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## A genome-scale shRNA resource for transgenic RNAi in *Drosophila*

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### Abstract

Existing transgenic RNAi resources in *Drosophila melanogaster* based on long double-stranded hairpin RNAs are powerful tools for functional studies, but they are ineffective in gene knockdown during oogenesis, an important model system for the study of many biological questions. We show that shRNAs, modeled on an endogenous microRNA, are extremely effective at silencing gene expression during oogenesis. We also describe our progress toward building a genome-wide shRNA resource.

Current *Drosophila* transgenic RNAi resources use long hairpins as silencing triggers<sup>1</sup>. However, for reasons unknown, long hairpins are ineffective for gene silencing in the female germline, a conclusion that we reached after extensive testing of various construct designs (Supplementary Fig. 1 and Supplementary Note 1). In *Drosophila*, RNAi can be triggered via distinct routes, each of which generates small silencing RNAs via discrete processing and loading machineries<sup>2</sup>. In particular, artificial microRNAs, referred to as shRNAs, have been shown to trigger effective silencing in somatic cells<sup>3</sup> and in the female germline in one case<sup>4</sup>. As shRNAs have not been used extensively and compared to long hairpins for their

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**Accession codes.** Gene Expression Omnibus: GSE27039 (small RNA sequences).

Note: Supplementary information is available on the Nature Methods website.

### AUTHOR CONTRIBUTIONS

J.-Q.N., R.Z. and B.C. carried out major experiments; L.-P.L., L.H., D.Y.-Z., H.-S.S., R.B., M.B. and L.A.P. produced the TRiP lines; P.K. performed the luciferase experiments in ovaries; D.H. and J.B. analyzed the piRNA pathway during oogenesis; and G.J.H. and N.P. supervised the project. R.Z., B.C., J.-Q.N., D.H., J.B., G.J.H. and N.P. wrote the manuscript.

### COMPETING FINANCIAL INTERESTS

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efficacies, we systematically evaluated their use as a transgenic trigger of RNAi in *Drosophila*.

We constructed Valium20, a vector that combines the optimized expression features of the previously reported Valium10 for somatic RNAi<sup>5</sup> with a modified scaffold of the microRNA miR-1 (Fig. 1a and Supplementary Fig. 2). Unique cloning sites allow the generation of shRNAs that accommodate the desired sequences, leading to a hairpin with perfect duplex structure, which favors shRNA loading into AGO2, the principal effector of RNAi in flies (Fig. 1a). To test the effectiveness of Valium20, we generated several fly lines containing shRNA constructs that target genes associated with either distinctive germline or maternal effect phenotypes. We induced shRNA expression specifically in the germline with *MTD-Gal4* (ref. 6), a line that carries three *Gal4* drivers expressed at various stages during oogenesis (Online Methods). For all examined lines, we recovered the expected oogenesis and maternal effect mutant phenotypes (Supplementary Note 1), indicating that shRNAs triggered potent gene knockdown during oogenesis (Fig. 1b,c). To determine whether maternal expression of shRNAs can also block expression of zygotically expressed genes, we generated shRNAs to a few zygotic genes that result in embryonic lethality when mutated. In all cases, we observed the expected phenotypes as shown for decapentaplegic (*dpp*; Fig. 1c). Finally, we tested the effectiveness of shRNAs expressed from Valium20 in somatic tissues. In general, the obtained phenotypes were stronger than those obtained with the long-hairpin-based vector Valium10 and resembled genetic null mutations for the respective genes (Fig. 1d,e and Supplementary Fig. 3).

Whereas shRNAs expressed from Valium20 generated effective knockdown phenotypes in germline and soma, the phenotypic penetrance in the germline was influenced by temperature and maternal age, indicating room for improvement (Supplementary Note 1). We therefore generated Valium22 (Supplementary Fig. 2) based on the UASp vector<sup>7</sup>, which is optimized for transgene expression in the female germline. Indeed, Valium22-mediated knockdowns in the germline were overall stronger than those generated using Valium20. We note, however, that Valium22 did not allow robust transgene expression in the soma, leading to incomplete somatic knockdowns (data not shown).

We chose the ovarian Piwi-interacting RNA (piRNA) pathway as a system to compare the efficacies and specificities of Valium20 and Valium22. Both somatic and germline cells of the *Drosophila* ovary produce piRNAs to silence transposable elements, but the pathway architecture differs in both cell types<sup>8</sup>. We generated multiple shRNA lines targeting proteins with a role in the piRNA pathway. Consistent with the strong knockdown observed for each target (Fig. 2a), RNAi phenotypes generated using the maternal *MTD-Gal4* line and Valium22 were highly reminiscent of each published null mutant. Depletion of the proteins Piwi, Aub, Spn-E or Armi resulted in complete sterility (Fig. 2b), and we observed strong derepression of three transposable elements known to be targets of the germline piRNA pathway by quantitative reverse transcription-PCR (RT-PCR) or by antibody staining (Fig. 2c,d). Loss-of-function mutations in *tudor* are known to have no impact on transposon silencing in the germline or on fertility, but these mutants are defective in Aub localization to the posterior end of the growing oocyte. This leads to a failure in the specification of primordial germ cells and results in a 'grandchild-less' phenotype. We observed all of these characteristics in the *shRNA-tud* (shRNA to the tudor gene) transgenic flies (data not shown). Typically, we observed similar phenotypes when we tested the same shRNAs with Valium20. However, Valium20-mediated knockdown of Piwi or Armi led to more severe phenotypes. This is in full agreement with data for their respective genetic null mutants, as it has been shown that the function of Piwi and Armi in the somatic support cells of the ovary is critical for germline development and therefore ovarian morphology. This indicates leaky shRNA expression from Valium20 transgenes in somatic ovarian cells (Supplementary

Table 1, Supplementary Fig. 4 and Supplementary Note 1). We suspect that a temperature-sensitive element present in the *hsp70* minimal promoter of Valium20 causes low-level expression that, in combination with basal Gal4 levels, leads to shRNA expression sufficient for gene knockdowns in the soma. Altogether, our results indicate that Valium22 is not only more effective for knockdowns in the germline but is also superior in terms of tissue specificity as the effects appear to be strictly restricted to the female germline.

It was important to understand the biogenesis requirements, processing accuracy and loading characteristics of shRNAs, which are designed to take advantage of different aspects of the mostly distinct fly miRNA and siRNA pathways to favor loading into AGO2, the major RNAi machinery in *Drosophila*. Our analysis indicated that shRNAs are processed by the sequential action of Drosha-Pasha and Dicer-1-LoqsPB complexes followed by loading into AGO1 and AGO2 proteins (Supplementary Figs. 5a,b and 6 and Supplementary Note 1). Loading into AGO2 required Dcr-2 and presumably also R2D2 and perfect base-pairing at positions 9 and 10 of the siRNA, as has been shown for endogenous siRNAs. Immunopurification experiments indicated that shRNAs are efficiently loaded into AGO2 as intended but that a fraction is also associated with AGO1. Small RNA sequencing efforts from cultured *Drosophila* cells transfected with three different Valium20 constructs showed that shRNAs are accurately processed at the intended 5' ends. Furthermore, the amount of mature shRNA strands accumulated at levels comparable to those of the most abundant cellular microRNA (miR) strands and exceed miR complementary strand (miR\*)-passenger strand levels several fold as predicted from the differences in thermodynamic stability of guide and passenger strand 5' ends.

RNAi approaches can suffer from unspecific targeting of genes exhibiting sufficient sequence complementarity to the experimental siRNA (referred to as off-target effects)<sup>9</sup>. In comparison to long-hairpin constructs, shRNA constructs are expected to be advantageous as they give rise to only two siRNA species, the guide and passenger strands (Supplementary Fig. 6 and Supplementary Note 1). We minimized off-target effects caused by extended complementarity by filtering out shRNAs whose guide or passenger strands have complementary matches of 16 nucleotides or more to the fly transcriptome. As shRNAs are partially loaded into AGO1, they could also produce off-target effects through fortuitous recognition of mRNAs via the 'seed' region leading to miRNA-like repression<sup>10</sup>. To address the problem of off-target effects experimentally, we generated several shRNA lines targeting genes that are not required for viability. Only one line out of eight tested was associated with unexpected lethality (Supplementary Table 2). This is similar to what has been observed with long hairpins<sup>5,11</sup>. Thus, off-target effects with shRNAs are still a matter of concern, and we suggest verification of obtained phenotypes using independent shRNA constructs to the same gene.

Encouraged by the remarkable silencing potency of shRNAs, we started to generate a large-scale community resource for transgenic RNAi in *Drosophila* (<http://www.flyrnai.org/TRiP-HOME.html>). As of March 2011, we have constructed over 2,900 fly stocks (an updated list is available at the Transgenic RNAi Project (TRiP) homepage), and they are available from the Bloomington *Drosophila* Stock Center. We aim to construct shRNAs to all 14,208 annotated *Drosophila* protein-coding genes (genome release 5). Toward this end, we predicted shRNA sequences for all genes using designer of small interfering RNA (DSIR)<sup>12</sup>, an algorithm trained on effective siRNAs (<http://biodev.extra.cea.fr/dsir/dsir.html>). DSIR has proven quite reliable for the prediction of shRNAs for effective knockdown in transgenic flies, and the vast majority of the shRNAs described in this paper were designed using the DSIR algorithm. We synthesized 83,256 unique shRNA oligonucleotides *in situ* on four custom glass-slide microarrays<sup>13</sup>. We amplified these as pools, and inserted them into Valium20 and Valium22. We analyzed ~160,000 individual clones per vector, and identified

accurate clones through either conventional sequencing or a two-step process involving DNA Sudoku<sup>14</sup> compression followed by Illumina sequencing. We anticipate that at least 8,000 constructs per year will become available from the TRiP for distribution to the community.

## ONLINE METHODS

### Drosophila strains

The maternal triple driver (*MTD*)-*Gal4* stock<sup>6</sup> was a gift from L. Cooley (Yale University). The stock contained homozygous insertions of three *Gal4* constructs, which together provide robust germline and maternal *Gal4* expression. The genotype was *P{COG-Gal4:VP16}; P{Gal4-nos.NGT}40; P{nos-Gal4-VP16}* (Bloomington stock 31777). *P{COG-Gal4:VP16}*<sup>7</sup> contained a promoter from the *otu* gene and the 3' untranslated region (UTR) from the *K10* gene. *Gal4:VP16* expression from this transgene was weak or absent in the germarium and robust beginning in stage-1 egg chambers. *P{nos-Gal4-VP16}* contained both the promoter and 3' UTR from the *nanos* gene<sup>15</sup> and was expressed throughout the germarium and in all stages of egg chambers, with lower expression in young egg chambers (~stages 2–6)<sup>7</sup>. *P{Gal4-nos.NGT}40* contained the *nanos* promoter and *αTub84E* 3' UTR<sup>16</sup>, and was made for maternal loading of *Gal4* to drive expression during embryogenesis.

*GMR-Gal4* and *C96-Gal4* were used to drive expression in the eye and wing, respectively, as described previously<sup>5</sup>. Their descriptions are available from FlyBase (<http://flybase.org/>). Details on the full genotype of all the lines used in this study are available on the TRiP website (<http://www.flyrnai.org/TRiP-HOME.html>).

### Phenotypic analyses

For DAPI staining, ovaries were dissected in PBS and fixed in 4% electron microscopy (EM)-grade para-formaldehyde (Electron Microscopy Sciences) diluted in PBS for 30 min. Ovaries were counterstained with DAPI (Invitrogen) for 10 min. Embryonic cuticles and wings were prepared as described previously<sup>17,18</sup>. For immunofluorescence, ovaries were dissected from 3–5-day-old flies into ice-cold PBS and subsequently fixed in 4% formaldehyde (Thermo Scientific) containing 0.15% Triton X 100 (Sigma-Aldrich), diluted in PBS, for 25 min. After three rinses with PBT (PBS with 0.3% Triton X 100) ovaries were blocked in BBX (PBS containing 0.3% Triton X 100 and 0.1% BSA) for 30 min at room temperature (20–22 °C). Ovaries were incubated with primary antibodies over night at 4 °C diluted in BBX (antibodies to Piwi, Aub and Ago3, 1:500; antibodies to Armi and I element, 1:1,000; antibodies to Tudor, 1:10; antibodies to Spn-E, 1:50). After four PBT washes secondary antibodies were incubated 5 h at room temperature diluted in BBX (1:500; Molecular Probes). Ovaries were stained with DAPI for 10 min in the second of four PBT washes. Antibodies used were: antibody to Piwi, antibody to Aub and antibody to AGO3 (ref. 19); antibody to Tudor, antibody to Spn-E<sup>20</sup>; antibody to Armi<sup>21</sup> and antibody to I element (gift from D. Finnegan; University of Edinburgh). For the sterility test, ten 3–5-day-old female flies were pre-mated with wild-type males overnight in small cages on apple juice plates with yeast paste. Apple juice plate was changed without anesthetizing flies. After 18 h at 25 °C, the flies were removed and the number of laid eggs was counted (typically ~200 eggs). Forty-eight hours later hatched and non-hatched eggs were determined.

Additional information on the phenotypic analyses of RNAi reagents is available in Supplementary Figures 7 and 8 as well as in Supplementary Table 3.

## Vector construction

For descriptions of vector construction, see Supplementary Note 2.

## $\beta$ -elimination

The chemical structure of the 3' termini of small RNAs was analyzed as described previously<sup>22</sup>. In brief, RNA from immunoprecipitates or 25  $\mu$ g of total RNA from S2 cells treated with the indicated dsRNAs (17.5  $\mu$ l total volume for each sample) was incubated at room temperature for 30 min with 5  $\mu$ l 5 $\times$  borate buffer (148 mM borax and 148 mM boric acid; pH 8.6) supplemented with 3.125  $\mu$ l freshly prepared 200 mM NaIO<sub>4</sub>. We added 5  $\mu$ l of 50% glycerol to quench nonreacted sodium periodate by incubating for an additional 15 min at room temperature. Samples were then vacuum-dried and dissolved in 60  $\mu$ l 1 $\times$  borax buffer (30 mM borax, 30 mM boric acid and 50 mM NaOH; pH 9.5).  $\beta$ -elimination was carried out by incubation for 2 h at 45 °C. RNAs were ethanol-precipitated and resolved in 1 $\times$  gel loading buffer.

## Northern blotting

Northern blotting was carried out as described previously<sup>23,24</sup>. In brief, total RNAs from knockdown cells were isolated using TRIzol (Invitrogen). We separated 30  $\mu$ g total RNAs from cultured cells (with or without  $\beta$ -elimination) or RNAs from immunoprecipitations on 15% denaturing poly-acrylamide gels and transferred to Hybond-N+ membranes (Amersham Biosciences) in 1 $\times$  TBE buffer. Small RNAs were UV-light cross-linked to the membrane and prehybridized in ULTRAhyb-Oligo buffer (Ambion) for 1 h. DNA probes complementary to the indicated strands were 5' radio-labeled and added to the hybridization buffer (hybridization for 6 h at 30 °C). Membranes were washed 4 times in 1 $\times$  SSC with 0.1% SDS at 30 °C and exposed to PhosphorImager screens (GE Healthcare) for 12–48 h. Membranes were stripped by heating in 0.2 $\times$  SSC containing 0.1% SDS in a microwave twice. Sequences of the oligonucleotide probes are listed in Supplementary Note 3.

## Immunoprecipitation

Cell extracts were prepared, evenly split and immunoprecipitated using antibodies to AGO1 (Abcam) or the Flag epitope (Sigma), respectively, as previously described<sup>23</sup>. RNAs were recovered from the immunoprecipitated samples using TRIzol and used for northern blotting.

## Transposon qPCR analysis

Total RNA was extracted from ovaries of 3–5-day-old flies using TRIzol. cDNA was prepared with random primers. qPCR was performed using Maxima SYBR Green/ROX qPCR Master mix (Fermentas). Calculation of steady-state RNA levels was calculated applying the  $2^{-\Delta\Delta C_t}$  method<sup>25</sup>. *Rp49* was used for normalization of all samples, and fold enrichments were calculated in comparison to an shRNA knockdown targeting the white gene. Fold changes in steady-state transcript levels and s.d. were calculated from three biological replicates. Primer sequences are available in Supplementary Note 3.

## Small RNA libraries

Small RNAs were cloned as described previously<sup>19</sup>. For this study, the following small RNA libraries from total RNAs were prepared: 19-nucleotide (nt) to 24-nt from S2 cells transfected with shRNA to *dlg1* (*shRNA-dlg1*); 19-nt to 24-nt from S2 cells transfected with *shRNA-N*; and 19-nt to 24-nt from S2 cells transfected with *shRNA-dpp*. For each construct,  $\sim 4 \times 10^6$  S2-NP cells were transfected with 2  $\mu$ g of Valium20-shRNA construct and 1  $\mu$ g of pMT-Gal4 plasmid. ShRNA expression in cells was induced by adding 500  $\mu$ M

CuSO<sub>4</sub> 2 d after transfection. Total RNA was isolated using TRIzol 24 h after induction. Libraries were sequenced in-house using the Illumina GA-II sequencing platform.

### Bioinformatic analysis of small RNA libraries

The analysis of small RNA libraries was performed as previously described<sup>26</sup>. Illumina reads were stripped of the 3' linker and collapsed, and the resulting small RNA sequences were matched without mismatches to the *Drosophila* release 5 genome and to the genomes of *Drosophila* C virus, Flock house virus and Cricket paralysis virus with up to three mismatches. Only reads that met these conditions were analyzed further. For annotations we used a combination of University of California Santa Cruz genome browser, miRBase and Flybase tracks for protein-coding genes, repeats or transposons, noncoding RNAs and microRNAs as well as custom tracks (for shRNAs, synthetic markers, endo-siRNAs from structured loci, miR and miR\* strands) with different priorities (annotation priority list is available upon request). For comparison of small RNA counts between libraries, reads were normalized to the same total number after bioinformatic removal of sequences matching to synthetic cloning markers or assumed degradation products of abundant cellular RNAs (rRNAs, snoRNAs and tRNAs). Heatmaps were computed by plotting the abundance and ratio of individual miR, miR\* and shRNA strands in each library.

### Construction of the shRNA library

An shRNA library representing 83,256 unique synthetic hairpins was synthesized on four custom 22K Agilent microarrays<sup>13</sup>. The library covered all 14,208 annotated genes (excluding small RNA and noncoding RNA genes) of the *Drosophila* release 5 genome with up to six shRNAs per gene (14,138 genes were covered by six hairpins, and 14,147 genes were covered by five hairpins). Hairpin constructs were based on the miR-1 backbone and essentially resembled those described above with perfect complementarity between guide and passenger strands. Additional sequence was attached on both ends for PCR amplification. In addition, to eliminate off-target effects only shRNAs that lacked sequence complementarity to annotated microRNA 'seed' sequences were considered. DNA pools from microarray chips were amplified<sup>13</sup> and cloned into Valium20 and Valium22 destination vectors. Plasmid DNA was transformed, clones were picked (160,000 individual clones per destination vector), and resulting transformants were multiplexed using DNA Sudoku<sup>14</sup> at Open Biosystems. Pools were barcoded via PCR, and amplicons were sequenced in-house using the Illumina GA-II sequencing platform. Positive clones were picked into 96-well plates and Sanger sequencing was carried out to validate correct shRNA sequences. Once shRNA clones in Valium20 and Valium22 are available, they will be openly available to the *Drosophila* community.

### Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

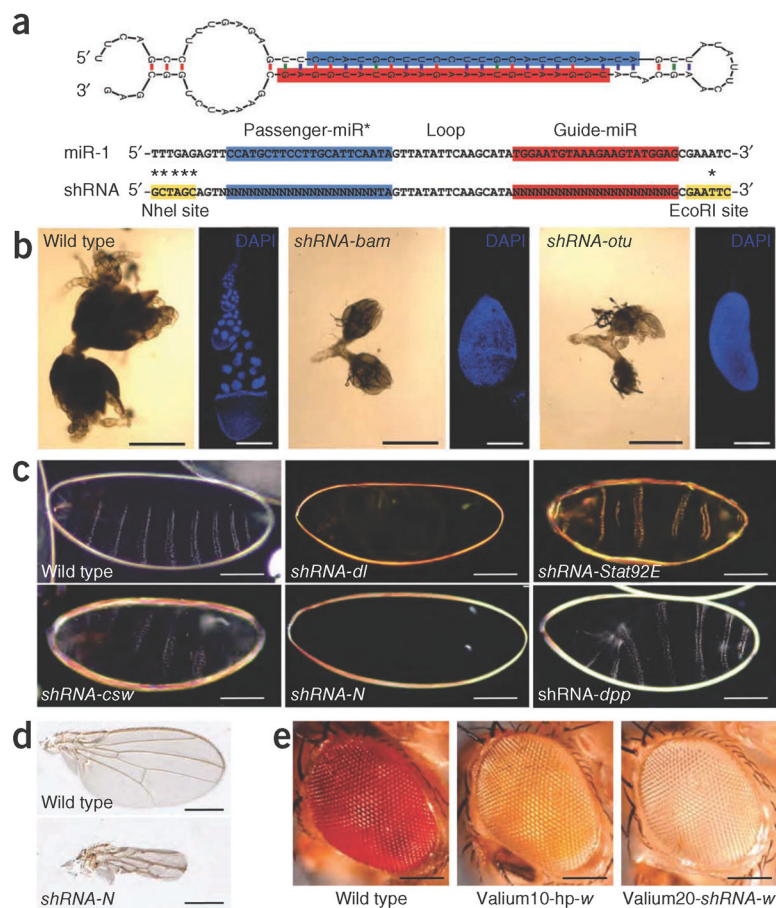
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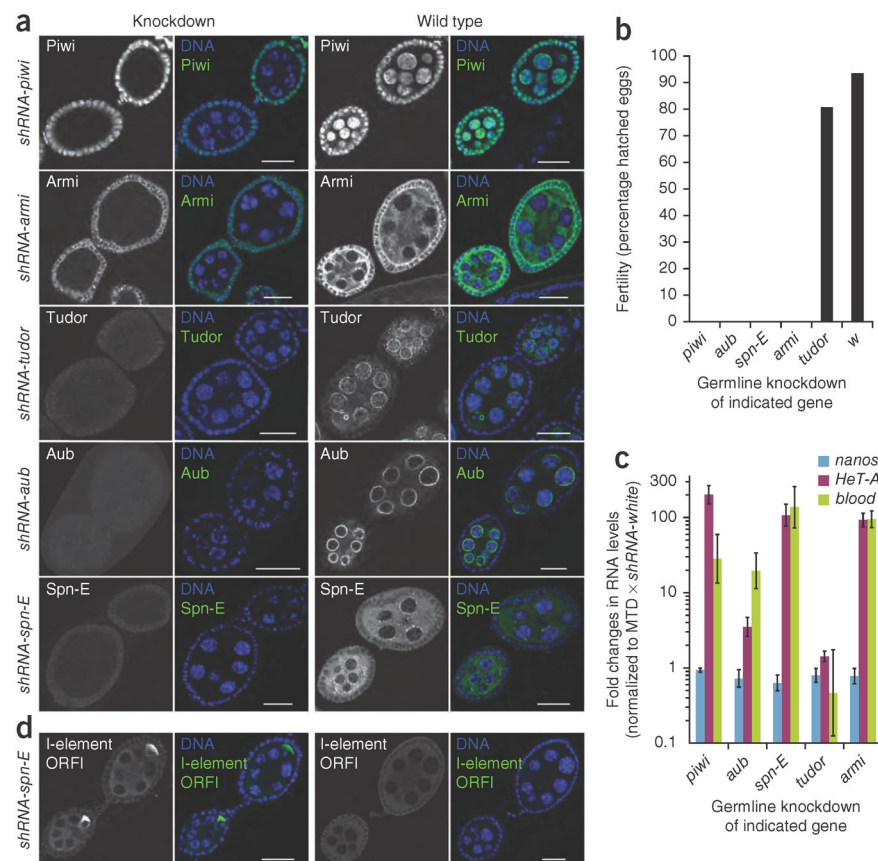
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**Figure 1.**

Design of shRNA constructs and phenotypes of shRNA-mediated gene silencing. **(a)** Structure of the *Drosophila* miR-1 and shRNA hairpins (miR-1 nucleotides replaced by the sequence of interest are indicated by N). **(b)** Phase contrast images showing ovary phenotypes associated with knockdown of bag-of-marbles (shRNA to *bam*, labeled *shRNA-bam*) and ovarian tumor (*shRNA-otu*) in *MTD-Gal4/UAS-shRNA* females (using Valium20). DAPI images show single tumorous egg chamber and wild-type egg chamber. Scale bars, 500  $\mu$ m (phase contrast) and 200  $\mu$ m (DAPI). **(c)** Dark field images of the cuticle of wild-type embryo and embryos derived from *MTD-Gal4/UAS-shRNA* females. Scale bars, 100  $\mu$ m. **(d)** Knockdown of Notch in the wing using *C96-Gal4/UAS-shRNA-N*. Scale bars, 400  $\mu$ m. **(e)** Knockdown of white using *GMR-Gal4*. In the labels, hp stands for hairpin. Scale bars, 100  $\mu$ m.



**Figure 2.**

Analysis of the piRNA pathway during oogenesis. **(a)** Immunofluorescence staining of early egg chambers showing depletion of the indicated piRNA pathway components using specific antibodies (green) upon shRNA expression via *MTD-Gal4* (using Valium22). DNA was visualized with DAPI (blue). Black and white images are of the antibody staining only. Scale bars, 20  $\mu$ m. **(b)** Fertility rates of females in which the indicated genes were knocked down with shRNAs in the germline via *MTD-Gal4* (using Valium22). For each knockdown 300–500 eggs were counted. **(c)** Fold changes in steady-state RNA levels of the transposable elements *HeT-A* and *blood* in comparison to the germline-specific *nanos* transcript upon knockdown of the indicated genes via shRNAs. The data were compared to a control sample in which the white gene was knocked down (*tp49* transcript levels were used for normalization). Data are averages of three independent biological replicates; error bars, s.d. **(d)** Immunofluorescence staining of early egg chambers with an antibody to the I-element ORF1p. Left two images are of flies expressing *shRNA-spnr-E* with *MTD-Gal4* (using Valium22); right two images are of wild-type flies. DNA was visualized with DAPI (blue).