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PIWI-piRNA pathway-mediated transposable element repression in *Hydra* somatic stem cells

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ABSTRACT

Transposable elements (TEs) can damage genomes, thus organisms use a variety of mechanisms to repress TE expression. The PIWI–piRNA pathway is a small RNA pathway that represses TE expression in the germline of animals. Here we explore the function of the pathway in the somatic stem cells of *Hydra*, a long-lived freshwater cnidarian. *Hydra* have three stem cell populations, all of which express PIWI proteins; endodermal and ectodermal epithelial stem cells (ESCs) are somatic, whereas the interstitial stem cells have germline competence. To study somatic function of the pathway, we isolated piRNAs from *Hydra* that lack the interstitial lineage and found that these somatic piRNAs map predominantly to TE transcripts and display the conserved sequence signatures typical of germline piRNAs. Three lines of evidence suggest that the PIWI–piRNA pathway represses TEs in *Hydra* ESCs. First, epithelial knockdown of the *Hydra piwi* gene *hywi* resulted in upregulation of TE expression. Second, degradome sequencing revealed evidence of PIWI-mediated cleavage of TE RNAs in epithelial cells using the ping-pong mechanism. Finally, we demonstrated a direct association between Hywi protein and TE transcripts in epithelial cells using RNA immunoprecipitation. Altogether, our data reveal that the PIWI–piRNA pathway represses TE expression in the somatic cell lineages of *Hydra*, which we propose contributes to the extreme longevity of the organism. Furthermore, our results, in combination with others, suggest that somatic TE repression is an ancestral function of the PIWI–piRNA pathway.

Keywords: PIWI; piRNA; Hydra; transposable elements; stem cells; aging

INTRODUCTION

The PIWI-piRNA pathway is best known for repressing transposable element (TE) expression in animal germlines (Siomi et al. 2011). The central effector complex of this pathway consists of PIWI proteins partnered with a class of small RNAs, the PIWI-interacting RNAs (piRNAs), which are typically 24–31 nucleotides (nt) in length (Aravin et al. 2006; Girard et al. 2006; Grivna 2006; Vagin 2006). TE repression is directed by cytoplasmic PIWI-piRNA complexes that target TE RNAs for degradation (Brennecke et al. 2007; Gunawardane et al. 2007). In some species, PIWIpiRNA complexes can also act in the nucleus to silence TE genomic loci (Watanabe et al. 2018; Batki et al. 2019). In addition to germline function, somatic expression and function for piwi genes and piRNAs has been demonstrated for many organisms, although in most cases the somatic targets have not been well explored (Mani and Juliano 2013; Ross et al. 2014; Ozata et al. 2019). In Drosophila, the *piwi* gene functions in several somatic tissues to repress TEs, including the ovary, fat body, and intestinal stem cells (Malone et al. 2009; Jones et al. 2016; Sousa-Victor et al. 2017). One possibility is that somatic TE repression is a derived function of the pathway in *Drosophila*, but two recent studies demonstrate widespread expression of PIWI family genes and TE-derived piRNAs in somatic tissues of arthropods and mollusks, suggesting that somatic repression of TEs is an ancestral function of the pathway in at least the protostome lineage (Jehn et al. 2018; Lewis et al. 2018).

To better understand the ancestral function of the PIWI-piRNA pathway in metazoans requires careful analysis of PIWI-piRNA pathway function in nonbilaterian taxa, such as cnidarians, which are the sister group to the bilaterians (Dunn et al. 2014). Recent work in the sea anemone *Nematostella vectensis*, a cnidarian, demonstrates that PIWI-piRNA pathway genes are expressed in both embryonic and adult somatic tissue and that a *piwi* gene is required for metamorphosis (Praher et al. 2017; Modepalli

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et al. 2018). Furthermore, degradome analysis demonstrated that the PIWI-piRNA pathway targets TEs in *Nematostella*, but the cell-type specificity of this targeting has not been characterized (Praher et al. 2017). Here we aim to identify somatic targets of the PIWI-piRNA pathway in *Hydra*. This cnidarian has several experimental advantages for such studies, including well-described somatic stem cell populations which express *piwi* genes, transgenic lines that allow for the collection of specific cell lineages for RNA analysis, and reagents for isolating PIWI-piRNA complexes (Juliano et al. 2014).

Hydra is composed of three cell lineages: ectodermal epithelial, endodermal epithelial, and interstitial; each lineage is supported by its own population of stem cells (Fig. 1A). The basic Hydra body plan consists of two epithelial monolayers (Fig. 1A). No fully undifferentiated stem cells exist to renew the epithelium. Instead, all body column epithelial cells are mitotically active, leading to the displacement of cells toward the extremities. At the oral and aboral ends, epithelial cells stop dividing and terminally differentiate to build the tentacles, hypostome, and basal disk. The mitotic ectodermal and endodermal body column cells therefore simultaneously function as epitheliomuscular cells and as epithelial stem cells (ESCs). Cells of the interstitial lineage include neurons, gland cells, the stinging nematocytes specific to cnidarians, and germ cells (Fig. 1A). This lineage is supported by an undifferentiated multipotent interstitial stem cell (ISC) population that continually produces neurons, gland cells, and nematocytes in a homeostatic adult animal. ISCs are also capable of producing germline stem cells (GSCs) when GSCs are experimentally depleted (Bosch and David 1987; Nishimiya-Fujisawa and Kobayashi 2012). Therefore, while ISCs have germline potential, ectodermal and endodermal ESCs are strictly somatic.

Hydra has two cytoplasmic PIWI proteins, Hywi and Hyli, which are located in perinuclear, cytoplasmic granules found in both ESCs and ISCs (Juliano et al. 2014; Lim et al. 2014). Hywi and Hyli participate in the conserved "ping-pong" piRNA biogenesis pathway that is well described in animal germlines. Analyzing piRNAs isolated from whole Hydra demonstrated that a large fraction of Hywi-bound piRNAs map to TEs in an antisense orientation, and a large fraction of Hyli-bound piRNAs map to TEs in a sense orientation. These piRNAs exhibit a typical "ping-pong" signature, which consists of a 10 bp overlap between the 5' end of Hywi-bound piRNAs (initiator piRNAs) and the 5' end of Hyli-bound piRNAs (responder piRNAs) (Juliano et al. 2014; Lim et al. 2014; Ozata et al. 2019). These data indicated that Hywi and Hyli likely participate in ping-pong mediated cleavage of TE RNAs, a conserved mechanism that occurs in the cytoplasmic granules of germ cells (Brennecke et al. 2007; Aravin et al. 2008). In addition, a recent study found evidence of phased piRNA production in Hydra, a conserved mechanism of piRNA biogenesis triggered by the initial ping-pong cleavage event (Gainetdinov et al. 2018). After cleavage, the remaining 3' portion of the target RNA is fragmented into head-totail phased trailing piRNAs, which are sense with respect to the target RNA (Han et al. 2015; Homolka et al. 2015; Mohn et al. 2015; Gainetdinov et al. 2018).

Hywi knockdown in Hydra ESCs led to loss of epithelial integrity and death within 12 d, thus demonstrating a necessary function for the PIWI-piRNA pathway in Hydra somatic stem cells (Juliano et al. 2014). However, our previous study focused on piRNAs isolated from whole animals and did not directly test the identity of PIWI-piRNA pathway targets in ESCs. Therefore, the identity of PIWIpiRNA targets in the somatic ESCs was unknown and was the subject of this study. Here we report that a primary target of the PIWI-piRNA pathway in Hydra ESCs is TE RNAs. This targeting occurs through the conserved ping-pong mechanism, which couples piRNA biogenesis to TE destruction (i.e., the RNA transcripts of TEs are processed into piRNAs). This is supported by our findings that TE-derived piRNAs are prevalent in Hydra ESCs and that TE transcripts are up-regulated in response to epithelial hywi knockdown. Using degradome sequencing, we find evidence of PIWI-directed cleavage products of TE transcripts in somatic cells using the ping-pong mechanism, and we demonstrate a direct interaction between Hywi protein and target TE RNAs in ESCs. These observations demonstrate that the PIWI-piRNA pathway represses TEs in the somatic stem cells of Hydra, which further supports that TE repression is an ancestral function of the pathway in not only the germline, but also in somatic tissues.

RESULTS AND DISCUSSION

Hydra vulgaris AEP piRNAs have conserved sequence characteristics

In our previous study, we isolated and sequenced Hywibound and Hyli-bound piRNAs from Hydra vulgaris strain 105 and mapped these to genome and transcriptome assemblies generated from the same strain. These data suggested that the Hydra PIWI-piRNA pathway targets TEs through the conserved ping-pong mechanism (Juliano et al. 2014). However, we were not able to discern cell lineage-specific roles for the pathway because piRNAs were isolated from whole Hydra vulgaris strain 105. TE repression is a conserved function of the PIWI-piRNA pathway in germ cells (Aravin et al. 2007; Brennecke et al. 2007; Carmell et al. 2007; Houwing et al. 2007; Das et al. 2008; Kuramochi-Miyagawa et al. 2008), but targets in somatic cells are less well understood (Ross et al. 2014). Therefore, in this study we focused on identifying targets of the pathway specifically in the ESCs.

To understand lineage-specific functions of the PIWIpiRNA pathway in *Hydra*, we used transgenic lines with



FIGURE 1. Epithelial Hydra piRNAs have sequence signatures typical of ping-pong biogenesis. (A) Schematic of the Hydra body plan and cell types adapted from Siebert (2018). Hydra is comprised of three cell lineages. The two epithelial cell lineages, endoderm (end, green) and ectoderm (ect, blue), form two monolayers separated by a basal lamina called the mesoglea. The cells of the interstitial lineage (pink) are embedded among the epithelial cells; this lineage is supported by a multipotent ISC which gives rise to three somatic cell types: neurons, gland cells, and nematocytes. Interstitial cells shown: progenitors (prog), nematoblasts (nb), nematocyte (nem), ganglion neuron (gn), sensory neuron (sn), and zymogen gland cell (zmg). The ISCs are also able to produce GSCs (not shown). In contrast, the epithelial cell lineages are strictly somatic. The epithelial cells of the body column (shown in lighter colors) are mitotically active, unipotent stem cells that express PIWI proteins (Juliano et al. 2014). Due to mitotic divisions, these cells are translocated toward the oral and aboral ends of the animal. When epithelial cells arrive at the extremities (shown in darker colors), the cells exit the cell cycle, lose PIWI expression, and terminally differentiate to build the hypostome and tentacles at the oral end and the peduncle (ped) and basal disc (bd) at the aboral end. (B) Experimental design for Figures 1, 2: piRNAs were extracted from either untreated (wild-type, contains germline-competent interstitial lineage) or colchicine-treated (epithelial, does not contain interstitial cells) Hydra by immunoprecipitation (IP) using Hywi- and Hyli-specific antibodies (Juliano et al. 2014). piRNAs were isolated from immunoprecipitated complexes, sequenced, and mapped to the Hydra transcriptome (Siebert et al. 2019). (C-J) The number of occurrences of complementary overlap between the 5' ends of Hywi- and Hyli-bound piRNAs; all four possible combinations were tested. "S" indicates piRNAs that map in a sense orientation to a transcript, and "AS" indicates piRNAs that map in an antisense orientation to a transcript. The high frequency of 10 bp overlap between Hywi- and Hyli-bound piRNAs is indicative of ping-pong biogenesis. This signal is present in piRNAs isolated from both wild-type (WT) (C-F) and epithelial (G-J) Hydra, indicating that the ping-pong biogenesis pathway is active in ESCs. The highest frequency of 10 bp overlaps occurs between antisense Hywi piRNAs and sense Hyli piRNAs in both WT (C) and epithelial Hydra (G). This is consistent with a previous study analyzing piRNAs isolated from WT Hydra vulgaris strain 105 (Juliano et al. 2014).

cell-lineage or cell-type specific expression of fluorescent proteins. These lines were generated using Hydra vulgaris strain AEP that produces eggs in the laboratory and is routinely used for the production of transgenic lines (Wittlieb et al. 2006). As this strain differs from the strain used in our previous study, Hydra vulgaris strain 105, which does not produce eggs in the laboratory and cannot be used to make transgenic lines, we began this study by testing if the two Hydra strains have the same piRNA characteristics. We first established that our Hywi and Hyli antibodies are able to immunoprecipitate PIWI-piRNA complexes from the AEP strain (Supplemental Fig. S1A-D). Next, we isolated and sequenced Hywi-bound and Hyli-bound piRNAs from the AEP strain, mapped these piRNAs to an AEP transcriptome (Fig. 1B; Siebert et al. 2019), and analyzed the sequences for evidence of ping-pong piRNA biogenesis. Similar to previous data from the 105 strain (Juliano et al. 2014; Lim et al. 2014), we found the following evidence of ping-pong in the AEP strain: (i) 86% of Hyli-bound piRNAs have an adenine at the 10th base from the



FIGURE 2. Epithelial piRNAs map predominantly to TE transcripts. Hywi- and Hyli-bound piRNAs isolated from (A) wild-type (WT) *Hydra* and (B) epithelial *Hydra* were mapped to the transcriptome. The piRNA mapping density (mapped piRNA counts per kb per million counts) is shown on the *y*-axis of the box and whisker plots. The results for piRNA mapping are subdivided by: (i) Hywi- or Hyli-bound piRNAs, (ii) sense (S) or antisense (AS) orientation, and (iii) transcript class (see legend). In all cases, TEs (red) have a significantly higher mapping density than other transcript classes as determined using Tukey's range test (*P* < 0.001; Supplemental Analysis 1).

5' end, indicating their likely identity as responder piRNAs (Supplemental Fig. S2; Ozata et al. 2019), and (ii) the presence of a 10 bp overlap between Hywi-bound and Hyli-bound piRNAs (Fig. 1C–F; Supplemental Fig. S3).

We used the piRNA mapping data to gain insights into possible targets. When mapping the piRNAs to the transcriptome, we considered both sense and antisense orientations with respect to the transcript (Supplemental Analysis 1). piRNAs mapping in an antisense orientation could indicate binding of PIWI-piRNA complexes to an RNA target, which could ultimately lead to cleavage of that target by the "slicer" activity of the PIWI domain (Saito et al. 2006; Gunawardane et al. 2007; Nishida et al. 2007). piRNAs mapping in a sense orientation could indicate that the RNA target has been directly processed into either responder piRNAs by the ping-pong mechanism or trailing piRNAs by the phasing mechanism (Brennecke et al. 2007; Han et al. 2015; Homolka et al. 2015; Iwasaki et al. 2015; Mohn et al. 2015; Ozata et al. 2019). Therefore, for antisense mapping we allowed three

> mismatches because cleavage of RNA targets can occur with imperfect base-pairing (Zhang et al. 2015), but for sense mapping, we did not allow any mismatches under the assumption that piRNAs derived directly from an RNA target should have the identical sequence. To understand piRNA mapping preferences we categorized transcripts as: (i) TE transcripts ("TE"), (ii) protein-coding transcripts with annotations ("gene"), (iii) unannotated transcripts with open reading frames of at least 100 amino acids ("uncharacterized"), and (iv) noncoding transcripts with open reading frames of less than 100 amino acids ("ncRNA") (see Materials and Methods and Supplemental Analysis 1 for further details on transcript annotations). The "uncharacterized" transcripts could include TEs that we were unable to annotate, we therefore kept these transcripts as a separate category. Similar to previous data from the 105 strain (Juliano et al. 2014; Lim et al. 2014), we found that AEP piRNAs map to TE transcripts at a higher density than non-TE transcripts (Fig. 2A; Supplemental Table S1).

> A previous study analyzing total small RNAs isolated from *Hydra vulgaris* strain 105 found evidence that sense-oriented piRNAs lie at regular intervals downstream from ping

pong cleavage sites; this is suggestive of phased piRNA biogenesis (Gainetdinov et al. 2018). In our mapping analysis of piRNAs isolated from Hydra vulgaris strain AEP we found Hywi-bound piRNAs adjacent to each other, such that the 3' ends of sense-oriented piRNAs map directly adjacent to the 5' ends of downstream mapped senseoriented piRNAs; this occurred at a higher rate than was expected by chance ($Z_0 = 5.67$, $Z_0 > 1.96$ corresponds to P < 0.05) (Supplemental Fig. S4). This indicates that Hywi protein participates in phased piRNA biogenesis and also suggests that pre-piRNAs are not significantly trimmed at the 3' end in Hydra. In contrast, we did not find a similar enrichment for Hyli-bound piRNAs ($Z_0 = 0.50$) demonstrating that Hywi is primarily responsible for phased piRNA biogenesis in Hydra (Supplemental Fig. S4). In summary, we concluded that the PIWI-piRNA pathway functions similarly in the two commonly used Hydra strains, with both pingpong and phased piRNA biogenesis occurring and with TE transcripts as the likely major target. Subsequent experiments in this study were done with Hydra vulgaris strain AEP, hereafter referred to simply as "Hydra".

Hydra piRNAs isolated from epithelial stem cells are enriched for TE sequences

To gain insight into the function of the PIWI-piRNA pathway in Hydra ESCs, we first isolated and analyzed piRNAs from epithelial Hydra, which do not contain any cell types of the interstitial lineage. To obtain epithelial Hydra, we used an established colchicine treatment protocol to remove ISCs (Campbell 1976), which leaves ESCs as the only PIWI-expressing cell type. To confirm ISC depletion in our hands, we performed colchicine treatments using a transgenic line expressing GFP under the control of the cnnos1 promoter, which drives expression in ISCs (Hemmrich et al. 2012). Animals were treated with 0.4% colchicine for 8 h followed by a 10-d recovery period. Immunoblot analysis confirmed the loss of GFP-positive ISCs (Supplemental Fig. S1E). Given that stress could lead to increased expression of TEs (Horváth et al. 2017), we assayed TE abundance in colchicine-treated animals by qPCR. We found that neither hywi expression or TE expression is increased in colchicine-treated animals, therefore TE-derived piRNAs are not artificially abundant in the epithelial Hydra used in this study (Supplemental Fig. S1F). Epithelial Hydra obtained from colchicine treatment lack the interstitial lineage but can be cultured if fed by hand and show no deficits in asexual reproduction or regeneration (Marcum and Campbell 1978). In summary, the epithelial lineages are able to return to normal function after the colchicine treatment, thereby providing a suitable system for studying PIWI-piRNA pathway function in the ESCs.

To isolate the piRNAs expressed in epithelial cells, we performed Hywi and Hyli IPs from lysates prepared from

colchicine-treated cnnos1::GFP Hydra (Fig. 1B). The isolated piRNAs were sequenced and the base distribution across the length of the piRNAs was analyzed. Similar to piRNAs isolated from WT animals, 90% of Hywi-bound piRNAs isolated from epithelial animals had a uridine at the 5' end and 79% of Hyli-bound piRNAs had an adenine at the 10th base from the 5' end (Supplemental Fig. S2). In addition, epithelial piRNAs showed the 10 bp overlap consistent with ping-pong piRNA biogenesis (Fig. 1G-J; Supplemental Fig. S3). The strongest 10 bp overlap was found between sense Hyli piRNAs and antisense Hywi piRNAs as expected. However, similar to piRNAs isolated from WT Hydra, other piRNA combinations showed some degree of 10 bp overlap. This included homotypic ping-pong, which also occurs in Drosophila, mice, and mollusks (Aravin et al. 2008; Zhang et al. 2012; Jehn et al. 2018). Overall, these data demonstrated that piRNAs expressed in the ESCs are processed by the ping-pong biogenesis pathway (Brennecke et al. 2007; Iwasaki et al. 2015). Furthermore, similar to our analysis for piRNAs isolated from Hydra that include the interstitial lineage, we found that Hywi-directed phased piRNA biogenesis occurs in ESCs ($Z_0 = 2.40$, Supplemental Fig. S4).

To identify PIWI-piRNA pathway RNA targets in ESCs, we mapped epithelial piRNAs to the Hydra vulgaris AEP stranded transcriptome (Siebert et al. 2019) using the same strategy described above for piRNAs isolated from untreated animals (Fig. 1B). Using this mapping strategy, we found that for sense and antisense mapping, both Hywi-bound and Hyli-bound piRNAs map to TE transcripts at a significantly higher density than to non-TE transcripts (Fig. 2B; Supplemental Table S1; Supplemental Analysis 1). This was likely a reflection of TE RNAs being processed into piRNAs by the ping-pong mechanism, (Fig. 1G-J) or by a Hywi-mediated phasing (Supplemental Fig. S4; Han et al. 2015; Homolka et al. 2015; Mohn et al. 2015). For piRNAs that map in an antisense orientation, 68.0% of piRNAs isolated from WT animals and 85.1% of piRNAs isolated from epithelial animals mapped with at least one mismatch (Supplemental Analysis 1). This indicated that the majority of antisense mapping piRNAs originated from a locus distinct from the target locus. For example, many likely originated from piRNA cluster loci, which have been documented in Hydra vulgaris 105 strain using the genome assembly of that strain (Lim and Kai 2015). When considering antisense mapping events to the transcriptome, most non-TE transcripts had a low mapping density. However, for both the Hywi-bound and Hylibound antisense mapping, there were groups of gene transcripts with high antisense mapping density, which could indicate the existence of gene targets (Fig. 2). Sense mapping of Hywi-bound or Hyli-bound piRNAs to gene transcripts may indicate that these transcripts are processed into either responder or trailing piRNAs (Fig. 2). In summary, these data suggested that TEs are a major

target of the PIWI–piRNA pathway in ESCs. In addition, this mapping strategy identified putative non-TE targets that we address further below.

To further test the lineage-specific functions of the PIWI-piRNA pathway, we analyzed total Hydra vulgaris AEP small RNAs that were isolated and sequenced from ectodermal, endodermal, and interstitial lineage cells sorted by FACS in our previous study (Juliano et al. 2014). We identified the piRNAs in the small RNA data sets by cross-referencing with the piRNAs sequenced from WT Hydra vulgaris AEP animals in this study (Supplemental Fig. S5A). Using this strategy, we produced a list of unique piRNA species and annotated the lineage(s) of origin for each sequence (Supplemental Fig. S5B-I). We next mapped the ectodermal and endodermal epithelial piRNAs to the Hydra transcriptome and, similar to our results using piRNAs isolated from epithelial animals, we found an enrichment for piRNAs mapping to TE transcripts as compared to other transcript categories (Supplemental Fig. S5J). In addition, this strategy allowed us to identify and map piRNAs specific to the interstitial lineage and we also found an enrichment for piRNAs mapping to TE transcripts as compared to other transcript categories (Supplemental Fig. S5K). An enrichment for transposon sequences in piRNAs expressed specifically in ISCs is expected because ISCs have germline potential.

RNAi knockdown of *hywi* in epithelial stem cells leads to the up-regulation of both TE and non-TE transcripts

Transcripts that are targeted by the PIWI-piRNA pathway should be up-regulated in response to down regulation of PIWI genes. To identify such transcripts, we used our previously validated hywi RNAi-1 transgenic line, which constitutively expresses a hywi hairpin under the control of an actin promoter that is not active in ISCs (Juliano et al. 2014). The hairpin construct also includes DsRed2, which is used to track transgenic tissue. The hairpin construct is integrated only in the interstitial lineage and is maintained through asexual propagation. In the asexually propagated hywi RNAi-1 transgenic line, the hairpin is not expressed in the PIWI-positive ISCs and therefore does not have a negative impact on the viability of the line. Integration of the hairpin construct in the interstitial lineage allows for germline propagation. After sexual reproduction, the transgene is now integrated into the genome of all cells in the offspring and the actin promoter drives expression in all epithelial cells and differentiated interstitial cells; the ESCs are the only piwi-positive cells that have RNAi transgene expression in the F1 generation. This allows us to analyze the effect of knocking down hywi in the ESCs using F1 animals. Using this strategy, we previously found that hywi knockdown in ESCs leads to death within 12 d of hatching (Juliano et al. 2014). In this study we performed RNA-seq and differential gene expression (DGE) analysis to compare gene expression between WT offspring (did not inherit the RNAi transgene) and *hywi* RNAi offspring (did inherit the transgene) 4 d after hatching. We identified 441 transcripts that were up-regulated in *hywi* RNAi animals as compared to WT siblings (Fig. 3A; Supplemental Tables S1, S2).

Given that epithelial piRNAs were enriched for TE sequences, we predicted that TEs would be up-regulated upon RNAi knockdown of hywi in ESCs. However, among the 441 transcripts up-regulated in response to hywi knockdown only 7% (33 out of 441) were annotated as TEs (Fig. 3B). To gain insights into the identity of the 238 transcripts in the gene category that were up-regulated in response to hywi knockdown (either due to direct or indirect effects), we performed GO-term analysis. We found seven terms (biological process) significantly enriched, including "defense response" and "innate immune response" (adjusted P-value <0.05; Supplemental Analysis 2). This result could suggest that TE derepression triggers an innate immune response in the affected cells, a phenomenon which has been observed in other organisms (De Cecco et al. 2019; Simon et al. 2019).

To test the reproducibility of the DGE results, we used a second *hywi* RNAi line (*hywi* RNAi-2), which targets a different portion of the *hywi* gene. We generated this transgenic line using the same strategy as the *hywi* RNAi-1 line (Juliano et al. 2014). RNA was extracted from both WT and *hywi* RNAi hatchlings produced from the *hywi* RNAi-2 parental line. qPCR analysis of these samples was used to test the expression levels of four TE transcripts and five gene transcripts identified by the DGE analysis (Fig. 3A). These transcripts were also up-regulated in the *hywi* RNA-2 line, which suggests that the results we obtained in the DGE analysis were likely due to RNAi knockdown of *hywi* and not off target effects of the *hywi* hairpins (Fig. 3C,D).

Our DGE analysis raised the possibility that the PIWI-piR-NA pathway has many non-TE targets in ESCs, but it was also possible that these were up-regulated due to secondary effects of TE up-regulation. To help discern between these two scenarios, we asked how many of the up-regulated transcripts had a high density of piRNAs mapping to them, which could indicate direct targeting by the PIWIpiRNA pathway. To identify transcripts with the highest density of mapped piRNAs, we ordered transcripts from the highest to lowest density of mapped epithelial Hywi piRNAs, and defined transcripts in the top 20% as "high mapping" (Supplemental Table S2). Of the 441 up-regulated transcripts, we found the following percentages to be "high mapping" in each category: (i) 97.0% (32 out of 33) for TE transcripts, (ii) 16.7% (six out of 36) for noncoding transcripts, (iii) 20.1% (27 out of 134) for uncharacterized transcripts, and (iv) 10.1% (24 out of 238) for gene transcripts. Therefore, we found that TE transcripts were



FIGURE 3. Both TE and non-TE transcripts are up-regulated in response to epithelial *hywi* RNAi. (*A*) DGE analysis was performed to compare transcript expression levels between *hywi* RNAi animals and WT siblings; knockdown animals carry the *hywi* RNA-1 transgene (Juliano et al. 2014). Plot showing 458 differentially expressed transcripts (red); 441 transcripts are up-regulated and 17 transcripts are down-regulated (adjusted *P*-value <0.05). (*B*) Transcript class breakdown of the 441 transcripts up-regulated in response to *hywi* knockdown. (*C*,*D*) A second RNAi transgenic line (*hywi* RNAi-2) was used to validate the DGE results (Juliano et al. 2014). qPCR was used to test the levels of select (*C*) TE transcripts and (*D*) gene transcripts identified by the DGE performed on the *hywi* RNAi-1 transgenic line. Error bars reflect standard deviation. These results indicate that at least some transcripts up-regulated in the *hywi* RNAi-1 transgenic line, suggesting the specific effect of *hywi* knockdown.

overrepresented in the "high mapping" category. In addition, we identified 24 genes that may be direct targets of the PIWI–piRNA pathway in ESCs because they were both in the "high mapping" category and were up-regulated in response to *hywi* knockdown (Supplemental Table S3).

In the *Hydra* epithelial lineages, PIWI expression is largely restricted to the body column ESCs. We therefore predicted that direct targets of the PIWI–piRNA pathway would have lower expression in the ESCs as compared to the differentiated cells of the tentacles, hypostome, and basal disc where *piwi* genes are not expressed. To determine which of our 24 putative gene targets exhibited this expression pattern, we interrogated the available singlecell expression data (Siebert et al. 2019). In these visualizations we included genes that were found to be expressed in at least 1% of the ectodermal or endodermal epithelial cells (Siebert et al. 2019). We found epithelial expression for 16 of the 24 putative gene targets: (i) thirteen gene transcripts were expressed in both the endodermal and ectodermal epithelial lineages, (ii) one gene transcript was expressed only in the ectodermal epithelial lineage, and (iii) two gene transcripts were expressed only in the endodermal epithelial lineage. No epithelial expression was found for eight of the gene transcripts, which could indicate low expression in a homeostatic animal. This could be due to transcript degradation by the PIWI-piRNA pathway or an overall low level of transcription. For the 16 gene transcripts for which we had expression data, we generated plots to visualize expression profiles in epithelial cells along the oral-aboral axis (Supplemental Analysis 5). We inspected these plots for expression patterns that could suggest targeting by the *piwi*-piRNA pathway: higher expression in the piwi-negative cells at the extremities as compared to the *piwi*-positive cells in the body column. Of the 16 transcripts, the expression pattern of t14391 was the most suggestive of Hywi-mediated repression, with higher expression at the extremities in both the ectoderm and endoderm (Supplemental Fig. S6). Swissprot BLAST analysis of this transcript suggested similarity to PARP-12. Interestingly, PARP family genes are involved in stress response, includ-

ing DNA damage repair, making this an interesting putative target for future study (Bai 2015).

PIWI-piRNA complexes cleave TE transcripts in epithelial stem cells

Our analysis of piRNA sequences isolated from ESCs strongly suggested that Hywi and Hyli repress TE RNAs by processing these RNAs into piRNAs using the pingpong mechanism. To further test this, we looked for evidence of cleaved TE RNAs using degradome sequencing, a strategy used to isolate and sequence cleaved RNAs by selecting for uncapped RNAs with a poly(A) tail (Addo-Quaye et al. 2008; German et al. 2008). To determine the RNA targets cleaved by PIWI proteins in *Hydra*, we

generated degradome sequencing libraries from both untreated and colchicine-treated cnnos1::GFP Hydra. Cleaved RNA targets of the PIWI-piRNA pathway display the ping-pong signature, which in Hydra most often manifests as a 10-bp overlap between the 5' end of antisenseoriented Hywi-bound piRNAs with the 5' end of sense-oriented Hyli-bound piRNAs and degradome fragments (Fig. 4A). We considered genes that have 10 or more of each of these species mapped and aligned in a "ping-pong" signature to be RNA targets of the PIWI-piRNA pathway. We identified 2047 such targets in WT animals and 254 such targets in epithelial animals (Supplemental Table S4). When considering the inverse of canonical pingpong, Hywi oriented sense and Hyli oriented antisense, we identified 709 targets for WT and 74 targets for epithelial animals; therefore, this did occur, but at a lower rate than canonical ping-pong targeting. In libraries produced from both whole animals and epithelial animals, TEs comprised the highest percentage of transcripts displaying this signature (Fig. 4B; Supplemental Table S4). The second highest category was uncharacterized transcripts, which may include some TE transcripts that we were unable to annotate (Supplemental Table S4). Of the 441 transcripts up-regulated in response to hywi knockdown, 46 were identified as targets in the WT degradome library (Supplemental Table S4). The majority of these (38 out of 46) were either TEs (22) or uncharacterized transcripts (16). For gene transcripts, only three of the 24 that were up-regulated in response to hywi RNAi and had a high number of piRNAs mapping, were also identified as ping-pong targets in the degradome analysis (Supplemental Table S3). However, we found that 19 of these 24 gene transcripts were in the top 5% for Hywi piRNAs mapping in the sense orientation (Supplemental Table S3). This high number of sense mapped Hywi piRNAs could indicate that these transcripts are a target of phased piRNA processing, but that we were not able to detect the ping-pong cleavage site that initiated the phased piRNA production. In summary, the degradome results further supported that a major function of the PIWI-piRNA pathway in Hydra ESCs is to cleave TE transcripts, but also revealed some possible non-TE targets for future study.

Finally, we tested whether Hywi-piRNA complexes physically associate with the somatic RNA targets identified by piRNA mapping, DGE analysis of *hywi* RNAi animals, and/or degradome sequencing. To this end, we performed RIP to isolate Hywi-piRNA-RNA ternary complexes from epithelial *Hydra* (Ilyin et al. 2017). We performed RT-PCR to test for the presence of the following transcripts in our immunoprecipitated complexes: (i) eight TE transcripts that were up-regulated in response to *hywi* knockdown, had a high number of piRNAs mapping, and were identified as targets in the degradome sequencing (Supplemental Table S3); (ii) seven gene transcripts that were up-regulated in response to *hywi* knockdown and had a



FIGURE 4. TE transcripts are cleaved by PIWI-piRNA complexes in Hydra ESCs. (A) Hywi-bound piRNAs, Hyli-bound piRNAs, and degradome reads were aligned to the transcriptome to identify likely direct targets of the PIWI-piRNA pathway. One example from the data is shown (t28405, P-element). The following pattern indicates cleavage: Antisense-oriented Hywi-bound piRNAs align with a 10-nt 5' overlap with both sense-oriented Hyli-bound piRNAs and sense-oriented degradome reads. For a transcript to be considered a target of the PIWI-piRNA pathway, we require a minimum of 10 reads of each species to map in such an arrangement. (B) The degradome sequencing from WT Hydra identified 2047 transcripts (Supplemental Table S4) as targets of the PIWI-piRNA pathway, and the transcript class distribution of these targets is shown. The degradome sequencing from epithelial Hydra identified 254 transcripts (Supplemental Table S4) as targets of the PIWI-piRNA pathway, and the class distribution of these targets is shown. In contrast to the class distribution of the transcriptome as a whole, TE transcripts and uncharacterized transcripts comprise the majority of targets in both WT and epithelial animals. (C) RNA immunoprecipitation (RIP) was used to identify transcripts that directly interact with Hywi-piRNA complexes in ESCs (for complete results see Supplemental Fig. S7). Following RIP, luciferase mRNA was added to the washed beads to act as a positive control for RNA extraction and cDNA synthesis. RT-PCR was used to test for the presence of target transcripts in complexes immunoprecipitated with a Hywi antibody. The results for four TE transcripts are shown, which were amplified in three biological replicates (Supplemental Fig. S7). The following transcripts did not associate with Hywi protein as expected: (i) The TRY3 transcript (t24687) is up-regulated in response to hywi RNAi but has a low piRNA mapping density; and (ii) the ITPA transcript (t34938) did not change in response to hywi RNAi and has a low piRNA mapping density.

high number of piRNAs mapping (one of these was also identified as a target in the degradome sequencing) (Supplemental Table S3); (iii) three gene transcripts that were up-regulated in response to hywi knockdown, but did not have a high number of piRNAs mapping; and (iv) six gene transcripts that were not affected by hywi knockdown and did not have a high number of piRNAs mapping to act as negative controls. We could reproducibly pull down four of the eight TE transcripts in all three biological replicates; the remaining four could be pulled down in at least one replicate. These data demonstrated a physical interaction between Hywi-piRNA complexes and TE RNAs (Fig. 4C; Supplemental Fig. S7; Supplemental Table S3). In contrast, we could not reproducibly pull-down transcripts from the remaining categories listed above (Supplemental Fig. S7). Three of the seven gene transcripts from the second category (t27033, t8142, and t5862) were pulled down in one or two of the three biological replicates (Supplemental Fig. S7; Supplemental Table S3). These three genes did not appear in the degradome sequencing, but do have a large number of sense mapping Hywi-bound piRNAs and therefore may be targeted by the phasing mechanism (Supplemental Table S3; Homolka et al. 2015; Pandey et al. 2017).

Conclusion

The mechanisms by which the PIWI-piRNA pathway repress TEs are best understood in the germline, but a large body of work demonstrates diverse functions for the pathway in somatic cells (for reviews, see Mani and Juliano 2013; Ross et al. 2014; Ozata et al. 2019). Here we find that the PIWI-piRNA pathway operates in the Hydra soma to degrade TE RNAs using the conserved pingpong mechanism. We demonstrated this by isolating degraded RNAs and PIWI-piRNA complexes specifically from somatic cells. We found evidence of ping-pong mediated degradation of somatic TE RNAs and association of somatic PIWI-piRNA complexes with TE RNAs. Furthermore, loss of *hywi* function led to TE up-regulation in somatic cells. Recent studies surveying a wide variety of mollusk and arthropod species found evidence for widespread somatic ping-pong as a TE repression mechanism (Jehn et al. 2018; Lewis et al. 2018). Furthermore, the *piwi* genes have widespread somatic expression in Nematostella, another cnidarian in which the pathway likely represses TEs in the soma (Praher et al. 2017). Therefore, our data in combination with these previously published studies suggest that the PIWI-piRNA pathway has ancestral roles in both germline and somatic TE repression. Furthermore, the somatic function of the pathway may have been lost in some lineages, such as in mammals, leaving germline TE repression as the most prominent function.

TE expression is up-regulated in aging metazoan somatic tissues and in aging yeast, suggesting that TE derepression is a conserved feature of cellular aging (Maxwell et al. 2011; De Cecco et al. 2013; Li et al. 2013; Van Meter et al. 2014; Patterson et al. 2015; Wood et al. 2016). Some studies have shown that repressing TEs can extend lifespan, thus understanding the mechanisms by which long-lived organisms repress TEs is of interest (Wang et al. 2011; Jones et al. 2016; Wood et al. 2016; De Cecco et al. 2019; Simon et al. 2019). A longevity study demonstrated that mortality in Hydra does not increase with age, suggesting a lack of senescence (Schaible et al. 2015), but the mechanisms by which Hydra escapes aging are not understood. It has been hypothesized that the PIWI-piRNA pathway contributes to Hydra longevity through somatic TE repression (Sturm et al. 2017). Although this is a difficult hypothesis to test, our study demonstrates that the basic premise of this hypothesis is plausible because the PIWI-piRNA pathway does repress TE expression in Hydra somatic tissue. Interestingly, Drosophila piwi is required to repress TE expression in the fat body and in intestinal stem cells and this has been linked to longevity in these somatic tissues (Jones et al. 2016; Sousa-Victor et al. 2017). For example, in aging flies, TEs are up-regulated in intestinal stem cells and overexpression of piwi prevents the age-associated decline of intestinal stem cell function (Sousa-Victor et al. 2017). It is however not clear why piwi fails to be effective in the somatic tissues of aging Drosophila, whereas hywi and hyli appear to indefinitely protect somatic cells in Hydra. Comparative studies of somatic PIWI-piRNA pathway function in animals with a variety of life history strategies could reveal new insights into the role the pathway may play in protecting somatic cells from TE-induced damage and how this may influence the rate of aging.

MATERIALS AND METHODS

Code availability

Analysis code, scripts, and Supplemental Analysis files that are needed to reproduce the analyses are available in a git repository: https://github.com/cejuliano/bteefy_piwi_transposon

1_piRNA_and_Degradome_Counts.Rmd

Generation of piRNA and degradome count files, transcript classification

2_Differential_Gene_Expression_GO_Analysis.Rmd

hywi knockdown—Differential Gene Expression analysis, GO-term enrichment

3_Ping_Pong_Phasing.Rmd

Ping-pong hit count files, ping-pong overlap frequency, ping-pong hit classification, piRNA phasing, Z-score analysis

- 4_Lineage_Sorted_piRNA_Mapping.Rmd Lineage-sorting and counting of piRNAs
- 5_Single_Cell_Data_Exploration.Rmd Assessment of homeostatic epithelial expression of putative *hywi* targets

Transgenic strains used in this study

Interstitial stem cell depletion was performed on cnnos1::GFP *Hydra* (Hemmrich et al. 2012). Differential gene expression (DGE) analysis was performed on our previously published *hywi* RNAi-1 animals (targeting bases 379–899 of the *hywi* gene) (Juliano et al. 2014). Validation was done by creating a second transgenic line using our previously published *hywi* RNAi-2 transgene (targeting bases 1557-2093 of the *hywi* gene) (Juliano et al. 2014). *Hydra* were cultured at 18°C and fed *Artemia nauplii* 1–3 times per week.

Colchicine treatment to remove the interstitial lineage

Colchicine treatment was performed as previously described (Campbell 1976). In brief, cnnos1::GFP Hydra (Hemmrich et al. 2012) were exposed to 0.4% colchicine dissolved in Hydra medium (HM) (0.3 mM CaCl₂, 0.3 mM MgSO₄, 0.5 mM NaHCO₃, 0.08 mM K2CO₃) for 8 h. After exposure, animals were rinsed with HM and transferred to fresh HM. Replicates were kept in separate dishes and HM was changed every day for 10 d after colchicine exposure while animals recovered. Ten animals were used per RIP experiment. To create degradome libraries, RNA was extracted from 10 Hydra per replicate using TRIzol (ThermoFisher Scientific) according to the manufacturer's instructions. Interstitial cell depletion was assayed using qPCR to detect levels of Cnnos1 and GFP transcripts using cDNA synthesized from 1 µg of TRIzol-extracted RNA. Interstitial cell depletion was also tested using an immunoblot to detect levels of GFP from three animals (see below for immunoblot details).

Immunoprecipitation of piRNAs with PIWI antibodies

Immunoprecipitation (IP) of PIWI proteins for piRNA isolation and sequencing was done using either 225 untreated Hydra vulgaris AEP or 225 colchicine-treated cnnos1::GFP Hydra. Hydra were centrifuged in a mini benchtop centrifuge (max speed 2000g), HM was removed, and animals were resuspended in 1 mL MCB buffer (50 mM HEPES, pH 7.5, 150 mM potassium acetate, 2 mM magnesium acetate, 10% glycerol, 0.1% Triton X-100, 0.1% NP-40, 1 mM DTT). The following were added fresh to the MCB buffer before use: (i) 1 tablet/10 mL of Pierce Protease Inhibitor Mini Tablets, EDTA-free (ThermoFisher Scientific); and (ii) 1 U/µL RNaseOUT Recombinant Ribonuclease Inhibitor (ThermoFisher Scientific). Hydra polyps in MCB buffer were homogenized on ice using a Dounce homogenizer. The resulting lysate was centrifuged at 20,000g for 10 min at 4°C and the supernatant was collected. Lysate was preincubated with 200 µL of rehydrated Protein A Sepharose CL-4B beads (GE Healthcare, Life Sciences) for 15 min at 4°C with rocking; to rehydrate beads, 100 mg of beads were washed and resuspended in 300 μL of MCB buffer. To remove beads, lysate was centrifuged at 20,000g for 10 min at 4°C and supernatant was collected. For each IP, 12 µg of purified Hywi or Hyli antibody (Juliano et al. 2014) was added to 500 µg of cleared protein lysate as measured by a NanoDrop ND-1000 Spectrophotometer 280 nM reading. The volume was then brought up to 1.2 mL per tube with MCB buffer. Next, the samples were incubated for 40 min with rocking at 4°C. Following antibody incubation, 60 µL of rehydrated beads were added to the lysate and rocked at 4°C for an additional 40 min. Beads were collected by centrifugation at 500g for 5 min at 4°C and then washed five times with 1 mL of MCB buffer. After the washes, the bead volume was brought up to 100 μ L with MCB buffer and 10 μL was removed for immunoblot blot analysis (see below for immunoblot details). To isolate piRNAs, 1 mL of TRIzol was added to the remaining beads and RNA extraction was performed according to the manufacturer's instructions. The RNA pellet was resuspended in 20 µL of DEPC-treated water (Ambion). For size analysis, 8 µL was removed for 5'-end labeling with [y-32P] ATP using polynucleotide kinase. Labeled RNAs were run on a TBE-Urea gel. The remaining 12 µL of RNA was used to prepare a sequencing library (see below for details on library preparation).

Immunoblotting

To detect GFP protein in cnnos1::GFP animals, 10 Hydra were homogenized in SDS-PAGE sample buffer for immunoblot analysis. Anti-GFP antibody was used at a 1:1000 dilution (Sigma, catalog #11814460001, Roche). All antibodies were diluted in blocking solution (3% w/v powdered milk dissolved in 0.1% TBS-Tween). For detection of Hywi and Hyli protein, 8 µg of purified anti-Hywi or anti-Hyli antibody in 50 mL of blocking solution was used for immunoblots. Anti-GFP was detected using goat antimouse IgG (H + L) Highly Cross-Adsorbed Secondary Antibody, Alexa Fluor Plus 800 (ThermoFisher Scientific, A32730). Anti-Hywi was detected using goat anti-Rabbit IgG (H + L) Superclonal Secondary Antibody, Alexa Fluor 680 (Thermofisher Scientific, A27042). Anti-Hyli was detected using goat anti-guinea pig IgG (H + L) Cross Adsorbed Secondary Antibody, DyLight 800 conjugate (ThermoFisher Scientific, SA5-10100). All secondary antibodies were diluted 1:10,000 in blocking solution. Immunoblots were imaged using the LI-COR Odyssey platform.

piRNA and degradome sequencing library preparation

Sequencing libraries were prepared from immunoprecipitated piRNAs using the NEXTflex Small RNA-seq Kit v3 (PerkinElmer cat # NOVA-5132-05). The manufacturer's protocol was followed, but with modifications to 3' and 5' adapter ligations: 3' adapter ligation was performed at 16°C overnight and 5' adapter ligation was performed at 20°C for 2 h. Following adapter ligation, libraries were amplified for 18 cycles and selected for a size of 108–180 bp using BluePippin (Sage Science). For degradome sequencing, RNA was isolated using TRIzol from 20 untreated and colchicinetreated cnnos1::GFP Hydra. Starting with 2 µg of total RNA, poly(A) enrichment was performed to capture mRNAs, which was followed by 5' adapter ligation. Ligation of the 5' adaptor was done without removing the 5' m^7G cap so that only cleaved mRNAs were available for ligation. The RNA was then chemically fragmented and 3' adapters were ligated. For ligations, 25% of the standard adapter mass was used. Libraries were prepared following the NEXTflex Small RNA-seq Kit v3 protocol. Libraries were amplified using 14 PCR cycles. Following amplification, the samples were sheared for 1 min at 94°C and size-selected for 125–200 bp using BluePippin. Degradome and piRNA libraries were pooled and sequenced using a HiSeq 4000 single end 50 bp run.

Analysis of sequenced piRNAs and degradome reads

Adapters were trimmed from piRNA and degradome sequencing reads using cutadapt (Martin 2011). After trimming, piRNAs and degradome reads were mapped to a previously published *Hydra vulgaris* AEP transcriptome (Siebert et al. 2019) using Bowtie v1.1.2 (Langmead et al. 2009). Mapping parameters were as follows: "-a –nofw -v 3 -S" was used for antisense mapping and "-a -k 40000 –norc -v 0 -S" was used for sense mapping. Degradome reads were mapped using the same parameters as sense piRNA reads. Multimappers were apportioned fractionally such that each piRNA mapping event was apportioned a single count according to the following formula:

$$C_i = \sum_{R \in R_i} \frac{1}{N}$$

where C_i is the count value for a transcript *i*, *N* is the number of times a particular piRNA maps to a unique location in the transcriptome, and R_i represents a piRNA mapping to transcript *i*.

Boxplots were generated in R using custom code (see Supplemental Analysis File "1_piRNA_and_Degradome_Counts. Rmd"). Ping-pong analysis was performed using custom scripts available in the accompanying git repository. Overlap plots, Z₁₀, and Z₀ scores were generated in R using custom code (see Supplemental Analysis file "3_Ping_Pong_Analysis.Rmd"). Mapping results were visually inspected using Integrative Genomics Viewer (IGV) (Robinson et al. 2011). The percentage of antisense piRNAs that map with mismatches was calculated using [(number of mapped piRNAs with up to three allowed mismatches) – (number of mapped piRNAs without allowed mismatches)]/(number of mapped piRNAs with up to three allowed mismatches).

Transcript annotation

Transposable element (TE): TEs were annotated in our transcriptome by using the Repbase, Swissprot, PFAM, and nr databases. First, BLAST against repeat class "Transposable element" available for Hydra vulgaris in Repbase (Bao et al. 2015) (https ://www.girinst.org/repbase/) was used to identify transcripts with TE identity (e-value cutoff of 1×10^{-5}). Next, BLAST against the Swissport database was used to add Uniprot protein descriptions to transcripts using the Uniprot Retrieve ID/mapping tool (https://www.uniprot.org/uploadlists/, cutoff e-value of 1 × 10⁻⁵). Transcripts with Uniprot protein descriptions containing the character strings "transpos", "jerky", or "mobile element" were classified as TEs. Finally, previously published PFAM annotations of our Hydra transcriptome (Siebert et al. 2019) were used to predict additional TEs. Transcripts predicted to encode domains containing "transposase", "THAP", or "_tnp_" in the domain descriptor were also classified as TEs. Gene: Transcripts with the following characteristics were placed in the "gene" category: (i) received a Swissprot or nr annotation and (ii) not classified as TE by the method described above. Uncharacterized: Transcripts with the following characteristics were placed in the "uncharacterized" category: (i) lack of Swissprot/nr/Pfam annotation, (ii) predicted ORF of 100 amino acids or greater, and (iii) not classified as a TE by the approach described above. This category contains taxonomically restricted genes. Noncoding (ncRNA): Transcripts with the following characteristics were placed in the "ncRNA" category: (i) lack of Swissprot hit, nr hit, or PFAM domain, (ii) ORF less than 100 amino acids, and (iii) not classified as a TE by the method described above. Annotation results described above are summarized in the Supplemental File "Table_S1"

GO-term enrichment analysis

GO-term enrichment analysis was performed on the 441 transcripts up-regulated in response to *hywi* RNAi-1 using GOATOOLS v0.6.10 (https://github.com/tanghaibao/goatools) (Klopfenstein et al. 2018). GO terms were considered enriched if the Bonferroni-corrected *P*-value was <0.05.

RNA immunoprecipitation (RIP) and RT-PCR

For each RIP experiment, 10 colchicine-treated cnnos1::GFP *Hydra* were used to prepare a lysate and perform an IP as described above for piRNA isolation. Prior to IP, 5% of the total cleared lysate volume was removed and saved as the "input" sample. After IP, 1 ng of Luciferase Control RNA (Promega L4561) was added to the washed beads as a positive control for subsequent RNA extraction and cDNA synthesis. RNA was extracted from the beads using TRIzol. cDNA was synthesized using M-MLV Reverse Transcriptase, RNase H Minus, Point Mutant (Promega, M3681), and random hexamers (ThermoFisher Scientific, N8080127). Three biological replicates were performed for each transcript tested (Supplemental Fig. S7). PCR was performed using primers spanning 75–150 bp (Supplemental Table S5) for each transcript using GoTaq Green Master Mix (Promega, M7122) for 35 cycles.

qRT-PCR

To test the effect of colchicine treatment on TE expression, RNA was extracted from 10 colchicine-treated cnnos1::GFP *Hydra* using TRIzol. To perform qPCR on *hywi* RNAi-2 hatchlings, RNA was extracted from 10 hatchlings using TRIzol. For all qPCR, cDNA was synthesized using M-MLV Reverse Transcriptase, RNase H Minus, Point Mutant (Promega, M3681), and random hexamers (ThermoFisher Scientific, N8080127). qRT-PCR was performed using SsoAdvanced Universal SYBR Green Supermix (BioRad,1725271) with primers spanning 75–150 bp for each transcript assayed (Supplemental Table S5). *RP49* was used for normalization. Fold change was calculated using the $\Delta\Delta C_t$ method.

Differential gene expression analysis

For DGE analysis, offspring from the *hywi* RNAi-1 transgenic line were collected 4 d post hatching. Offspring that did not inherit the transgene were considered "WT." Three WT and three *hywi* RNAi replicates were collected consisting of 10 animals each and RNA was extracted using TRIzol. Total RNA libraries were prepared using the Tru-Seq stranded RNA Kit (Illumina RS-122-2201). Libraries were sequenced on one lane of an Illumina HiSeq 2000 using a single-end 50 bp sequencing strategy. Illumina TruSeg3 Adapters were trimmed using trimmomatic (2:30:10 LEADING:3 TRAILING:3 SLIDINGWINDOW:4:15 MINLEN:36). RSEM v1.2.31 (Li and Dewey 2011), bowtie v1.1.2 (Langmead et al. 2009), and the transcriptome reference was used to estimate expression levels. Differential gene expression analysis was performed using edgeR, v 3.20.9 (Robinson et al. 2010). After expression normalization, replicates were contrasted for variance using function plotMDS. One replicate from each treatment was identified as an outlier and excluded from downstream analysis. Analysis code is available in file "2_Differential_ Gene_Expression_GO_Analysis" in the accompanying git repository.

Small RNA and piRNA filtering

Small RNA libraries used for sorting piRNAs by lineage were generated in our previous study (Juliano et al. 2014); NCBI BioProject PRJNA213706. Analysis code is available in file "4_Lineage_ Sorted_piRNA_Mapping" in the git repository.

Single cell data exploration

Single-cell sequencing data for epithelial cells from homeostatic *Hydra* were interrogated for expression of putative PIWI targets (Siebert et al. 2019). URD spline objects for endoderm and ectoderm are available from Dryad: https://doi.org/10.5061/dryad .v5r6077 (file Hydra_URD_analysis_objects). These objects contain expression data for genes that are expressed in at least 1% of the ectodermal or endodermal epithelial cells. Analysis code is available in file "5_Single_Cell_Data_Exploration" in the git repository.

DATA DEPOSITION

Raw reads from piRNA sequencing, degradome sequencing, and lineage-sorted mRNA sequencing have been submitted to the GEO repository GSE135440.

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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