Silencio/CG9754 connects the Piwi–piRNA complex to the cellular heterochromatin machinery

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The repression of transposable elements in eukaryotes often involves their transcriptional silencing via targeted chromatin modifications. In animal gonads, nuclear Argonaute proteins of the PIWI clade complexed with small guide RNAs (piRNAs) serve as sequence specificity determinants in this process. How binding of nuclear PIWI–piRNA complexes to nascent transcripts orchestrates heterochromatin formation and transcriptional silencing is unknown. Here, we characterize CG9754/Silencio as an essential piRNA pathway factor that is required for Piwi-mediated transcriptional silencing in Drosophila. Ectopic targeting of Silencio to RNA or DNA is sufficient to elicit silencing independently of Piwi and known piRNA pathway factors. Instead, Silencio requires the H3K9 methyltransferase Eggless/SetDB1 for its silencing ability. In agreement with this, SetDB1, but not Su(var)3-9, is required for Piwi-mediated transcriptional silencing genome-wide. Due to its interaction with the target-engaged Piwi-piRNA complex, we suggest that Silencio acts as linker between the sequence specificity factor Piwi and the cellular heterochromatin machinery.

[Keywords: H3K9 methylation; Piwi; transposon silencing; heterochromatin formation; piRNA pathway; transcriptional silencing]

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In analogy to heterochromatin formation in *S. pombe*, the PIWI–piRNA complex likely provides the sequence-specific information to target chromatin-modifying enzymes to TE loci for their silencing. However, how recruitment of nuclear Piwi clade proteins to nascent transcripts leads to heterochromatin formation is unknown.

In *S. pombe*, the adaptor protein Stc1 is reported to bridge the Ago1/siRNA complex to the methyltransferase Ccr4 [Bayne et al. 2010], and the chromodomain protein Chp1 is implicated in tethering the target-bound Ago1/siRNA complex to chromatin [Ishida et al. 2012]. However, neither of these factors is conserved in animals. In turn, two proteins with largely unknown functions that are required for piRNA-guided transcriptional silencing in animals [Gtsf1/Asterix and Maelstrom] [Sienski et al. 2012; Donertas et al. 2013; Muerdter et al. 2013; Ohtani et al. 2013] are absent in *S. pombe*, highlighting significant molecular differences in these two systems. Moreover, the Piwi–piRNA complex fails to mediate transcriptional silencing if recruited artificially [not involving a piRNA–target interaction] to a reporter RNA in a *Drosophila* cell culture model that expresses a fully functional piRNA pathway centered on nuclear Piwi [Post et al. 2014]. Whether this reflects the necessity for a conformational change in Piwi upon piRNA–target binding or is due to the requirement of a minimum threshold of target-engaged Piwi molecules to initiate silencing is unclear. Similarly unclear is the identity of the factors that establish and maintain piRNA-specified heterochromatin. For example, evidence for an involvement of the two H3K9 methyltransferases Su(var)3-9 and Eggless/SetDB1 has been reported [Fritsch et al. 2010; Rangan et al. 2011; Huang et al. 2013; Bulut-Karslioglu et al. 2014], but it is not known which roles these enzymes have in piRNA-guided silencing and what they contribute to heterochromatin formation at TE insertions.

Here, we establish an in vivo assay to test the sufficiency of piRNA pathway factors in mediating transcriptional silencing and heterochromatin formation. This led to the discovery of CG9754/Silencio [Czech et al. 2013; Handler et al. 2013; Muerdter et al. 2013] as a key factor that links target-engaged Piwi to the cellular heterochromatin machinery. Using this assay and genome-wide profiling of the H3K9me3 mark, we demonstrate an integral role of Eggless/SetDB1 in piRNA-guided silencing and show that Su(var)3-9 contributes to spreading of the H3K9me3 mark to TE-flanking domains. Our work provides an entry point into the mechanistic dissection of piRNA-guided silencing and clarifies the roles of H3K9 methyltransferases in this process.

### Results

**Recruitment of CG9754 to RNA or DNA elicits potent transcriptional silencing**

Several studies indicate that binding of the Piwi–piRNA complex to a nascent target RNA causes heterochromatin formation and transcriptional silencing [e.g., Sarot et al. 2004; Sienski et al. 2012; Le Thomas et al. 2013; Post et al. 2014]. In agreement with the study by Lau and colleagues [Post et al. 2014], we found that recruitment of Piwi to a reporter construct via a sequence stretch complementary to endogenous piRNAs elicits potent reporter silencing in a Piwi-dependent and target orientation-dependent manner [Supplemental Fig. S1A]. In contrast, artificial recruitment [tethering] of Piwi via the λ-5 boxB system [thereby bypassing the requirement for piRNA target sites] to a reporter RNA expressed from a transiently transfected plasmid does not affect reporter expression [Supplemental Fig. S1B; Post et al. 2014].

To determine whether the λ-5 boxB assay failed due to the reporter plasmid not being properly chromatinized, we established an in vivo RNA-tethering system in the *Drosophila* ovary [Fig. 1A; Supplemental Fig. S1C,D]. This assay builds on the α-tubulin promoter driving the ubiquitous expression of a GFP reporter, which harbors five boxB sites in its 3′ untranslated region (UTR), 1.5 kb downstream from the transcriptional start site (TSS). Expression of λ-5 HA-tagged Gawky/GW182 [a cytoplasmic factor that elicits post-transcriptional gene silencing] [Jonas and Izaurralde 2015] with the germline-specific MTD-Gal4 driver leads to potent reporter silencing in germline but not somatic cells of the ovary [Fig. 1A,B]. Tethering of Piwi to the reporter instead fails to induce silencing despite λ-5-targeted Piwi being abundant in the nucleus [Fig. 1A,B, Supplemental Fig. S1C]. This indicates that Piwi is only able to signal to the downstream silencing machinery if engaged with a target via its bound piRNA.

We reasoned that an adaptor protein might link target-engaged Piwi to the silencing machinery and therefore asked whether any factor that has been implicated in Piwi-mediated transcriptional silencing [Fig. 1C] triggers repression if recruited ectopically to the reporter. While Gtsf1/Asterix and Maelstrom were inert in this assay, tethering of CG9754, an uncharacterized protein with a suggested role in Piwi-mediated silencing [Czech et al. 2013; Handler et al. 2013; Muerdter et al. 2013], results in potent reporter repression that was dependent on the λ peptide on CG9754 [Fig. 1D].

Silencing induced by tethered CG9754, but not by GW182, results in strongly reduced steady-state reporter RNA levels [Fig. 1E] and an approximately threefold reduction of RNA Pol II occupancy at the α-tubulin promoter of the reporter construct. The endogenous α-tubulin promoter instead [identical in sequence except for a small stretch of SNPs] exhibits constant Pol II occupancy [Fig. 1F]. These findings indicate that CG9754 is sufficient to induce transcriptional silencing if targeted to a nascent RNA. In support of this, ectopic recruitment of CG9754 to DNA upstream of the TSS of a different GFP reporter leads to similarly potent silencing [Fig. 1G; Supplemental Fig. S1E,F]. Also in this assay, Piwi, Gtsf1, and Maelstrom are incapable of inducing silencing.

We next combined the λ-5 CG9754-mediated silencing assay with potent germline-specific RNAi to establish a genetic hierarchy between CG9754 and other piRNA pathway factors [Supplemental Fig. S1G]. Depletion
Figure 1. CG9754 elicits transcriptional silencing upon recruitment to nascent RNA or DNA. [A] The cartoon depicts the GFP-5xboxB reporter used for RNA tethering. It harbors the ubiquitous α-tubulin promoter, the GFP-coding sequence, and a 3’ UTR harboring five boxB sites. The confocal image below depicts GFP fluorescence signal in an egg chamber expressing the GFP-boxB reporter. Throughout this study, we converted GFP signal to monochrome images with an inverted black/white color scheme for displaying optimal contrast. Soma and germline tissue is indicated. Images at the right depict the expression of λN-HA fusion proteins under control of the germline-specific MTD-Gal4 driver (λN-GW182: cytoplasmic; λN-Piwi: nuclear-enriched). [B] GFP fluorescence in egg chambers expressing the ubiquitous GFP-boxB reporter and the indicated λN fusions in the germline. [C] Simplified cartoon of the linear piRNA pathway feeding into Piwi, also indicating key players involved at the indicated steps. (TGS) Transcriptional gene silencing, [D] GFP fluorescence in egg chambers expressing the ubiquitous GFP-boxB reporter and the indicated λN fusions in the germline. Expression of Lacl-CG9754 served as a negative control. [E] Shown are changes in GFP-boxB steady-state RNA levels in ovaries expressing the indicated λN fusions (normalized to rp49). n = 3. Error bars indicate SD. [F] Shown is RNA Pol II occupancy at the indicated loci determined by chromatin immunoprecipitation (ChIP) followed by quantitative PCR using ovaries expressing the indicated λN fusions. n = 3. Error bars indicate SD. Signals were normalized to those at the TSS of the act 5C gene, arrestin is not expressed in ovaries. [G] The cartoon depicts the 8x lacO-GFP reporter used for DNA tethering. This reporter is expressed in germline and somatic cells of the ovary (nanos promoter coupled to intronic piwi enhancer) (Hayashi et al. 2014). The confocal image below depicts GFP fluorescence signal of the reporter. The middle image depicts the expression of Lacl-CG9754 under control of the germline-specific MTD-Gal4 driver. Images at the right show GFP fluorescence signal of the reporter in egg chambers expressing the indicated Lacl fusions in the germline. Note that Lacl-Piwi accumulates mostly in the cytoplasm (potentially due to the reported oligomerization of Lacl) (Supplemental Fig. S1E) and thus cannot be interpreted unambiguously. [H] Shown are GFP fluorescence signals in egg chambers expressing the GFP-boxB reporter, λN-CG9754, and the indicated shRNA constructs in the germline.
of λN-CG9754/C CG9754 results in robust reporter derepression, demonstrating the feasibility of this approach (Fig. 1H). Depletion of Piwi, Zucchini (an essential piRNA biogenesis factor for Piwi), Gts1/Asterix, or Maelstrom instead had no impact [Fig. 1H]. Considering that CG9754 is required for all aspects of Piwi-mediated silencing, this factor acts most likely downstream from all known piRNA pathway factors.

Taken together, our data indicate that recruiting CG9754 to chromatin—directly to DNA or indirectly via a nascent RNA—elicits transcriptional silencing of the target locus that is reminiscent of piRNA-guided silencing yet does not require known piRNA pathway factors.

**CG9754 is an essential piRNA pathway factor in the ovarian germline and soma**

CG9754 was identified genetically as a factor required for TE silencing in the ovary using transgenic RNAi (Czech et al. 2013; Handler et al. 2013; Muerdter et al. 2013). We used CRISPR/Cas9 (Jinek et al. 2012; Gokcezade et al. 2014) to generate two mutant alleles of CG9754 that harbor nucleotide insertions/deletions in the coding sequence close to the N terminus. Both alleles result in frameshifts and are expected to be strong mutant alleles [Fig. 2A]. In support of this, CG9754 protein is not detectable in ovarian lysates from homozygous mutants by Western blot analysis using a monoclonal antibody raised against CG9754 [Fig. 2B].

Flies homologous for either mutant allele as well as transheterozygous flies are viable yet exhibit oogenesis defects and severe fertility defects [Fig. 2C,D]. In this, they resemble *piwi* mutant flies (Cox et al. 2000). In addition, patterns of TE derepression [for soma- and germline-specific TEs] are very similar between CG9754 mutants and *piwi* mutants [Fig. 2E]. In support of previous findings (Handler et al. 2013), complete loss of CG9754 does not affect levels and nuclear accumulation of Piwi in germline and somatic TEs.

![Figure 2](image-url)
cells [Fig. 2B,F], indicating that CG9754 is specifically required for Piwi-mediated transcriptional target silencing (also, localization and levels of Aub and Ago3 in germline cells are unaffected) [Supplemental Fig. S2].

CG9754/Silencio is required for H3K9 methylation and transcriptional silencing of Piwi-repressed TE insertions

In order to directly probe a role of CG9754 in heterochromatin formation and transcriptional silencing downstream from Piwi, we used cultured ovarian somatic cells (OSCs) [Niki et al. 2006; Saito et al. 2009]. The detailed knowledge of their TE insertion landscape [Sienski et al. 2012] and the ability to efficiently deplete factors of interest using RNAi [Fig. 3A] make these cells an ideal system for genome-wide studies. To determine changes in H3K9me3 patterns and RNA Pol II occupancy, we focused on 381 euchromatic insertions of the 11 TE families that are strongly repressed by the piRNA pathway in OSCs. The genomic regions flanking these TE insertions are genome-unique anchor points that allow extracting levels and spreading of the H3K9 methyl mark as well as the transcriptional activity of each individual TE insertion [Fig. 3B].

Depletion of CG9754 leads to a strong reduction in H3K9me3 levels at Piwi-repressed TE insertions and to directional transcriptional readthrough into flanking genomic regions [Fig. 3C]. The resulting patterns are highly similar to those obtained from Piwi-depleted cells [Sienski et al. 2012]. Similarly, patterns of TE derepression as well as changes in the expression of protein-coding genes are highly correlated between CG9754-depleted versus Piwi-depleted cells [Fig. 3D–F]. As observed previously [Sienski et al. 2012], many of the up-regulated genes are flanked by Piwi-repressed TE insertions, leading to their silencing in wild-type cells by the spreading of heterochromatin [e.g., expanded] [Fig. 3F]. Altogether, our data from flies [Fig. 2] and OSCs [Fig. 3] assign a specific and essential role to CG9754 in local heterochromatin formation and transcriptional silencing downstream from Piwi. We therefore named CG9754 “silencio.”

CG9754/Silencio associates with target-engaged Piwi–piRNA complexes

As Silencio is required for Piwi-mediated silencing and induces silencing if tethered ectopically to an RNA, we tested for an interaction between Piwi and Silencio. Reciprocal coimmunoprecipitation (co-IP) experiments indicated that a large part of the nuclear Silencio pool interacts with Piwi [Fig. 4A]. Because of Silencio’s silencing capacity, we wondered whether it interacts with a distinct subset of nuclear Piwi. We therefore sequenced small RNAs isolated from Silencio immunoprecipitations as well as from Piwi immunoprecipitations and compared both with the population of cellular piRNAs from total small RNA libraries. The majority of the small RNAs obtained from the Silencio immunoprecipitation resemble mature piRNAs, as evidenced by their average length of 26.7 nucleotides [nt] and their tendency for a terminal 5’ uridine (~80%). Interestingly, piRNAs originating from 3’ UTRs of cellular mRNAs [Robine et al. 2009; Saito et al. 2009], but not from TEs, are specifically depleted in the Silencio immunoprecipitation library but not in the Piwi immunoprecipitation library [Fig. 4B; Supplemental Fig. S3A,B]. As 3’ UTR-derived piRNAs have no widespread role in target identification due to the absence of highly complementary RNAs, these results were a first hint that target-engaged Piwi–piRNA complexes might be enriched in Silencio immunoprecipitations.

Upon closer inspection of the small RNA populations in the Silencio immunoprecipitation, we noticed a considerable portion of RNA reads that map in sense orientation to Piwi-repressed TE sequences. These TE sense reads are specifically enriched in Silencio immunoprecipitations but not in Piwi immunoprecipitations from the same nuclear OSC lysate [Fig. 4C]. In this context, it is important to note that TE-derived piRNAs in OSCs are almost exclusively antisense to the active transposon transcript (e.g., mdg1, 99.9%; gypsy, 99.4%; 412, 99.99%).

Several observations suggest that the observed sense RNAs represent footprints of endogenous piRNA targets. First, their size profile mirrors that of mature piRNAs yet, on average, is 1–2 nt larger, suggesting ~1-nt extensions on both sides of a small RNA duplex [Fig. 4C], whether this size profile of sense reads also exists in vivo or is generated by nucleases during the experiment is currently unclear. Second, the sense reads lack a uridine bias at the 5’ end, arguing that they are not mature piRNAs [Fig. 4C,D]. Instead, their three terminal nucleotides at the 3’ end are enriched for adenosine residues, which would be expected due to the 5’ uridine bias of complementary piRNAs. Third, the sense reads map predominantly to TEs that are repressed by Piwi–piRNA complexes in OSCs [e.g., gypsy, mdg1, 412, and Tabor] [Fig. 4E]. Finally, the sense read profiles are essentially a mirror image of antisense piRNA populations. piRNAs targeting mdg1, for example, exhibit a gap in the 5’ portion of the TE sequence, and this gap is also devoid of corresponding sense reads [Fig. 4F].

These observations suggest that target-engaged Piwi–piRNA complexes interact with Silencio, further supporting a central role for this factor in the effector step of piRNA-guided transcriptional silencing.

CG9754/Silencio induces heterochromatin formation, and its transcriptional silencing capacity depends on SetDB1/Eggless

A hallmark of piRNA-guided transcriptional silencing in flies and mice is the local establishment of an H3K9me3 domain. To test whether experimental tethering of Silencio to the RNA of the GFP-boxB reporter also leads to H3K9me3, we performed chromatin immunoprecipitation (ChiP) followed by quantitative PCR (qPCR) from ovaries where An alone, An-GW182, An-Piwi, or An-Silencio was expressed in germline cells [Fig. 5A]. Only tethering of Silencio results in H3K9me3 specifically at the reporter locus (TSS and internal) to levels that are comparable with those at the light locus, which maps to heterochromatin. In contrast, the endogenous α-
tubulin promoter that is identical in sequence to the reporter except for a few SNPs does not change methylation status. Surprisingly, tethering of HP1a/Su(var)205, a factor that is known to induce potent transcriptional silencing and heterochromatin formation if targeted to DNA (Ayyanathan et al. 2003; Li et al. 2003), does not trigger silencing or H3K9me3 [Fig. 5A].

We obtained very similar results when targeting Silencio to DNA upstream of the lacO-GFP reporter (Fig. 5B). In this assay, HP1a was equally potent to induce silencing and H3K9me3. This strongly indicates that Silencio not only orchestrates heterochromatin formation [like HP1a] but also integrates unknown functions that allow it to effectively signal to the underlying DNA locus.
when targeted to a nascent RNA, which is only transiently associated with chromatin via the transcribing polymerase.

The trimethylation of H3K9 triggered by Silencio recruitment to the GFP reporter prompted us to investigate which histone methyltransferase mediates this
Modification. To do so, we depleted the *Drosophila* H3K9 methyltransferases Su(var)3-9 and Eggless/SetDB1 in the germline using transgenic RNAi in the background of the GFP-boxB reporter silenced by λN-Silencio. Loss of SetDB1, but not of Su(var)3-9, results in derepression of the boxB reporter in the germline (Fig. 5C). Similar results were obtained when Silencio was targeted to the reporter at the DNA level (Fig. 5D). SetDB1 is also essential for HP1a-induced reporter silencing. This instead is independent of Silencio (Supplemental Fig. S4)—an expected result, as it is a piRNA pathway-specific factor, while HP1a is a general heterochromatin effector in all cells. Based on these findings, we conclude that SetDB1 is the central effector methyltransferase that acts downstream from Silencio and HP1a.

**Distinct roles of SetDB1 and Su(var)3-9 at piRNA targeted TE insertions**

Previous studies have implicated both Eggless/SetDB1 and Su(var)3-9 in the *Drosophila* piRNA pathway (Rangan et al. 2011; Huang et al. 2013). However, no systematic survey on the role of these two chromatin modifiers downstream from Piwi has been reported. We therefore used OSCs and established potent siRNA-mediated depletion conditions for Egg/SetDB1, Su(var)3-9, and G9a, an

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**Figure 5.** Silencing and heterochromatin formation by CG9754/Silencio depends on the H3K9 methyltransferase SetDB1. (A) Shown are GFP fluorescence signals in egg chambers expressing the GFP-boxB reporter and the indicated λN fusions. The bar diagram shows corresponding H3K9me3 levels at the indicated loci, measured by ChIP followed by qPCR (values were normalized to a gene desert; light is a heterochromatic control locus). Displayed is the mean of three independent experiments. Error bars indicate SD. The cartoons detail the reporter locus and the endogenous α-tubulin locus in scale and depict the location of used primer pairs. (B) Everything is as in A except for the lacO-GFP reporter experiment. (C) Shown are GFP fluorescence signals in egg chambers expressing the GFP-boxB reporter, λN-CG9754, and the indicated shRNA constructs in the germline [two independent shRNA lines for Su(var)3-9 were used with similar results]. (D) As in C except for the lacO-GFP reporter.
enzyme that might contribute to H3K9 methylation (Supplemental Fig. S5A,B; Stabell et al. 2006). Genome-wide profiling of the H3K9me3 mark as well as of RNA Pol II occupancy shows that only depletion of SetDB1 impairs heterochromatin formation and transcriptional silencing at piRNA targeted TE insertions (Fig. 6A). In contrast,

Figure 6. Distinct roles for Eggless/SetDB1 and Su(var)3-9 at piRNA-repressed TE insertions. (A) Heat maps showing H3K9me3 signal (green) and RNA Pol II occupancy (red) on the 10 kb flanking the 381 euchromatic TE insertions targeted by Piwi in OSCs. Signals from GFP-depleted cells (control) were compared with cells depleted for CG9754, SetDB1, Su(var)3-9, G9a, or HP1a. All heat maps were sorted for the average H3K9me3 intensity in the control experiment. Below each heat map, area charts represent the average signal. (B) Box plot showing changes in H3K9me3 levels at euchromatic H3K9me3 peaks (enriched in Piwi-repressed TE insertions) and heterochromatic H3K9me3 peaks in OSCs depleted for the indicated factors. The identities of analyzed H3K9me3 peaks are as in Sienski et al. (2012). (C) Bar graph showing mean levels of piRNAs derived from flamenco and cluster 20A (each bar is the average value of piRNA densities in 40 nonoverlapping 1-kb windows) in OSCs depleted for the indicated factors (values were normalized for those present in GFP knockdown control cells). Error bars correspond to SEM. (D) Jitter plot showing fold changes in mRNA levels of TEs in the indicated knockdowns. The top panel displays TEs regulated by Piwi. Fold change, >4, n = 11. (****) P-value < 0.0001, according to Mann-Whitney test. The bottom panel displays TEs notregulated by Piwi. n = 36. (E) Metaprofile showing HP1a occupancy at the genomic regions flanking the 146 euchromatic gypsy insertions in OSCs depleted for the indicated factors.
H3K9me3 levels within constitutive heterochromatin, which are established independently of Piwi (Sienski et al. 2012), are insensitive to SetDB1 depletion and instead are moderately affected in Su(var)3-9-depleted cells (Fig. 6B), supporting a role of Su(var)3-9 in the maintenance of constitutive heterochromatin. The impact of SetDB1 loss on TE silencing in OSCs is not a result of reduced piRNA levels (Rangan et al. 2011), as depletion of SetDB1 does not impair piRNA production from flamenco or cluster 20A, the two dominant piRNA sources in OSCs (Fig. 6C). Thus, at least in somatic ovarian cells, SetDB1 acts specifically in target repression and is not required for piRNA cluster transcription.

In addition, transcriptome profiling by RNA sequencing (RNA-seq) indicates that the steady-state transcript levels of Piwi-repressed TEs are only increased upon depletion of SetDB1 (Fig. 6D, Supplemental Fig. S5C). We note that TE desilencing in SetDB1-depleted cells is considerably weaker than in Piwi-depleted cells. As the simultaneous knockdown of SetDB1 and either of the two other methyltransferases does not lead to higher TE derepression (Fig. 6D, Supplemental Fig. S5C), this most likely does not reflect redundancies among the methyltransferases. Instead, it could be due to the different kinetics of H3K9me3 loss in SetDB1 versus Piwi-depleted cells.

Further support for the integral role of the H3K9me3 mark in piRNA-guided TE silencing stems from the observation that depletion of HP1α, a major reader of the H3K9me3 mark, leads genome-wide to strong transcriptional increases at piRNA targeted TE insertions—again, without affecting global piRNA levels (Fig. 6A,C; Supplemental Fig. S5D; Ohtani et al. 2013). Consistent with this, HP1α is present at piRNA-targeted TE insertions as well as at the flanking genomic regions in a Piwi-dependent manner (Fig. 6E,F).

Although loss of Su(var)3-9 does not impair Piwi-mediated TE silencing, a closer inspection of the H3K9me3 profiles at piRNA-repressed TE insertions reveals impaired spreading of the H3K9me3 mark to flanking genomic regions in the absence of this enzyme (Fig. 6A,G; Supplemental Fig. S5F). A calculation of the spreading index (distance from TE insertion at which the signal is 50% of the peak) indicates nearly twofold-reduced H3K9me3 spreading in Su(var)3-9-depleted cells (upstream, 1.1 kb vs. 1.8 kb, downstream, 1.6 kb vs. 2.8 kb). This is most likely a direct effect, as Su(var)3-9 is enriched at piRNA targeted TE insertions (including their immediately flanking genomic regions) to levels that are comparable with those found in constitutive heterochromatin (e.g., light and parp) (Fig. 6F). Similar roles in heterochromatin formation and spreading have been attributed to SetDB1 and Su(var)3-9 in mouse embryonic stem cells (Fritsch et al. 2010, Bulut-Karslioglu et al. 2014), indicating a conserved division of labor between these two H3K9 methyltransferases in heterochromatin biology.

Our data support a model in which Piwi–piRNA complexes target SetDB1 in a sequence-specific manner to TE insertions in order to initiate a repressive heterochromatin state, while Su(var)3-9 assists in local bidirectional spreading of the methyl mark, probably by binding to pre-existing H3K9me3 marks via its own chromodomain or HP1α (Lachner et al. 2001; Nakayama et al. 2001).

Discussion

The formation of heterochromatin [with the resulting transcriptional silencing of the locus] downstream from nuclear Argonaute proteins is an intriguing process. On the one hand, this is because the target RNA, which is essential for Argonaute targeting, will be suppressed in its own production. On the other hand, the process must deal with the fact that transcripts are only transiently attached to their originating chromatin locus, transcript maturation or transcription termination should in principle lead to the release of the target RNA, including bound Argonaute complexes. A wealth of pioneering work in S. pombe on centromeric heterochromatin formation downstream from Ago1 has provided significant insight into these questions [see above for references]. However, how these findings relate to the molecular steps that are involved in transcriptional target silencing downstream from animal PIWI clade proteins is only poorly understood. In fact, the involvement of several nonconserved proteins between yeast and animals suggests significant differences.

Here, we characterize CG9754/Silencio as an essential piRNA pathway factor required for nuclear Piwi–piRNA complexes to mediate transcriptional silencing and heterochromatin formation. The central conclusions from our work are as follows: (1) Silencio is the first protein in the piRNA pathway that triggers heterochromatin formation and silencing if tethered ectopically to RNA or DNA. In this, our work provides a powerful entry point into the mechanistic dissection of the silencing process in a controlled setting. (2) We show that the two central H3K9 methyltransferases, Su(var)3-9 and Eggless/SetDB1, are acting downstream from Piwi yet in different and nonredundant manners: While SetDB1 is required for heterochromatin formation downstream from Piwi, Su(var)3-9 appears to be involved in spreading the H3K9me3 to flanking genomic areas [see also Brower-Toland et al. 2009].

The molecular function of Silencio

The primary sequence of Silencio does not reveal insights into its molecular roles in the piRNA-guided silencing process. Silencio does not harbor any domain of known function. Also, orthologs of Silencio are not identifiable outside Drosophilids. However, considering that heterochromatin formation downstream from nuclear PIWI proteins also occurs in mammals (e.g., Pezic et al. 2014), the existence of a factor with equivalent function seems very plausible outside Drosophilids.

How Silencio triggers heterochromatin formation is unclear. Our data suggest that it directly or indirectly recruits SetDB1 to target loci or licenses SetDB1’s methylation activity. The specificity for this process almost certainly relies on Piwi–piRNA complexes. In fact, Silencio interacts with target-engaged Piwi. In this sense,
Silencio could be considered as an adaptor protein that bridges Piwi to basic cellular activities involved in the establishment and maintenance of heterochromatin. Such a model is highly reminiscent of the biology of TE silencing in mammals, where, besides the gonadal piRNA pathway, a large family of KRAB-type zinc finger transcription factors recruits the adaptor protein KAP1/Trim28 to target TE sequences [Rowe et al. 2010]. KAP1 acts as a binding platform for various chromatin effector proteins, including SetDB1 and HP1a. The concept of an adaptor protein is also evident for small RNA-guided post-transcriptional silencing, where GW182 bridges the sequence-specific AGO–miRNA complex to the cellular pathways that mediate mRNA turnover and translational repression [Behmann et al. 2006].

The most remarkable aspect about Silencio is its ability to trigger heterochromatin formation if tethered to the nascent RNA. In fact, overcoming the transient interaction of the target RNA with chromatin does not seem to be a trivial task, as tethering of HP1α, which is a potent silencer if recruited directly to promoter regions, is unable to induce heterochromatin formation and silencing under these conditions. Whether Silencio links to activities that tether the target RNA to chromatin or whether this indicates that Silencio and HP1α signal to different downstream effectors with different silencing capacities or kinetics is unclear. In this respect, it is interesting to note that tethering of Ago1 as well as other RITS (RNA-induced transcriptional silencing) complex components to nascent transcripts in S. pombe is very inefficient in initiating heterochromatin as well [Buhler et al. 2006]. A recent report indicates that impairing transcription elongation by loss of the Paf1 complex enables trans-silencing by RITS, presumably as the residence time of the nascent transcript at the target locus is increased [Kowalik et al. 2015].

The role of SetDB1 and Su(var)3-9 in piRNA-guided heterochromatin formation

Our data point to a major role of Eggless/SetDB1 in the establishment of H3K9me3-marked heterochromatin downstream from nuclear Piwi and Silencio. Consistent with this, loss of SetDB1, but not of Su(var)3-9, leads to the derepression of piRNA targeted TEs as well as a genome-wide loss of H3K9me3 at targeted TE insertions. In contrast to Su(var)3-9, which harbors an H3K9me3-binding chromodomain, SetDB1 seems to require other specificity determinants as a guide (e.g., piRNAs and KRAB zinc fingers). Whether SetDB1 is required for only the initiation or also the maintenance of the H3K9me3 domain is currently unclear and will require sophisticated experimental systems similar to those established in yeast [Audergeron et al. 2015; Ragunathan et al. 2015]. In terms of TE repression, Su(var)3-9 plays a less important role downstream from Piwi. Instead, it seems to primarily mediate the spreading of the H3K9me3 mark to genomic regions that are flanking piRNA targets. This is very consistent with a central role of Su(var)3-9 in position effect variegation [Tschiersch et al. 1994; Rea et al. 2000; Schotta et al. 2002]. Also, phenotypically, the described division of labor between these two methyltransferases is consistent: eggless/SetDB1 mutant flies are subviable, and escapers exhibit severe oogenesis defects and are sterile [Clough et al. 2007]. Instead, Su(var)3-9 mutant flies are viable and fertile.

The precise role of H3K9 methylation in piRNA-guided heterochromatin formation is not clear. It might serve as a binding platform for downstream effector complexes such as chromatin remodelers or histone deacetylase complexes, as has been shown previously [Czernin et al. 2001; Sugiyama et al. 2007]. In mammals, it is probably also involved in guiding the de novo DNA methylation at CpG dinucleotides [Matsui et al. 2010]. Alternative/additional roles of the H3K9me3 mark could be to aid higher-order chromatin structures [Sun et al. 2001] or control the subnuclear localization of silenced loci [Towbin et al. 2012] as well as prevent the ectopic recombination of TE loci that are identical in sequence at many genomic locations [Ellermeier et al. 2010].

Clearly, we are only beginning to reveal the molecular events that occur downstream from a Piwi–piRNA complex upon interacting with a complementary target transcript at chromatin. Understanding the mechanistic logic of piRNA-guided heterochromatin formation promises to provide a more general insight into the connections and interplay between heterochromatin formation and transcription.

Materials and methods

Genetics and transgenes

The RNA-tethering constructs (subcloned into pUASP/attB or act15C vectors) express the N domain [Baron-Benhamou et al. 2004] followed by a single HA tag fused to the N terminus of the full-length ORFs of GW182, Piwi, Gtsf1, Maelstrom, CG9754, or HP1α. The empty N-HA vector served as a control. The DNA-tethering constructs express LacI fused to GFP, Piwi, Gtsf1, Maelstrom, CG9754, or HP1α. For the boxB sensor construct, the α-tubulin-84D promoter drives GFP cDNA followed by five boxB sites in the 3′ UTR. In the DNA sensor construct, eight lacO sites precede a nanos promoter expressing a GFP-Pwi cDNA (including the first intron of piwi, which causes expression in somatic cells) [Hayashi et al. 2014]. The UASP-N and UASP-LacI transgenes were inserted into attP40, and the sensor constructs were inserted at attP2 [Markstein et al. 2008]. Other transgenes were MTD-Gal4 and shRNAi lines [all inserted into attP2] targeting white, piwi, zuc, gtsf1, maelstrom, CG9754, eggless, and Su(var)3-9 [Ni et al. 2009; Olivieri et al. 2012; Sienski et al. 2012; Donertas et al. 2013; Mohrb et al. 2014]. Novel CG9754 alleles were isolated in a white1118 background isogenic for the second and third chromosomes using two independent guide RNAs (Supplemental Table S2) and were compared with Cas9-generated piwi alleles that have the same genetic background [Senti et al. 2015].

Antibodies

Mouse monoclonal antibodies (used for Western blotting and immunoprecipitations) against His-tagged full-length CG9754/Silencio were generated by the Max F. Perutz Laboratories
Monoclonal Antibody Facility. Rabbit antisera (used for immunofluorescence) were raised against the MEAPMKLEVKENVY-VECGI NC peptide of CG9754. Rabbit anti-Piwi is described in Brennecke et al. (2007), rabbit anti-Rbp3 was raised against full-length Drosophila Rbp3, and rabbit anti-Eggsense was raised as described in Clough et al. (2007). The commercially available antibodies used were MAb 16B12 anti-HA (Covance), MAb anti-LacI (Abcam), anti-H3K9me3 39161 [ActiveMotif], anti-RNA Pol II 8WG16 (Abcam), MAb CIA9 anti-HPLa [Developmental Studies Hybridoma Bank], and rabbit polyclonal PRB291C anti-HPLa (Covance). Rabbit anti-Su(var)3-9 was a kind gift of G. Reuter (Schotta et al. 2002), and rabbit ant-dG9a was a kind gift of M. Yamaguchi (Kato et al. 2008).

Cell culture

OSCs were cultured as described (Niki et al. 2006; Saito et al. 2009) and transfected with Cell Line Nucleofector kit V (Amamax Biosystems, program T-029). The reporter constructs were cloned into pAcM (Saito et al. 2010) and cotransfected with siRNA tar-

Transposon qPCR analysis

Primer sequences used for RT-qPCR are listed in Supplemental Document S1.

RT-qPCR

Table S1.

Primer sequences used for RT-qPCR are listed in Supplemental Document S1.

Computational analyses

For details regarding the computational analyses, see Supplemental Document S1.

Accession numbers

Illumina deep sequencing data sets were deposited at NCBI Gene Expression Omnibus (GSE74097).

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Note added in proof

Hannon and colleagues (Yu et al. 2015) independently identified CG9754 as a central bridging factor between Piwi and the heterochromatin machinery and named this factor Panoramix.
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