Heterochromatin Protein 1a (HP1a).

Testing various features of *1360*, we found that *1360*-sensitive PEV at site 1198 does not require the inverted repeats at the ends of the element, nor internal transcription start sites, but is impacted by small deletions over sites homologous to PIWI-interacting RNAs (piRNAs), a small RNA population initially generated in germ-line tissue and loaded into the zygote. This observation was extended to the retrotransposon *Invader4*, suggesting that the mechanism is broadly applicable. Thus, sites with homology to piRNAs may be *cis*-acting targets for heterochromatin assembly, likely in the early embryo (a stage when piRNA pathway components are abundant), with this decision subsequently propagated in somatic tissue by chromatin-modifying factors.

Results

***1360*-Sensitive Silencing Is Observed in a Subset of Euchromatic Domains.** We generated a collection of landing pad lines by mobilizing a *P* element construct containing attP sites flanking a Flippase FLP-excisable *1360*, cloned upstream of an *hsp70-white* reporter (Fig. 1A). Mobilization led to the recovery of 38 variegating landing pad lines (~5% recovery) with inserts mapping to unique genomic sites (Fig. 1B). Many (34%) mapped to the telomere-associated repeats (TASs) present on 2R and 3R. The subtelomeric repeats are known targets for *P* element insertion (6). These genomic regions are enriched for Polycomb-group proteins (PcG) and the associated trimethylation of histone H3 lysine 27 (H3K27me3) mark (7, 8). In 60% of the variegating lines (including those in TASs), the construct mapped to regions with a high repeat density (>30%), a property of canonical heterochromatin (9). (See Dataset S1 for a list of all lines.) In many of these cases, the construct insertion sites lie in the pericentric heterochromatin of 2L and 2R, in the fourth or the Y chromosome. A subset of lines have insertion sites in regions of low repeat density (<10%) that are enriched for chromatin marks associated with transcriptional activity, such as H3K4me2 and H3K9ac, in both S2 (undifferentiated male embryo cell line) and BG3 (differentiated neuronal cell line) cells (Dataset S1) (7). The distribution of variegating insertions within transcriptionally active domains is not random; all lie within two divisions (or numerical cytological positions) of pericentric or telomeric heterochromatin. These lines represent a unique resource with which to study position effects.

To determine which variegating landing pad reporters are *1360*-sensitive, eye pigment assays were performed comparing 3- to 5-d-old adults with the *1360* element removed by FLP-mediated excision with their sibs with the *1360* element present, i.e., ^{+/−}*1360*. Fourteen lines exhibited pigment levels that were increased in the absence of *1360*, a suppression of PEV (Fig. 1B; also see Dataset S1 and Fig. S1). Some of these lines had

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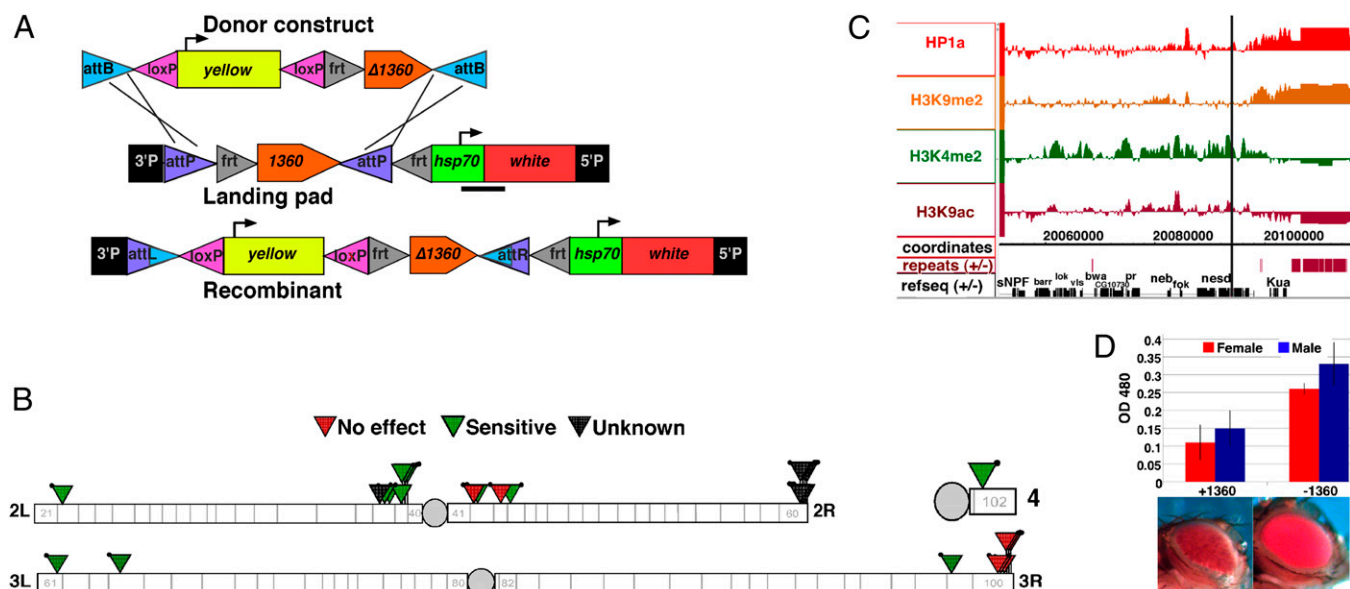


Fig. 1. Sensitivity of PEV to *1360* can occur at euchromatic sites. (A) Schematic of phiC31-mediated cassette exchange between the donor construct and a landing pad, resulting in the recombinant product at the same genomic site. The *yellow* gene, used as a marker to recover recombinants, was removed by *loxP* excision before all assays. The bar below *hsp70-white* denotes the region assayed by ChIP-qPCR in Fig. 2. (B) Map of the landing pad insertions exhibiting variation. Those showing suppressed variation on excision of *1360* are marked by green triangles; those with no change, red triangles (see also Dataset S1). (Black triangles indicate that the line was not assayed.) (C) Chromatin state of the 1198 insertion site based on modENCODE ChIP-array data from third instar larvae (active marks H3K9ac and H3K4me2 and silent marks H3K9me2 and HP1a are shown). A vertical line denotes the P element insertion site within *nesd*. (D) Pigment assays ± 1360 for females (red) and males (blue), with representative eye pictures (see also Fig. S1). Bars represent the mean from two biological replicates (\pm SEM). Excision of the *1360* element results in a loss of silencing.

a reporter inserted within or close to genes, in repeat-poor regions enriched for euchromatic marks in S2 and BG3 cells (7). Thus, *1360* can support heterochromatin formation at a wider variety of sites than previously recognized (4, 10). We chose the *1360*-sensitive landing pad line 1198, with an insert at the base of 2L, distal to a block of heterochromatin (cytological position 38B6, 2L:20094149) for further investigation. This line exhibits >twofold change ± 1360 in eye pigment, a range sufficient to detect partial suppression of *1360*-sensitive silencing (Fig. 1D).

The construct is inserted within the gene *nessun dorma* (*nesd*). A survey of the chromatin environment at the insertion site (in the absence of the P element insert) shows that the region is enriched for marks associated with transcription (H3K4me2 and H3K9ac) in S2 and BG3 cells, as well as in third instar larvae (11, 12) (Fig. 1C). The 20-kb region surrounding the reporter has a repeat density of 2%, consistent with the notion that the reporter is not in canonical heterochromatin. Notably, the eye phenotype in the absence of *1360* corresponds with that observed on insertion of the reporter into a euchromatic environment, displaying a solid red eye; variegation is only observed in the presence of the *1360*. However, in the presence of the *1360* element, pigment levels become comparable to some lines present in annotated heterochromatin (e.g., line 1310; compare line 1198 with *1360* to line 1310 without *1360*; see Fig. S1), indicating that in the presence of a *1360*, the site may be more representative of heterochromatin. The degree of silencing is greater in the female than the male, as is typical of PEV (Fig. 1D). This line gives us an opportunity to study *1360*-sensitive ectopic silencing of a reporter.

1360 Supports Ectopic Heterochromatin Assembly. Heterochromatin assembly is associated with the presence of H3K9me2/3, deposited by a histone methyltransferase (HMT), typically (but not always) SU(VAR)3-9, and the chromo domain protein HP1a, encoded by *Su(var)205* in flies. We assessed eye pigment levels for the 1198 reporter ± 1360 when mutant for either *Su(var)3-9* or *Su(var)205* to test whether these components are required for

1360-sensitive PEV. In the presence of *1360*, a >twofold increase in pigment is detected with each mutant in both sexes, whereas no significant change is observed in the absence of *1360* in males (a minor quantitative change is seen in females), corroborating that *1360*-sensitive PEV is dependent on heterochromatin components (Fig. 2A). Importantly, this also indicates that in the absence of *1360*, HP1a-dependent silencing is not supported at this site.

To determine whether *1360*-sensitive PEV is a consequence of a change in the chromatin landscape, we performed chromatin immunoprecipitation quantitative PCR (ChIP-qPCR) using antibodies for HP1a and H3K9me2, assaying the promoter region of *hsp70-white* (as marked in Fig. 1A). This demonstrated that the local chromatin environment is substantially enriched for HP1a and H3K9me2 in the presence of *1360*, whereas in the absence of *1360*, levels were low (Fig. 2B). No significant difference in the euchromatic mark H3K4me2 was detectable ± 1360 . Thus, the *1360*-sensitive PEV exhibited by line 1198 reflects HP1a-dependent heterochromatin assembly at a normally euchromatic site in the genome.

Cis-Acting Sequence Element in the Right Half of 1360 Is Necessary and Sufficient for 1360-Sensitive PEV. Several features present in *1360* could be used as a signal for heterochromatin assembly. Possibilities include the terminally inverted repeats; putative transcription start sites (TSSs) found within the right half of the element; and regions with similarity to piRNA reads. Terminally inverted repeats can contribute secondary structure or serve as binding sites for transposon-derived proteins, both mechanisms used in plants (13, 14). Present within the right half of *1360* are three putative TSSs, with similarity to the *1360* transcription initiation sites suggested to produce antisense *Suppressor of stellate* transcripts in the *D. melanogaster* male germ line (15). If the integrated copy of *1360* is a target for the transcription machinery, this might cause inappropriate local transcript production, targeting the site for silencing by a similar mechanism.

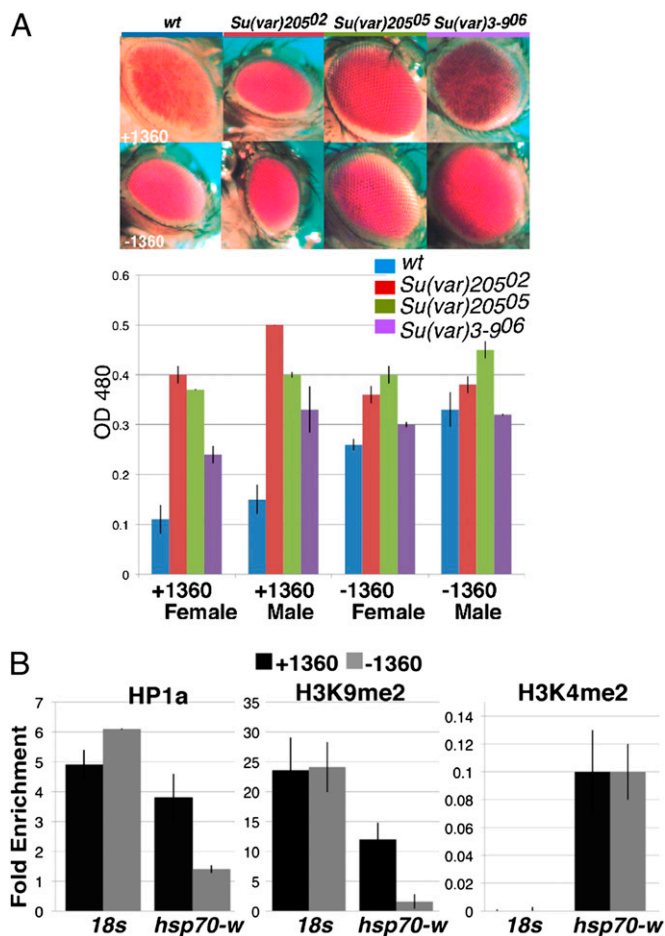


Fig. 2. *1360* promotes heterochromatin assembly. (A) Eye phenotypes (Upper) of line 1198 ^{+/-1360}, comparing the starting line (wt) (Left) with those mutant for *Su(var)205* (HP1a) or the HMT *Su(var)3-9*. (Lower) Corresponding pigment assay readings (OD₄₈₀). Bars give the mean from two biological replicas (\pm SEM). (B) ChIP assessment of levels of HP1a, H3K9me2, and H3K4me2 at the *hsp70-white* promoter for line 1198 +1360 (black bars) and -1360 (gray bars). qPCR primers bracket the *hsp70-white* promoter (see Fig. 1A). All data were normalized to input; fold enrichment over the α -actinin locus is shown (\pm SEM).

The right half also contains sites that map to piRNA reads with an antisense bias; these sites could be targets for antisense piRNAs bound to Piwi, possibly targeting sense transcripts of the integrated *1360* (Fig. 3A).

To identify which sequence features in *1360* contribute to *1360*-sensitive PEV, we tested *1360* fragments lacking the terminal inverted repeats (*1360* Δ IR), left half (*1360* Δ L), or right half (*1360* Δ R) for induction of silencing, as measured by pigment assays (Fig. 3A). Deletion of the right half of *1360* has the most pronounced effect on *1360*-sensitive variegation, resulting in a twofold increase in eye pigment (Fig. 3B; compare red bars). Lines lacking the inverted repeats or the left half of *1360* retained *1360*-sensitive silencing, indicating that these sequence elements are dispensable for *1360*-sensitive PEV.

Because each *1360* element is flanked by flippase recognition target (FRT) sites (Fig. 1A), we can test whether silencing for each line is dependent on the presence of the *1360* Δ construct. FLP-mediated excision of *1360* Δ IR and *1360* Δ L leads to a loss of silencing comparable to the loss of the full-length *1360* donor, whereas excision of *1360* Δ R produced no change in pigment levels (Fig. 3B). Thus, deletion of the left half or inverted repeats of *1360* does not impact *1360*-sensitive PEV, whereas deletion of the right half does, suggesting that sequence feature(s) within the

right half of the element, independent of the terminally inverted repeat, are necessary and sufficient for *1360*-sensitive PEV.

Putative Transcription Start Sites Present in *1360* Are Not Required for *1360*-Sensitive PEV. Although there are no universal core promoter elements in *D. melanogaster*, there are some canonical elements, spanning \sim 80 bp, that are common targets for the RNA Pol II transcription machinery, including the TATA box and the Initiator (Inr) motif (16). The *1360* element in our construct contains one TATA box with a downstream Inr motif close to one of the mapped *Su(ste)* promoter sites, whereas the other two sites have nearby Inr motifs. To test whether these putative promoters are required, we made three 80-bp deletions in *1360* Δ L, each spanning one of these sequence elements, to make *1360* Δ TSS (Fig. 3A). The resulting construct maintains *1360*-sensitive PEV, indicating that the putative TSSs are dispensable for silencing (Fig. 3B). FLP-mediated excision of *1360* Δ TSS led to a \sim twofold increase in pigment levels, confirming that the sequences in *1360* Δ TSS are sufficient to impart silencing (Fig. 3B). Thus, an alternative sequence element(s) must be responsible.

Disruption of Sites Matching Antisense piRNA Reads Compromises TE-Dependent Variegation. The piRNA pathway, previously implicated in *1360*-sensitive PEV (4), suppresses TEs in the germ line through posttranscriptional targeting of sense and antisense transcripts by the slicer-mediated activities of Aubergine (AUB) and Argonaute (AGO)3, using a ping-pong mechanism (17). The repertoire of defective and active TE copies present in the genome can skew the piRNA population to bias against active TEs, necessary to prevent further TE colonization of the genome (18, 19). A third component, Piwi, is a nuclear protein that binds mostly antisense piRNAs, a subset of which are generated by ping-pong cycle-dependent piRNA biogenesis (17, 20). Piwi interacts with HP1a and has been implicated in a transcriptional silencing role in the germ line (21–23). These piRNA components are loaded into the oocyte (17, 24); their mode of TE repression in the early embryo has been relatively unexplored but could include a role in establishing heterochromatin domains. Heterochromatin formation, and the boundary between heterochromatic and euchromatic domains, is established in the syncytium, at nuclear cycle 10–14, and must be propagated by chromatin-modifying mechanisms present in somatic tissue (25).

Large-scale sequencing of piRNA populations has revealed that individual TE families have distinct patterns of sense and antisense piRNAs (20, 26). To determine whether the right half of the *1360* element has a distinguishing piRNA distribution that could indicate key sites for silencing, we mapped the piRNA read density from wild-type (wt) ovary piRNA data generated by Li et al. (20), against the *1360* copy present in our construct, *1360/11503*. Both halves of the element exhibit matches to piRNA reads (Fig. 3A), but antisense peaks (one with a 10-nt overlapping sense and antisense peaks indicative of ping-pong cycle biogenesis) were present at two positions only, in the right half of the element (Fig. 3A). We tested whether these regions contributed to *1360*-sensitive PEV by removing the signature ping-pong overlapping peaks (36-bp deletion) and the neighboring shorter antisense peak (10-bp deletion) of *1360* Δ L to make *1360* Δ piRNA. The new junctions formed by the deletions did not result in new piRNA matches. Indeed, these relatively small deletions, 46 bp in total, significantly compromised the *1360* effect, although a complete loss of *1360*-sensitive silencing was not observed (Fig. 3B).

Whether the observed loss of silencing was directly associated with the *1360* Δ piRNA construct was tested by assaying PEV after FLP excision of this *1360* fragment. Indeed, a 1.3-fold change was observed, indicating that much of the capacity to silence had been lost and that the residual silencing was attributable to the *1360* remnant (Fig. 3B). The results indicate that this 46-bp

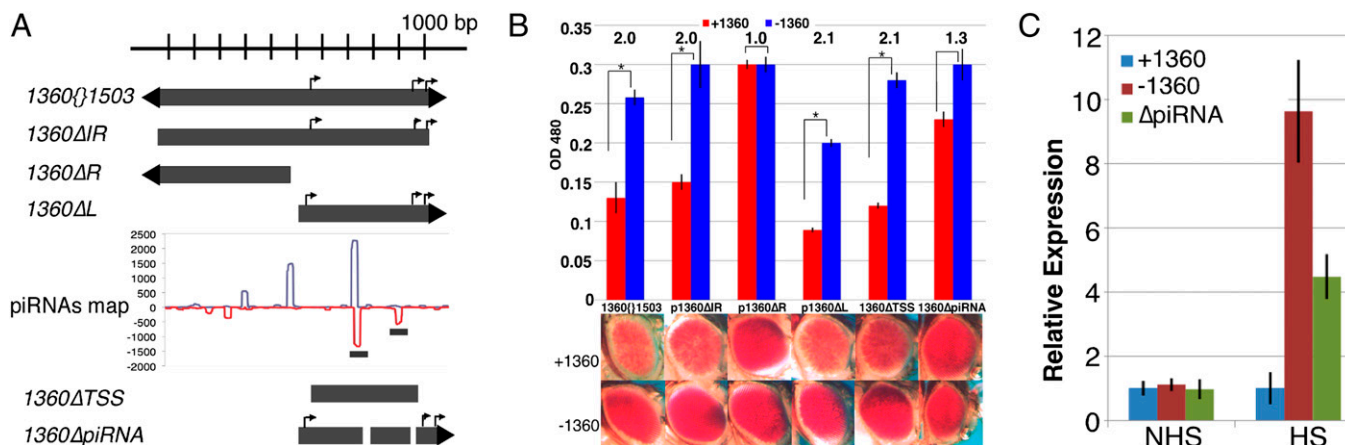


Fig. 3. piRNA sequence elements in the right half of *1360* support *1360*-sensitive silencing. (A) The *1360* variants used in donor constructs for phiC31-mediated cassette exchange. Bent arrows indicate putative transcription start sites. Frequencies of sense (blue line) and antisense (red line) piRNA reads (20) that map to *1360Δ1503* are shown. The bars below illustrate the deletions for the *1360ΔpiRNA* constructs. (B) Representative eye pictures for each *1360Δ* recombinant line are shown under the pigment assay data (from male flies) using OD₄₈₀ (± SEM). Asterisks indicate a significant change +*1360* (red) vs. –*1360* (blue) with a *P* value of <0.05 (Student's *t* test); numbers indicate fold change. Loss of the right half of *1360Δ1503* or loss of the piRNA hotspots results in a significant loss of silencing (no or reduced change on excision of the *1360* remnant). (C) RT-qPCR of *hsp70-w* transcript levels in non-heat-shocked (NHS) and heat-shocked (HS) adult flies heterozygous for the *P* element. Shown is fold expression (normalized to *RpL32*) relative to full-length *1360* recombinant lines (± SEM).

sequence component contributes significantly to *1360*-dependent silencing but is not the sole sequence responsible for the effect. To verify that the PEV assay data were truly a reflection of a change in silencing (from less capacity for transcription +*1360* to more capacity for transcription –*1360*), we performed RT-qPCR for *hsp70-w* driven *white* transcripts in 3- to 5-d-old non-heat-shocked (NHS) and heat-shocked (HS) adults, comparing the full-length and *1360ΔpiRNA* recombinant lines. Transcript levels for *hsp70-w* at 25 °C are sufficient to generate the eye phenotype. Heat shock increases the level of the appropriate transcription factor and provides a more robust test of the accessibility of the promoter region, producing levels of transcript suitable for a quantitative assay (3, 4). Indeed, the presence of *1360* significantly decreases the capacity for expression under heat shock; this capacity is restored on loss of *1360* and partially restored with the *1360ΔpiRNA* construct (Fig. 3C). These findings indicate that piRNA sequence elements present in *1360* contribute to *1360*-sensitive heterochromatin assembly.

To explore whether piRNA pathway components could impact *1360*-sensitive PEV at this site, we tested reporter line 1198 both ^{+/–}*1360* in the presence of *piwi* and *aub* mutant alleles. Mutations in both genes dominantly suppressed *1360*-sensitive PEV (Fig. S2). This finding implicates an RNAi mechanism in establishing this *1360*-sensitive silencing.

To determine whether this phenomenon is unique to element *1360*, we swapped *1360* with the long terminal repeat (LTR) element *Invader4* (Fig. 4A). *Invader4* was chosen for two reasons. First, many variegating reporters generated for this study (Fig. 1B) are located within or in close proximity to an *Invader4* element. [Similar criteria were used initially to select *1360* as a potential target for heterochromatin assembly (10).] Second, *Invader4* is regulated similarly to *1360* by piRNA pathway components in that transcription from both elements is up-regulated in *aub*, *ago3*, and *piwi* ovaries (17, 20). When *Invader4* replaces *1360* in the test construct, we found an equivalent level of variegation (Fig. 4B); this loss of expression was also suppressed by mutations in heterochromatin and piRNA components (Fig. S3). Deletion of sites with homology to piRNA reads (*Invader4ΔLTR*; Fig. 4A) resulted in an increase in pigment levels, similar to the loss of full-length *Invader4* [compare silencing by the full-length element (Left) with that by the truncated element (Right) in Fig. 4B]. This evidence supports the suggestion that piRNA sequence

elements can promote silencing; importantly, the phenomenon is not exclusive to *1360* and can be observed using either a DNA transposon- or a retrotransposon-derived element.

Work from plants and fungi argues that such a recognition event would likely be based on RNA-RNA base pairing (27), suggesting a requirement for transcription over the transposable element to generate a target template. Although the putative TSSs in *1360* are dispensable for *1360*-sensitive PEV at this insertion site, read-through transcription could occur from a nearby promoter. In this case, the *P* element is inserted downstream of the transcription start site for *nesd*. This gene is expressed at low levels in the female ovary and at moderately high levels in the early embryo (28). To determine whether read-through transcription of the *P* element is occurring, we looked for transcripts across the *P* element junction by RT-PCR in 0- to 10-h embryos; such transcripts are detected (Fig. 4C). The combined results support a model in which a small RNA-targeting event using this transcript participates in the HP1a-dependent assembly of heterochromatin at this site in the early embryo, with the consequences of that event being evident in the tissues that go on to form the eye (see model in Fig. S4).

Discussion

Our screen to identify a *1360*-sensitive phiC31 landing pad site generated many landing pad lines with inserts in unique chromatin domains, including several on the fourth chromosome and in repetitious elements such as the telomere-associated sequences on chromosomes 2R and 3R (Fig. 1). This study adds useful resources to the collection of phiC31 lines currently available (29). We find that reporters in a wide range of chromatin domains are sensitive to *1360*; thus, the influence of *1360* is not limited to repeat-rich, heterochromatic sites. However, proximity to repeat-rich regions appears to be important, because most *1360*-sensitive reporters are close to the base of the euchromatic arms or are telomere-proximal (Fig. 1B). Genetic analysis and ChIP experiments show that *1360*-sensitive PEV is representative of HP1a-dependent heterochromatin assembly (Fig. 2). The use of phiC31 recombineering technology allowed us to show that a unique sequence element within the right half of *1360*, independent of the inverted repeat terminal sequences and of putative transcription start sites, is required for optimal *1360* impact in line 1198 (Fig. 3). Importantly, an unrelated transposable element (*Invader4*) has

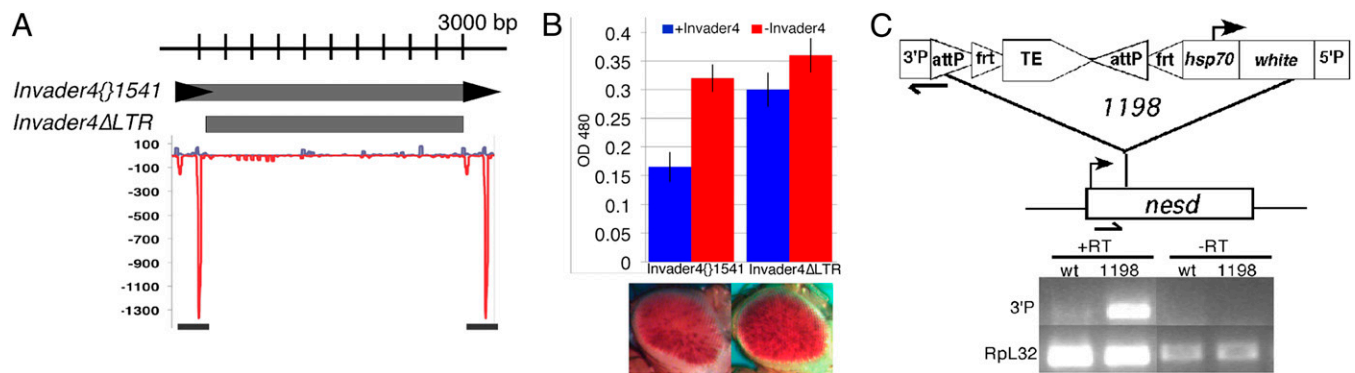


Fig. 4. *Invader4* piRNA sequence elements support PEV. (A) *Invader4* variants used in donor constructs for phiC31-mediated cassette exchange are shown. Frequencies of sense (blue line) and antisense (red line) piRNA reads mapping to *Invader4*{1541} are shown. The bar below denotes regions of *Invader4* deleted in *Invader4*ΔLTR. (B) Pigment assay data using OD₄₈₀ (± SEM) is shown for ^{+/-}*Invader4* test constructs. Loss of regions with homology to piRNA reads (*Invader4*ΔLTR) results in a loss of *Invader4*-sensitive PEV. (C) Diagram of the P element insertion site for line 1198 in the 5' end of *nesd*. Black bars represent primers used for RT-PCR (Upper). Read-through transcripts spanning the 3'P region of the P element (3'P) and control *RpL32* products are shown (Lower). Lines used were wt control (no P element insert) and 1198, where ^{+/-}RT indicates the addition or omission of reverse transcriptase.

the same effect as *1360*, implicating a mechanism that can be applied to other TEs (Fig. 4A and B). In both cases, the deletion of sites with homology to antisense-oriented piRNAs compromised silencing at this ectopic site. This suggests that a small RNA-directed, RNA-RNA targeting event is contributing to transposon-sensitive heterochromatin assembly, potentially using a mechanism similar to that documented in *Schizosaccharomyces pombe* and plants (Fig. S4) (30–33).

The ability to trigger ectopic HP1a assembly appears to be limited to genomic sites close to heterochromatic masses, because the distribution of *1360*-sensitive sites is limited (Fig. 1B). Indeed, half of *1360*-sensitive sites are in annotated heterochromatin (chromatin states 7/8, BG3 cells), whereas most sites inducing variegation but lacking *1360* sensitivity (without a change in pigment levels ^{+/-}*1360*) are in PcG-enriched regions, frequently in telomere-associated repeats (Dataset S1). Thus, *1360*-sensitive PEV appears to be an HP1a-dependent phenomenon. Prior reports, as well as this study, find that single repetitive elements within the euchromatic arms do not trigger detectable reporter silencing (4). The high density of repetitive elements in heterochromatic domains is a fundamental characteristic of these regions, suggesting that these repeats may cooperatively participate in stabilizing heterochromatin factors. Cooperative function of interspersed signals has been reported for other domain-wide chromatin structures, for example that developed to provide dosage compensation (34). Many of the *1360*-sensitive sites identified here are surrounded by repeats of different types; thus at different insertion sites, additional mechanisms could be at work. In the future it will be of interest to determine if our observations extend to other *1360*-sensitive sites, both those adjacent to and within annotated heterochromatin. It is probably no coincidence that our experimentally recovered test site (Fig. 1C) resembles the classical rearrangements that result in PEV, having a reporter in a normally euchromatic domain close to a heterochromatic mass.

Evidence for piRNA-mediated effects on chromatin assembly in flies is mixed, with data suggesting both silencing and activating roles, depending on reporter location (4, 35, 36). Mutations in piRNA pathway components are weak suppressors of PEV for a reporter in a repeat-rich environment, with and without *1360* present, suggesting that both local heterochromatin and *1360*-sensitive silencing are impacted (4). Other TEs with piRNA signals include *HeT-A*, *I* element, and *copA* (20). All three transposon families are silenced by HP1a and H3K9me2/3 assembly over their promoter elements in the female germ line by a mechanism dependent on RNAi components (22, 37). piRNA biogenesis is most active at this stage, and it has been

argued that its effects may be limited to germ cells. For example, the RNA helicase Spindle E, required for piRNA biogenesis, is necessary for TE repression in the female germ line but not in mature somatic tissues (no-ovary carcasses) (37). Mutations in many of the RNAi components and chromosomal proteins result in female sterility, making the issue difficult to study.

Both piRNA and chromatin structural proteins (and/or their mRNAs) are synthesized during oogenesis, loaded into the ovary, and present in the early embryo (0–6 h) (24). Their presence overlaps the early stages of heterochromatin formation (cycles 11–14) and of zygotic transcription (nuclear cycle 14) (25). The latter coincides with the appearance of the histone H3K4 demethylase SU(VAR)3-3, without which HP1a and H3K9me2 levels in the heterochromatin become substantially reduced, whereas levels for the activating H3K4me2 mark become high (immunofluorescent staining of nuclei in cycle 14 embryos), resulting in suppression of PEV. Generation of wt clones from heterozygous *Su(var)3-9* lines suggests that the chromatin patterns established in the late embryo persist during differentiation (25). Thus, piRNA components could play a role in initiation of heterochromatin formation in the early embryo at select sites (e.g., a subset of TEs) in the genome, with the structure maintained during development by chromosomal protein interactions. However, whether or not depletion of these products in the female germline/early embryo alone is sufficient to see an impact in the larvae/adult remains to be tested.

Although the impact on silencing of deleting the piRNA sites in *1360* and *Invader4* is striking, full suppression of the PEV induced by *1360* was not observed (Figs. 3B and 4B), which suggests that additional sites may be critical. Such sequences could be missing from the small RNA library used for our analysis. Additional piRNA libraries (17) examined did not identify additional sites within the right half of the *1360* element. The piRNA libraries reported to date do not achieve saturation, so it is possible that additional piRNA targets remain.

Alternatively, a different mode of targeting could also be in effect, unrelated to a piRNA-mediated event. Such dual mechanisms have been documented for heterochromatin formation in *S. pombe*, a system in which RNAi-mediated heterochromatin targeting is well-established (38, 39). In many eukaryotes, transposases have been found to be sources of DNA binding domains with chromatin-modifying abilities (1). For example, *Drosophila* BEAF-32, derived from the *hAT* transposase, is a chromatin insulator that binds the *scs* chromatin boundary element (40). Similar mechanisms may have evolved in *Drosophila* to specifically

target heterochromatin factors to TEs. However, no candidate proteins have emerged to date.

Evidence for an RNAi-based mechanism that contributes to heterochromatin assembly has been found in several model systems (worms, plants, fission yeast). In *S. pombe*, targeting of the HP1 homolog Swi6 and the H3K9 HMT Ctr4 depends on the processing of RNA Pol II transcripts (generated from heterochromatic loci) by RNAi components. The RNAi-induced transcriptional silencing complex (RITS) contains the chromo domain protein Chp1, as well as the RNAi component Ago1, which binds small RNAs with homology to target sites (e.g., *dg/dh* repeats) in pericentric heterochromatin (38). In the system described here, the read-through transcripts of the *nesd* gene detected in early embryo samples (Fig. 4C) are plausible targets for Piwi associated with *1360* or *Invader4* piRNAs, using a base-pairing interaction. Analysis of EST libraries from *Drosophila* embryos indicates that most TE families are transcribed (41). However, determining the origin of these transcripts is a challenging task given the high sequence similarity among TE family members. The current experiments have not allowed us to determine whether the *1360*-sensitive silencing observed is transcription-dependent, as would be required for an RNA-RNA recognition event for targeted silencing; such a test will be critical for future work to establish the mechanism involved. The present results support a mechanism that uses piRNAs, possibly ping-pong cycle-derived [as are *1360* and *Invader4* piRNAs (20)], for transposon-sensitive targeting of HP1a, likely early in development, with persistent effects observed using reporters in adult tissues.

Materials and Methods

Fly Stocks. All *Drosophila* stocks and crosses were maintained at 25 °C on cornmeal sucrose-based media (42). Fly stocks were from the Bloomington *Drosophila* Stock Center unless otherwise indicated. Transgenesis of the landing pad construct into the starting stock *yw*^{67c12} was carried out by Genetic Services. Stocks and constructs used, plus the mobilization and mapping procedures, are described in detail in *SI Materials and Methods*.

Chromatin Immunoprecipitation. Chromatin isolation and immunoprecipitation from third-instar larvae were carried out as described previously (12). The antibodies used were HP1a W191, Abcam 1220 H3K9me2, and Millipore 07-030 H3K4me2. Antibodies were validated by us and by others: see the Antibody Validation Database (<http://compbio.med.harvard.edu/antibodies/about>) (43). Quantitative PCR (see Table S1 for the list of primers) was performed using Bio-Rad iQ SYBR Green Supermix in a Cepheid SmartCycler. Two biological replicates, each consisting of two technical replicates, were assayed for each immunoprecipitation assay.

Mapping piRNA Reads and Assessing RNA Products. Direct-sequence mapping was carried out using small RNA sequence reads derived from wt Oregon R ovaries (20) retrieved from the National Center for Biotechnology Information (NCBI) trace archives with accession number SRP000458 (Figs. 3A and 4A). For RT-PCR, RNA was isolated from 0- to 10-h embryos (Fig. 4B) or 3- to 5-d adult flies, with or without heat shock (Fig. 3C), DNase I-treated, and reverse-transcribed using random hexamer primers (Fig. 4B) or oligo dT (Fig. 3C). See *SI Materials and Methods* for details.

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