The miR-310/13 cluster antagonizes β-catenin function in the regulation of germ and somatic cell differentiation in the *Drosophila* testis

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SUMMARY
MicroRNAs (miRNAs) are regulators of global gene expression and function in a broad range of biological processes. Recent studies have suggested that miRNAs can function as tumor suppressors or oncogenes by modulating the activities of evolutionarily conserved signaling pathways that are commonly dysregulated in cancer. We report the identification of the miR-310 to miR-313 (miR-310/13) cluster as a novel antagonist of Wingless (*Drosophila* Wnt) pathway activity in a functional screen for *Drosophila* miRNAs. We demonstrate that miR-310/13 can modulate Armadillo (Arm; *Drosophila* β-catenin) expression and activity by directly targeting the 3′-UTRs of arm and pangolin (*Drosophila* TCF) in *vivo*. Notably, the miR-310/13-deficient flies exhibit abnormal germ and somatic cell differentiation in the male gonad, which can be rescued by reducing Arm protein levels or activity. Our results implicate a previously unrecognized function for miR-310/13 in dampening the activity of Arm in early somatic and germline progenitor cells, whereby inappropriate/sustained activation of Arm-mediated signaling or cell adhesion may impact normal differentiation in the *Drosophila* male gonad.

KEY WORDS: Armadillo, Beta-catenin, Wingless, MicroRNA, Stem cells

INTRODUCTION
MicroRNAs (miRNAs) are an abundant class of small, non-coding RNAs that function as global regulators of gene expression (Bushati and Cohen, 2007). At the molecular level, they act by complementary base pairing between the 3′ untranslated region (3′-UTR) or coding sequence of target mRNAs and a ‘seed sequence’ in the miRNA (Hutvagner and Zamo, 2002; Martinez and Tuschi, 2004; Schnall-Levin et al., 2010). Mutations or misexpression of miRNAs strongly correlate with various human cancers, supporting the notion that they may function as tumor suppressors and oncogenes (Calin and Croce, 2006). miRNAs have also been shown to repress the expression of important cancer-related genes that encode key members of signaling pathways (Croce, 2009). The activity of these pathways might be particularly susceptible to miRNA control as they are subject to multiple levels of regulation leading to specific, dose-dependent phenotypes. We hypothesized that miRNAs might influence oncogenic processes by regulating the activity of crucial signaling pathways that are dysregulated in human cancers.

The evolutionarily conserved Wnt/Wingless (Wg) pathway is one of the most prevalent cancer-associated and developmentally important signaling pathways (Nusse, 2005; Polakis, 2000). A crucial effector of the Wnt pathway is β-catenin (β-cat), or *armadillo* (*arm* in *Drosophila*). Its functions are highly conserved as a nuclear transcription factor and as a key component of E-cadherin (E-cad)-mediated cell-cell adherens junctions (AJs) (Cox et al., 1996; Hülsken et al., 1994). In the absence of the Wnt signal, β-catenin (β-catenin) is targeted for ubiquitin proteasome-mediated degradation. Stimulation of the pathway by Wnt ligands activates a signal transduction cascade that inhibits DC function. Consequently, β-catenin accumulates in the cytosol and translocates into the nucleus, where it activates transcription of target genes together with the lymphoid enhancer factor (LEF)/T-cell factor (TCF) family of transcription factors, BcI9 and Pygopus (Hoffman et al., 2005).

Small variations in cellular β-catenin levels have been suggested to have a significant impact on cellular responses by modulating either transcriptional targets or cell adhesion (Goentoro and Kirschner, 2009). We speculated that the fine-tuning of β-catenin activity might be subject to miRNA regulation. Although selected miRNAs have been shown to be capable of modulating the Wnt/Wg pathway (Kennell et al., 2008; Saydam et al., 2009; Silver et al., 2007; Thatcher et al., 2008), a comprehensive assessment of miRNAs specifically targeting the nuclear β-catenin transcriptional complex has not been attempted. Here, we report a targeted screen to identify specific miRNA modulators of *arm*. We report the identification and characterization of the miR-310/13 cluster (which encompasses miR-310 to miR-313) as a novel antagonist of the Wg pathway that directly targets *arm* and/or *pangolin* (pan, or *dTCF*) 3′-UTRs. Importantly, the miR-310-13 loss-of-function (LOF) phenotype *in vivo* is consistent with increased Arm activity and reveals a novel role for miRNA-mediated regulation of Arm/Pan in the normal proliferation and differentiation of early germ and somatic progenitor cells in the *Drosophila* testis.
**MATERIALS AND METHODS**

**Cell culture and high-throughput screen (HTS)**

For the HTS, the Wg pathway was activated in *Drosophila* Clone 8 (C18) and S2R+ cells [grown as described by DasGupta et al. (DasGupta et al., 2005)] by introducing *Axin* double-stranded RNA (dsRNA), which resulted in a robust, ligand-independent activation of the Wg-responsive dTF12 reporter (DasGupta et al., 2005) (Fig. 1A). We screened a library of miRNA expression constructs [UAS-dsRED-pri-miR (Silver et al., 2007)] that consisted of 75 previously screened pri-miR constructs (Silver et al., 2007) plus 115 as yet unscreened pri-miR plasmids for their ability to suppress dTF12 activity downstream of the DC in this transcriptionally sensitized background. A total of 190 screen-ready plasmids were plated using a Janus MDT automated workstation (Perkin Elmer) in 5 µl aliquots as quadruplicates arranged in a quadrant on a set of three 384-well plates. Several quadrants of four replica wells were left empty for the addition of assay-specific controls. *Axin* dsRNA was generated using the Megascript kit (Applied Biosystems) using the following primers (5’-3’): forward TATACGACTCATAGGGAAGACAAAAGCCTGGC and reverse TAATACGACTCACTATAGGG (capital letters) TAATACGACTCACTATAGGGagaccaaacgccgcaccgctcgcc and reverse /H11032.

Cells were suspended at 20,000 cells/well for S2/S2R+ and 40,000 cells/well for C18. The dTF12-luciferase (TOP12-Ffl) reporter and Pol III-pan reporter (Promega) were co-transfected for 5 days and luciferase levels assessed using the Promega Dual-Glo kit (Promega). Cells were incubated post-transfection for 5 days and luciferase levels assessed using the Promega Dual-Glo kit (Promega).

For screen data analysis, Firefly luciferase activity values were normalized to those of *Renilla* luciferase for each replicate. Each plate contained multiple wells treated with empty vector control (pAct or pUASt) and with *arm* and *pan* dsRNA and the dynamic range was consistent with previous observations (DasGupta et al., 2005). Each screen data point was converted to a log score value using the following formula: log score (miR-_pan-UTR) = log[Nexp(X)/Nplate median]. Thus, the obtained log scores could be compared among several plates and different cell lines. The log scores were subjected to uncentered correlation metric cluster analysis using Gene Cluster 3.0 (http://bonsai.hgc.jp/~mdehoon/software/cluster/software.html) and MatLab (MathWorks).

**Drosophila stocks and genetics**

Transgenic flies were obtained from BestGene using P-element insertion of UAS-dsRED-miR-310/313 (UAS-miR-310/13). The following stocks were obtained from the Bloomington *Drosophila* Stock Center: C96-GAL4, ptc-GAL4, UAS-AxinGFP, UAS-Arm*s10, c587-GAL4, UAS-AxinGFP. UAS-RNAi lines were obtained from the Transgenic RNAi Project (TRIP) at Harvard Medical School.

The mosaic analysis with a represippable cell marker (MARC) technique (Lee and Luo, 2001) was utilized to generate *Axin* null clones overexpressing either the control UAS-GFP transgene alone or together with UAS-miR-310/13. *Axin* 5044210 FRT82 flies were obtained from Nichoals Tolwinski (Tolwinski et al., 2003). Hsflp, tub-GAL4 UAS-GFP; FRT82, tubGAL80, CD2/TM6c flies for MARCM experiments and Wg-lacZ flies were a gift from Jessica Tesiman (New York University School of Medicine). GMR-GAL4 UAS-Wg flies were obtained from Ken Cadigan (University of Michigan, Ann Arbor). *Arm* overexpression clones were generated by heat shocking flies expressing actin->STOP->GAL4 UAS-GFP and UAS-miR-310/13 and/or UAS-Arm*, hsflpMKRS/TM6c. For the MARCM experiments, larvae were heat shocked 48-72 hours after egg lay (AEL) at 38°C for 60 minutes. For the flip-out experiments, flies were heat shocked 120 hours AEL at 38°C for 20 minutes.

GFP control and miR-312 GFP sensor flies were obtained from Paul Macdonald (Reich et al., 2009). Flies expressing a short/long mir-310-313 encompassing fragment inserted in the attP1 sites were obtained from Eric Lai (Memorial Sloan-Kettering Cancer Center, New York). UAS-Arm RNAi, UAS-E-cad RNAi and UAS-Pan RNAi were obtained from DRSC (TRIP). Bam-GFP flies were generated by the McKearin laboratory (Chen and McKearin, 2003). The d59 deletion stock was generated using P-element imprecise excision. Forward primer GAACCCAAATTC-ACACCTCTT and reverse primer CACCAAAGTGCACAGATTGA were utilized in diagnostic PCR to validate the 1 kb deletion in d59 flies.

For fertility analysis, wild-type (WT) females (n=62) were crossed to d59 males (n=44) and eggs were collected on an agar-apple juice plate. Similar egg collection was performed with a WT cross for comparison. Flies were allowed to lay eggs overnight and 200 eggs were collected and incubated at 25°C for 24 hours for each cross. The number of hatched eggs was recorded after the 24-hour incubation, and the above steps were repeated for 7 days. A separate set of crosses was performed similarly that involving crossed d59 females (n=100) to WT males (n=50), compared with a WT cross.

**Antibodies and immunostaining**

Standard procedures were employed for immunostaining of third instar larval imaginal discs and the *Drosophila* testes. Primary antibodies were diluted in block solution as follows: rabbit anti-RFP (Chemicon) 1:350; guinea pig anti-Senseless (Nolo et al., 2001) 1:1000; mouse anti-β-galactosidase 1:50 [Developmental Studies Hybridoma Bank (DSHB)]; mouse anti-Armadillo 1:200 (DSHB); rabbit anti-cleaved caspase 3 1:200 (Cell Signaling); mouse anti-GFP 1:1000 (Invitrogen); rabbit anti-Vasa 1:1000 (gift of Ruth Lehmann, New York University School of Medicine); guinea pig anti-Tj 1:3000 (gift of Dorothea Godt, University of Toronto); and mouse anti-Eya 1:20 (DSHB). *Drosophila* testes were stained as described (Flaherty et al., 2010). Bright-field and fluorescent images were captured using a Zeiss TE2000UFS microscope and Nikon Elements software or using a Zeiss LSM 510 META confocal microscope and Zeiss LSM software versions 4.2. SP1 and Zen. Clonal area quantitation was performed using Nikon Elements and MatLab. Cell perimeter analysis in the *Drosophila* testes was performed using ImageJ (NIH).

**DNA constructs**

The SV40 promoter of psiCheck-2 (Promega) was excised by digestion with *BglII* and *Nhel* and replaced with the heat shock minimal promoter cloned by DasGupta et al. (DasGupta et al., 2005). *arm* 3’-UTR and pan 3’-UTR fragments were cloned into the polylinker site modified by the Eric Lai laboratory using the 5’ *NotI* and 3’ Xhol sites. Primers were: pan, forward ATAGAATTCGGCCGCGCAATCCAGCGCAAGATTATGGAC, reverse CGCGTGCGATGTCGGTGATGAATGAC and arm, forward ATAGAATTCGGCCGCGCATAGATGGCGGCT and reverse CGCGTGCAGATGAGGCGCAAGGATAGCA. For the cloning of hsa-miR-25 we utilized the forward CGCGGCGCATTCTCACGCTGCAAG and reverse TCTAGATGATCCTACACTGCT primers to amplify the precursor miR-25 and insert it into the NotI and Xhol sites of pcDNA3.1(+)(Invitrogen). The genomic DNA utilized was obtained from Applied Biosystems. Human MIR-92a and MIR-18 primary transcript cDNAs were obtained from Cell Bionces. The Wnt reporter for mammalian cells was STF16 (DasGupta et al., 2005).

**Western blotting**

S2R+ cells in 12-well plates were transfected with 60 ng actin-GAL4, 240 ng UAS-miR and 4 µg *Axin* dsRNA using the Effectene kit. Standard PAGE western protocols were used for cell lysis, preparation of extracts, protein quantification and gel electrophoresis. Primary antibody against Arm (N2 7A1 from DSHB) was used at 1:200. Secondary antibodies exhibiting infrared fluorescence with emission wavelengths of 700 and 800 nm were utilized for secondary detection. Detection and quantitation of band intensities were performed using Li-COR Odyssey.

**RESULTS**

A screen for miRNA modulators of Wg signaling

To identify miRNAs specifically targeting β-cat activity downstream of the DC, we used a *Drosophila* cell-based screening platform developed for the discovery of novel small-molecule inhibitors of the Wnt pathway (Fig. 1A) (see Materials and methods for a detailed screen protocol). The design of the primary screen allowed us to screen for miRNAs which when overexpressed would specifically modulate the activity of a Wnt/β-cat-responsive
Fig. 1. Identification of miR-310/13 in an RNAi-based targeted screen for miRNAs that suppress Wg pathway activity downstream of Axin. (A) The primary screen. miRNAs were tested for their ability to modulate Wg reporter (dTF12) activity in Clone 8 and S2R+ cells, where the pathway was ectopically activated by Axin dsRNA. (B) Unbiased cluster analysis of averaged log normalized scores for each screened miRNA in Clone 8 and S2R+ cells. (C) Highlighted clusters of strong inhibitors of the Wg reporter (top panel; orange box in B), including the previously reported Wg antagonist miR-8. Functional clustering of members of the miR-310/13 cluster. Also highlighted are potent activators of the Wg reporter (bottom panel; red box in B), including the previously published Wg agonist miR-315. (D, D') Epistasis analyses. miR-310/13 strongly inhibits the dTF12-luciferase reporter when the Wg pathway is activated by dsRNA-mediated knockdown of Axin or by cDNA expression of ΔNLRP6 or Dsh. No significant inhibition by miR-310/13 was observed upon pathway activation with the constitutively active S37Aβ-cat. (E) Alignment of mature regions of *Drosophila* miR-310, miR-311, miR-312 and the human ortholog hsa-miR-25. (F, G) Alignments of multiple *arm* (F) and *pan* (G) 3'-UTR transcripts in various *Drosophila* and insect species revealing a conservation of miR-310/13 binding sites in *arm* and *pan* miRNA. (H) Predicted binding sites of miR-310/13 components in the 3'-UTR of *arm* and *pan*. (E-H) Gray shading highlights the conserved regions. (I, J) *arm* 3'-UTR (I) and *pan* 3'-UTR (J) containing luciferase sensor reporters are significantly downregulated in the presence of miR-310/13, compared with control sensor containing full-length cDNA of *arm* lacking the 3'-UTR (Δ3'-UTR, J). (K) Western blot (WB) showing the ability of the miR-310/13 cluster and its individual components to significantly downregulate levels of Arm protein in S2R+ cells treated with Axin dsRNA. (L) hsa-miR-25 inhibits Wnt3a-induced activation of the SuperTOPFlash/SSTF 16 reporter in human embryonic kidney (HEK293T) cells. Error bars indicate s.d. (n=4). P-values by Student’s t-test. RLU, relative luciferase units.
luciferase reporter gene in cells where the pathway is artificially activated using dsRNA-mediated knockdown of Axin (Fig. 1A).

Primary screen results consisting of control-normalized log scores in quadruplicate for each miRNA (supplementary material Tables S1, S2) were subjected to hierarchical clustering (Fig. 1B,C), which grouped miRNAs into clusters related by similar potency in modulating Wg activity (strong inhibitors in the orange box and strong activators in the red box in Fig. 1B). Corroborating screen robustness, we identified previously characterized miRNA modulators of the pathway, miR-315 (Silver et al., 2007) and miR-8 (Kennell et al., 2008), as potent activators and inhibitors, respectively, of the dTF12 reporter (Fig. 1C).

Interestingly, the Wg pathway inhibitor miR-8 functionally clustered together with miR-310, miR-311, miR-312, miR-313 (the miR-310/13 cluster), miR-92a and miR-92b (Fig. 1C). miR-310/13 and miR-92a/b are components of a highly conserved family, with orthologs in mouse (not shown) and human (Fig. 1E) (Tanzer and Stadler, 2004).

miR-310/13 targets arm and pan 3′-UTRs

The mature sequences of each individual miRNA of the miR-310/13 cluster are very similar and share identical seed sequences (Fig. 1E), which is suggestive of a set of overlapping targets. Querying the miRNA target prediction algorithms TargetScan and MIRANDA identified highly conserved binding sites for miR-310/13 in the 3′-UTRs of arm and pan (Fig. 1F-H). To test the functionality of these putative binding sites, we generated luciferase sensors containing the arm and pan 3′-UTRs. We observed a significant decrease in luciferase levels in the presence of miR-310/13 compared with the control for both constructs (Fig. 1J). Importantly, the activity of a control luciferase sensor containing arm cDNA (Arm FL) lacking the 3′-UTR and any predicted miR-310/13 binding sites was not affected by miR-310/13 misexpression (Fig. 1J'). Western blots of cells treated with Axin dsRNA showed that transfection of the miR-310/13 cluster and its individual members led to a significant decrease in Arm protein levels compared with the control (Fig. 1K).

We speculated that if miR-310/13 was capable of directly modulating Arm levels, then its clonal overexpression should resemble arm LOF. We thus generated GFP-marked flip-out (FO) clones expressing miR-310/13 in the wing imaginal disc. Clones lacking wild-type (WT) arm are difficult to recover because Arm influences both cell proliferation/growth and cell-cell AJs, and cells delaminate and die in the absence of AJs. Similar observations have been made in LOF clones for components of Wg signaling and Arm-mediated cell adhesion, including pan, arr, fz/fz2 and DE-cad (shotgun—FlyBase) (Peifer et al., 1991; Widmann and Dahmann, 2009). However, arm−/−, arr−/− or DE-cad−/− clones can be forced to survive by expressing p35, an inhibitor of apoptosis. These ‘undead’ cells delaminate and are extruded towards the basal side of the epithelium (Widmann and Dahmann, 2009).

To test whether the toxicity of miR-310/13 is a function of its inhibitory activity on Arm levels, we co-expressed a 3′-UTR-deficient form of activated Arm (ArmS10) (Pai et al., 1997) and observed a significant rescue of FO clone viability (Fig. 2C, quantification in 2D). Furthermore, we were also able to rescue UAS-miR-310/13 FO clones by co-expressing p35, similar to the observations made by Widmann and Dahmann (Widmann and Dahmann, 2009) (compare Fig. 2E with 2B). Interestingly, miR-310/13, p35 clones (Fig. 2E') appeared to be much smaller than WT clones (Fig. 2A). x-y sections of these clones revealed that they were composed of very few cells (one to three, as judged by counting nuclei; red and yellow arrowheads in Fig. 2F), and appeared to be delaminating from the epithelial plane (red and white arrowheads in Fig. 2G), suggesting defective proliferation and compromised adhesion. Consistent with the observed phenotype, early clones expressing miR-310/13, p35 displayed reduced Arm expression (Fig. 2H-H').

Together, these results suggest that the miR-310/13 cluster might directly modulate Arm and/or Pan levels by targeting the 3′-UTRs of their transcripts, thereby influencing their function in cell signaling and adhesion.

Overexpression analysis of the miR-310/13 cluster in vivo

We utilized the GAL4-UAS system to drive the expression of miR-310/13 and co-cistronic dsRED in the Drosophila leg and wing imaginal discs, and assessed its effect on modulating the endogenous response to Wg signaling (Fig. 3). In the leg disc, Wg signaling in the ventral compartment restricts dpp expression to the dorsal leg primordia, as highlighted by Dpp-lacZ staining (Fig. 3). Antagonizing Wg signaling leads to the derepression and expansion of Dpp in the ventral compartment of the disc (Estella and Mann, 2008). Consistent with its putative function as an antagonist of Wg signaling, expression of miR-310/13 under patched (ptc)-GAL4 (indicated by dsRED staining at the anteroposterior boundary of the leg disc; Fig. 3B') resulted in significant ectopic expression of Dpp-lacZ in the ventral compartment (yellow arrowhead in Fig. 3B, compare with control in 3A); 3A', additional examples are shown and quantified in supplementary material Fig. S1). Mild expansion of the endogenous expression of Dpp-lacZ was also evident in the dorsal compartment (red arrowheads in Fig. 3A',B').

Misprediction of the miR-310/13 cluster along the presumptive wing margin (dorsoventral boundary) of the wing imaginal disc strongly suppressed expression of the Wg target gene senseless (sens) (Basler and Struhl, 1994), in a manner similar to that observed for known negative regulators such as Axin (Fig. 3C-F).

Sens reduction in larval tissues correlated with loss of wing margin sensory bristles and notchig of the adult wing (Fig. 3H-I, compare with WT in 3G,G'). Co-expression of an active form of Arm (Arm*), the transcript of which lacks the endogenous 3′-UTR
(\(3^{\prime}\)UTR), with the miR-310/13 cluster rescued wing notching and the loss of sensory bristles (Fig. 3J), resulting in wings resembling those derived from flies overexpressing Arm* alone (Fig. 3K). Interestingly, closer inspection of the dorsal wing margin bristles revealed that co-expression of miR-310/13 with Arm* inhibited specification of the extra wing margin bristles observed in animals expressing Arm* alone (Fig. 3J, K). These observations suggest that miR-310/13 might rescue the Arm* overexpression phenotype by regulating endogenous \(pan\) activity (Fig. 1G-I), which is required to specify the sensory wing margin bristles by activating Sens (Olson et al., 2011).

Additionally, we used MARCM to assess whether misexpression of miR-310/13 could suppress the ectopic activation of Sens in clones of cells lacking WT Axin (supplementary material Fig. S1). \(Axin^{-/-}\) cells that express high levels of GFP, signifying robust expression of miR-310/13 (arrows in supplementary material Fig. S1K-M and ratiometric image in S1M) compared with Sens with GFP), displayed reduced levels of Sens expression compared with control.
clones not expressing miR-310/13 (arrows in supplementary material Fig. S1I,J).

These data confirm our hypothesis that the miR-310/13 cluster functions downstream of the DC, most likely at the level of nuclear transcription. Our results also indicate that misexpression of miR-310/13 in vivo can phenocopy the loss of Arm/Wg signaling activity, both in wing and leg imaginal discs.

Loss-of-function analysis of miR-310/13
To explore the physiological function of the miR-310/13 cluster in Drosophila, we generated a deletion mutant (d59) by P-element-mediated imprecise excision of a 1.1 kb fragment encompassing the

Fig. 3. Overexpression of the miR-310/13 cluster in Drosophila imaginal discs phenocopies Wg loss of function. (A–A') Dpp-lacZ expression is restricted by Wg to the dorsal compartment of WT Drosophila leg discs (yellow arrowhead in A', ns*, non-specific staining. (B–B') Misexpression of miR-310/13-dsRED using ptc-GAL4 leads to Dpp-lacZ derepression and expansion to the ventral compartment of the leg disc (yellow arrowhead in B*, quantified in supplementary material Fig. S2). Note the expression of miR-310/13 in the ptc expression domain, visualized by dsRED expression in B'. Red arrowheads (A*B') indicate mild expansion of the endogenous expression of Dpp-lacZ evident in the dorsal compartment. (C–I') Effect of misexpression of miR-310/13 on sens, a Wg target gene, and wing patterning/growth of the wing imaginal disc. Sens is expressed in two rows at the dorsoventral boundary of the wing disc and is used as a readout for Wg signaling activity (C). C96-GAL4-driven expression of Axin leads to downregulation of Sens (D, arrowhead), adult wing notching, and depletion of sensory bristles (H,H', black arrowhead), compared with the WT control (C,G,G'). Misexpression of miR-310/13-dsRED in the C96 domain (note dsRED expression in E) phenocopies Wg overexpression, both in its ability to repress Sens (F, arrowheads) and by causing wing notching in the adult (I', black arrowheads). (J,K) Co-expression of arm lacking a 3'-UTR together with miR-310/13 leads to significant rescue of wing notching and bristle loss, in addition to the formation of ectopic sensory bristles (J) similar to those observed in flies overexpressing Arm alone under C96-GAL4 (K). (G'-J',K') Magnification of dorsal wing margins from WT flies (G'), flies that co-express an active form of Arm (Arm*) and miR-310/13 (J'), and those expressing Arm* alone (K'). Note that the increase in dorsal sensory wing margin bristles in flies expressing Arm* alone (K') is rescued by the co-expression of miR-310/13 (J'). Boxes indicate the regions magnified.

pri-miR-310/13 coding region (supplementary material Fig. S2A,B). d59 homozygous (d59/d59) mutant animals were viable and lacked any conspicuous defects, similar to what has been reported by Tsurudome et al. (Tsurudome et al., 2010) for independently generated deletion mutants of the mir-310-313 locus. However, closer inspection revealed that less than 3% of over 200 counted eggs laid by d59/d59 flies hatched and developed into adults, while the majority of the eggs appeared unfertilized (data not shown). We tested whether this sterility phenotype was gender associated by measuring the hatching rate of the eggs laid following crossing of d59/d59 females or males to WT counterparts, as compared with the hatching rate of eggs from a purely WT cross.
Fig. 4. See next page for legend.
Embryos deposited by d59/d59 mothers hatched at only a slightly lower rate than WT embryos (Fig. 4B). By contrast, d59/d59 males appeared to have severe fertility defects, with significantly lower hatching rates of their progeny compared with eggs fertilized by WT males (Fig. 4A).

Since d59/d59 males displayed a sterility phenotype, we explored whether the miR-310/13 cluster was expressed in the Drosophila testis (Fig. 4D-E’). Examination of testes of miR-310/13 sensor flies expressing tubulin-GAL4-UAS-GFP flanked by hexamerized binding sites for miR-312 (Reich et al., 2009) revealed markedly lower GFP levels (Fig. 4E’E) compared with control sensor flies that express GFP lacking miR-312 binding sites (Fig. 4D,D’; quantification in supplementary material Fig. S2I-K). These results indicate that the miR-310/13 cluster, all components of which share a common seed sequence, is expressed and functional in the Drosophila testis. Importantly, this sensor did not appear to be downregulated in other tissues such as larval imaginal discs (data not shown), which could explain the lack of any discernible phenotypes in adult appendages or imaginal discs of d59/d59 animals (data not shown).

The stem cell niche of the Drosophila testis, called the hub, resides at the apical tip and secretes the self-renewal signal Unpaired to adjacent germline stem cells (GSCs) and somatic cyst stem cells (CySCs) (Kiger and Fuller, 2001; Morrison and Spradling, 2008; Tulina and Matunis, 2001). GSCs divide asymmetrically, generating another GSC that remains at the hub and a daughter gonialblast (GB) that is enveloped by two somatic cyst cells (Fig. 4C). Whereas somatic cells undergo one round of division (Lindsay and Tokuyasu, 1980), dividing GBs divide four times to form a cyst of 16 spermatogonial cells, which undergo meiosis and eventually differentiate into spermatocytes (Fig. 4C). Dividing germ cells do not undergo complete cytokinesis, but are linked by actin-rich cytokinetic structures called fusomes (Lin et al., 1994). It has been shown that the interactions between somatic cyst cells and germ cells are crucial for a proper differentiation program (DiNardo et al., 2011; Flaherty et al., 2010; Issigonis and Matunis, 2012; Leatherman and DiNardo, 2008; Lim and Fuller, 2012). Understanding the exact mechanisms of this signaling crosstalk remains an area of active study.

Consistent with the observed sterility, testes of d59/d59 males exhibited abnormal accumulations of what appeared to be large cysts of overproliferating, undifferentiated spermatogonial/germ cell clusters (germ cells are marked by vasa in Fig. 4G-G’). These clusters (within green dashed lines in Fig. 4G,G’) resided far from the hub, and the number of germ cells within each cluster far exceed (>16) those found within normally differentiating WT germline cysts (Fig. 4F’) (see the testis phenotype modeled in Fig. 6B). There was also a marked increase in the number of somatic cyst cells, as evident by an excess of cells positive for Traffic jam (Tj), a CySC/early cyst marker (Li et al., 2003), in d59/d59 testes that resided far from the hub (compare Fig. 4G’ with 4F’). A majority of these cyst cells also expressed Zn finger homeodomain 1 (Zfh1), a marker that is predominantly expressed in CySCs and at lower levels in early cyst progenitors (Fig. 4I, compare with WT control in 4H; supplementary material Fig. S2F; n=28) (Leatherman and DiNardo, 2008). Moreover, most of the Zfh1+ cells did not express the cyst differentiation marker Eyes absent (Eya) (white arrowheads in supplementary material Fig. S2F), thereby suggesting a specific expansion of early cyst cells in testis lacking miR-310/13 function. That said, some cyst cells co-expressed Zfh1 and low levels of Eya, indicating that some of these cells can also differentiate (yellow arrowheads in supplementary material Fig. S2F; n=19). Interestingly, the abnormal germ cell clusters in the d59/d59 testes were found to typically associate with at least two or more Tj+ cyst cells (yellow arrowheads in Fig. 4G’), which in many cases also expressed Zfh1 (arrowheads in Fig. 4I; supplementary material Fig. S2F; M; 22 of 28 analyzed testes). However, some of the germ cell clusters associated with the Zfh1+ cyst cells were also found to be engulfed by Eya+ cyst cells (red arrowheads in supplementary material Fig. S2F; n=19). Nonetheless, these observations suggested that the increased numbers of Zfh1+ CySC/early cyst cells could be inhibiting differentiation in the associated clusters of germ cells, thereby resulting in the accumulation of large undifferentiated germ cell cysts in d59/d59 testes.

Concordant with our hypothesis that the increased Zfh1+ cyst cells could be inhibiting differentiation in the germ lineage, the expression of the differentiation marker Bag of marbles (Bam) in the large germ cell clusters was either low or undetectable in the d59/d59 testes (arrowheads in supplementary material Fig. S2H,H’) (McKearin and Spradling, 1990). Bam protein expression in these clusters was similar to that typically observed in early, undifferentiated germ cells in WT testes (compare Bam staining in d59 mutants versus WT, supplementary material Fig. S2H,H’ versus S2G,G’) (Insco et al., 2009), suggesting that they fail to undergo normal differentiation and instead might go through an increased number of transit amplifying (TA) divisions. Indeed, EdU staining...
revealed that the germ cells in these abnormal clusters continue to proliferate synchronously (Fig. 4K, compared with WT in 4J; n=32). Moreover, 1B1 (Hu li tai shao – FlyBase) staining showed that the germ cells within each cluster are connected by a shared cytoplasm, as evidenced by the presence of large branched fusomes as opposed to the ‘dot’ fusomes characteristic of GSCs and GBs (Fig. 4L-M). Based on the condensed nuclear morphology of spermatogonial cells within the large ectopic germ cell clusters (see DAPI staining in Fig. 4G, white arrowheads, compared with the early germ cells in WT testis in 4F), the proliferation status (Fig. 4K), the presence of branched fusomes (Fig. 4M) and their inability to activate the differentiation program (supplementary material Fig. S2H,H′), we conclude that the spermatogonial cells within these clusters represent early germ cell progenitors but are distinct from GSCs/GBs. Importantly, the d59/d59 testes phenotype could be rescued by reintroducing a large genomic construct spanning the mir-310-313 region (miR-310/13L), indicating that the observed phenotype was specific to the loss of miR-310/13 function (Fig. 4N-N′).

Reduction of Wg/Arm activity rescues the mir-310/13 LOF phenotype

Although the Drosophila testis is a well-studied model system for stem cell maintenance, little is known about the contribution of Wg...
signaling to this process. Recent evidence has highlighted important functions of E-cad and Arm: GSCs require E-cad for their attachment to the hub and to ensure asymmetric cell division, thus maintaining self-renewal ability (Leatherman and DiNardo, 2008; Leatherman and DiNardo, 2010). Moreover, Arm binds E-cad at AJs and is highly expressed in the hub, which might suggest important functions for Arm-mediated cell adhesion in stem cell homeostasis (Yamashita et al., 2003).

Apart from its function in cell adhesion, Arm may also regulate Wg-dependent transcription of target genes in the testis. In support of this notion, we observed high levels of expression of an in vivo Wg transcriptional reporter Fz3RFP (Olson et al., 2011) in CySCs/early somatic cyst cells (Fig. 5A,A′, somatic lineage stained for eys). As has been reported previously, we noted that Wg appears to be specifically transcribed in somatic cells in the vicinity of the hub, and its expression gradually decreases as cells move away from the hub (Fig. 5B,B′) (Leatherman and DiNardo, 2008). Notably, the expression of the Fz3RFP reporter was markedly elevated in d59/d59 testes, especially in the somatic cyst cells that are associated with poorly differentiated germ cell clusters (yellow arrowheads in Fig. 5D,D′), compared with reporter expression in heterozygous sibling control flies (Fig. 5C,C′). Ectopic expression of the Fz3RFP reporter was also detected in d59/d59 germ cells, albeit to a much weaker extent (white arrowheads in Fig. 5D′). These data are consistent with a putative function of the miR-310/13 cluster in directly modulating or buffering Wg signaling activity in the Drosophila testis.

Next, we explored whether the miR-310/13 deletion phenotype was functionally dependent on increased activity of the Wg pathway in the Drosophila testis. Indeed, removal of one copy of each of the two Frizzled receptors (termed fzf2), a manipulation shown to inhibit Arm-mediated Wg signaling (Chen and Struhl, 1999), in d59/d59 flies largely restored normal germ cell differentiation and morphology, although the accumulation of somatic cyst cells persisted (supplementary material Fig. S3D-D″). Consistent with the morphological rescue, the fertility of these flies also appeared to be partially, but significantly, restored (supplementary material Fig. S3E).

To determine whether miR-310/13 regulation of the Wg pathway is required in the somatic or germ lineage, we expressed either cDNAs or shRNAs directed toward Wg pathway components using drivers for either lineage (Fig. 5; supplementary material Fig. S3). To address whether the d59/d59 phenotype was a consequence of elevated Arm levels, we depleted Arm in the somatic lineage either by expressing AxinGFP (Fig. 5E,E′) or an shRNA against arm (Fig. 5G,G′) under the control of C587-GAL4. Somatic overexpression of AxinGFP and RNAi-mediated knockdown of Arm led to a significant rescue of both germ cell differentiation and accumulation of Tj+ cyst cells in the d59/d59 mutant testes (Fig. 5E,G), indicating that forced reduction of Arm protein levels in the soma of miR-310/13-deleted testes can restore WT testis morphology. RNAi-mediated depletion of E-cad using C587-GAL4 also rescued the abnormal germ cell accumulations (Fig. 5I,I′), albeit to a lesser extent (compare Fig. 5E,G with 5I). Control testes expressing the aforementioned transgenic constructs alone did not display any discernible phenotypes (supplementary material Fig. S3A–C′). The expression of AxinGFP (Fig. 5F,F′) or of shRNA against arm (Fig. 5H,H′) or E-cad (Fig. 5J,J′) in d59/d59 testes under the germline-speciﬁc driver nanos-GAL4 (nos-GAL4) also resulted in a significant rescue of the accumulation of undifferentiated germ cell clusters and somatic cyst cells.

The genetic interaction data suggest that the miR-310/13 LOF phenotype might be attributable to an elevation in Arm function in nuclear signaling and cell adhesion. Our observation that loss of Arm or E-cad in either cell lineage rescued the d59/d59 phenotype supports the notion that cell-cell contacts, in addition to Arm-mediated signaling, might be important for the regulation of proliferation/differentiation in both lineages. To test this, we co-expressed activated Arm in both early somatic (Tj-GAL4) and germ cells (nos-GAL4). Strikingly, we observed that Arm expression in both lineages can result in the formation of ectopic germ cell clusters that are associated with a distinct set of Tj+ CySC/early cyst cells (Fig. 5K–K′), which is somewhat reminiscent of the phenotypes observed in miR-310/13-deleted testes. These clusters resembled an ectopic hub surrounded by germ cells (two-cell clusters to be marked with yellow dashed lines in Fig. 5K′) and Tj+ somatic cells (Fig. 5K″). However, we did not observe a specific hub structure within these ectopic clusters.

The idea that cell adhesion might impact germ and somatic cell differentiation is particularly exciting. Since somatic cells can non-autonomously affect the differentiation program of the germ cells they envelop (Leatherman and DiNardo, 2008; Lim and Fuller, 2012; Tran et al., 2000; Voog et al., 2008), germ cell differentiation may be regulated by means of modulating adhesive interactions between germ and somatic cells. Based on the observation that miR-310/13-deficient testes exhibit a striking accumulation of Zfh1+ early cyst progenitor cells, and that downregulation of components of cell adhesion and/or signaling in either lineage can at least partly rescue the phenotype, we propose that somatic support cells may non-autonomously influence germ cell differentiation by modulating both cell adhesion and signaling (see working model in Fig. 6; supplementary material Fig. S4). It is possible that increased Arm levels or activity promotes greater cell-cell adhesion between the somatic cyst cells that encapsulate the germ cells, as evidenced by robust and extensive electron-dense ‘adhesion-like’ structures observed in transmission electron microscopy images of d59/d59 versus WT testis (supplementary material Fig. S4). This in turn could prevent the germ cells from exiting the ‘extended niche’ provided by the increased number of Zfh1+ CySC/early cyst cells, which instead continue to undergo TA divisions.

To investigate the Pan-dependent signaling functions of Arm in the germ and somatic cell lineages, we expressed shRNA directed against pan using either nos-GAL4 or C587-GAL4 in d59/d59 testis (supplementary material Fig. S3M–N′). The results revealed that whereas Pan knockdown under nos-GAL4 significantly rescued germ cell differentiation (supplementary material Fig. S3N,N′), its knockdown in the somatic lineage had a minimal effect on the germ cell clustering phenotype (supplementary material Fig. S3M,M′). These data suggest a prominent function for Pan-mediated nuclear signaling of Arm in the germ lineage. The phenotypes observed in the somatic lineage might be reflective of Arm function in cell adhesion, perhaps via E-cad (supplementary material Fig. S4). These results concur with the genetic interaction we observed with fzf2, where the reduction of FzFz2 levels resulted in the rescue of germ cell differentiation but did not appear to significantly rescue the somatic cell phenotype (increased Tj+ cells in supplementary material Fig. S3D′), corroborating the hypothesis that the Pan-dependent activity of Arm might have a more prominent function in germ cells.

Taken together, these observations suggest that miR-310/13 might function to fine-tune cell signaling and adhesion activities in both lineages by modulating the levels/activity of Arm, thereby
DISCUSSION
The current paradigm regarding the conserved function of Wnts in metazoans involves a highly regulated dosage of Wnt pathway activity, with too little or too much leading to abnormal cellular changes and diseases such as cancer. Through their ability to fine-tune protein levels, miRNAs are ideal candidates to refine the gradient of the Wnt response characteristic of numerous developmental contexts (van Amerongen and Nusse, 2009). In this study, we describe the identification of the miR-310/13 cluster and its human ortholog hsa-miR-25 as a conserved antagonist of the Wnt/Wg signaling pathway. Our results suggest that the miR-310/13 cluster can modulate Wg signaling activity by directly targeting the arm and pan 3′-UTRs. We show that misexpression of miR-310/13 within clones can modulate endogenous Arm levels (Fig. 2H). We also demonstrate that in vivo overexpression of miR-310 can phenocopy Arm/Wg LOF phenotypes in wing and leg imaginal discs, and that co-expression of arm cDNA lacking its 3′-UTR can rescue the miR-310/13 misexpression phenotype.

In addition to its previously reported role in neuronal synapses at neuromuscular junctions (Tsurudome et al., 2010), we have uncovered a novel function of miR-310/13 in the regulation of germ cell differentiation.
and somatic cell homeostasis in the *Drosophila* testis. LOF phenotypes of miR-310/13 in the male gonad revealed a crucial function for miRNA-mediated regulation of Wg/Arm activity in the control of normal differentiation and/or proliferation in both somatic and germ cell lineages. Specifically, we demonstrate that: (1) Wg-lacZ is expressed in CySCs and early cyst somatic cells, thereby suggesting a function for Arm/Wg activity in the regulation of germ and cyst cell differentiation (Fig. 5B'); (2) Fz3RFP reporter expression is markedly upregulated in miR-310/13-deficient testes (Fig. 5D'); (3) there is a significant increase in the number of Zfh1+ Tj+ CySC/early cyst cells (Fig. 4G','I; supplementary material Fig. S2F) and a striking accumulation of large, poorly differentiated, early germ cell clusters in miR-310/13-deficient testes (Fig. 4G'; supplementary material Fig. S2H'); (4) the miR-310/13 LOF phenotype can be significantly rescued by inhibition of Arm levels/activity; and (5) simultaneous misexpression of Arm in both germ and somatic lineages can somewhat recapitulate the miR-310/13-deficient phenotype in the *Drosophila* testis (Fig. 5K-K'').

We note that the differences in the phenotypes of Arm overexpression in both lineages compared with miR-310/13 LOF could result from a variety of factors, including the use of lineage-specific drivers that are spatially/temporally regulated versus a complete LOF phenotype, and the changes in the levels of Arm and/or Pan (as *pan* also appears to be regulated by miR-310/13, Fig. 1) in *arm* transgenic versus *miR*-deleted testes. In spite of the differences in phenotype, the important similarity is that the ectopic germ cell clusters in Arm-expressing testes appear to be restricted to an early stage of differentiation (Fig. 5K'), similar to what we observe in the miR-310/13-deleted flies (Fig. 4G). Additionally, it has previously been shown that Bam expression levels determine the number of TA mitotic divisions, thus regulating proliferation/differentiation in the adult germ cell lineage. Bam is not expressed in two-cell cysts, but its expression increases in four- to eight-cell cysts and drastically drops after the 16-cell stage (Gönczy et al., 1997; Inesco et al., 2009). These results, together with our observation that the differentiation marker Bam is either very low or completely absent in the large germ cell clusters, their inability to exit the cell cycle and that they display compact nuclear morphology in miR-310/13-deficient testes, are consistent with the notion that ectopic increases in Arm activity can result in the failure of differentiation of germ cells beyond the two- to early four-cell cyst.

Intriguingly, the germ cell clustering phenotype in d59 mutants is somewhat reminiscent of reduced TGFβ signaling activity. The main difference, however, is that whereas Bam expression is robustly upregulated in saxophone (sax) mutant (TGFβ signaling-deficient) testis (Li et al., 2007), it was absent or undetectable in d59/d59 mutants (supplementary material Fig. S2H'). Previous reports have indicated that BMP ligands, such as Dpp and Gbb, secreted from the hub and CySCs are required to inhibit differentiation of GSCs by inhibiting Bam expression (DiNardo et al., 2011; Flaherty et al., 2010; Li et al., 2007). Moreover, once Bam expression is turned off, TGFβ signaling is no longer required for the continued proliferation of the germ cells (DiNardo et al., 2011). We evaluated the levels of phosphorylated Mad (p-Mad) as a readout for TGFβ signaling in the d59 mutant testes. We noted an increase in the number of p-Mad+ early germ cells and cyst cells (white and blue arrowheads, respectively, supplementary material Fig. S5B,B') compared with WT testes (supplementary material Fig. S5A,A'). However, p-Mad immunostaining was undetectable in the large abnormal germ cell clusters (yellow arrowheads in supplementary material Fig. S5B,B'; n=20). We postulate that the increased number of Tj+, and in many cases Zfh1+, early cyst cells is a consequence of dysregulated Arm/Wg activity, which in turn prevents differentiation of the associated germ cells (perhaps by activating TGFβ signaling in the early germ cells (DiNardo et al., 2011)). However, the sustained proliferation observed in the germ cell clusters in d59/d59 mutants is likely to be due to increased Pan-dependent Arm signaling, as judged by the marked rescue of the clustering phenotype in testis knocked down for Pan function (supplementary material Fig. S3N,N'). We postulate that Wg-mediated control of Arm levels or activity might have important functions in the proliferation and differentiation of both somatic and germ cells, and that the fine-tuning of Arm activity by miR-310/13 might serve as an important mechanism to control the robust and stereotypical program of proliferation and differentiation of germ and somatic lineage progenitors in the *Drosophila* testis.

Finally, multiple studies suggest that the cytosolic availability of β-cat can be influenced by the abundance of E-cad, which has been shown to sequester transcriptionally prone β-cat to the plasma membrane (Cox et al., 1996; Fagotto et al., 1996). In stem cell maintenance or tumor establishment, the equilibrium between β-cat-mediated adhesion and transcription is synonymous to homeostasis or cancer progression. Recent evidence from studies in embryonic stem cells (ESCs) also suggests crucial functions of cell adhesion through cadherin-catenin complexes in stem cell homeostasis (Lyaeshenko et al., 2011). We propose that miRNAs such as miR-310/13 may regulate the Wnt response by modulating not only gene transcription, but also cell-cell adhesion. Such miRNA modulation of β-cat function in cellular adhesion might represent a previously unexplored mechanism of stem cell maintenance and differentiation.

Acknowledgements

We thank Dr Chi Yun and Shauna Katz at the NYU RNAi Core Facility (supported by the NYU Cancer Institute and the Kimmel Center for Stem Cell Biology) for screening support during the primary miRNA screen; Dr Nicholas Tolwinski for the UAS-AxinGFP fly stock; Dr Norbert Perrimon for the UAS-RNAi (TRIP) lines; and Dr Alfonso Martinez-Arias for critical reading of the manuscript.

Funding

This work was partially supported by the March of Dimes [Research Grant #FY10-363] and American Cancer Society [Research Scholar Grant #RSG-11-108-01-CDD]. Work in the E.C.L. group was supported by the National Institutes of Health (NIH) [R01-GM083300]. TRIP at Harvard Medical School is funded by the NIH/National Institute of General Medical Sciences (NIGMS) [R01-GM084947]. Deposited in PMC for release after 12 months.

Competing interests statement

The authors declare no competing financial interests.

Author contributions


Supplementary material

Supplementary material available online at http://dev.biologists.orglookup/suppl?doi:10.1242/dev.092817/-/DC1

References


Fig. S1. Overexpression of miR-310/13 in wing and leg imaginal discs. Related to Fig. 2. (A-D) Quantification of Dpp-lacZ in imaginal discs in Fig. 3A” (A) and 3B” (B). Integrated fluorescence intensity (FI) quantification of ventral compartment (VC) cells (C), demarcated in A and B by the white dashed line. Integrated fluorescence quantification of dorsal compartment (DC) cells (D), demarcated in A and B by the red dashed line. (E-H) Examples of wild-type imaginal discs (E,F) and imaginal discs overexpressing the miR-310/13 cluster under ptc-GAL4 (G,H). Discs overexpressing the miR-310/13 cluster show an expansion of lacZ staining into the ventral compartment compared with wild-type discs (green arrowheads). (I, J) Ectopic expression of Sens (A) within Axin⁻/⁻ MARCM clones positively marked with GFP (yellow arrowheads in J). Note that expression of GFP is not as robust as ectopic expression of Sens because of perdurance of the GAL80 inhibitor after recombination, which prevents expression of GAL4-driven UAS-GFP. (K-M’’) Cells expressing the miR-310/13 cluster inhibit the expression of Sens in Axin⁻/⁻ MARCM clones. Note that cells expressing high levels of GFP indicate high expression levels of miR-310/13; these cells display a more robust loss of Sens compared with cells expressing lower levels of GFP (see white arrows in K-M’’). (M’’) Ratiometric image to quantify the fluorescence intensities of red:green (Sens:GFP) channels. Arrows indicate regions of high green and low red, coincident with the loss of Sens in cells expressing higher levels of miR-310/13.
Fig. S2. Generation and characterization of miR-310/13 deletion flies. Related to Fig. 4. (A) EP(2)2586 flies harboring a P-element insertion in the vicinity of mir-310-313 were isogenized and crossed to the D2-3 transposase. The d59 imprecise excision line contained a 1.1 kb deletion that did not affect any of the neighboring genes. Red arrowheads indicate primers utilized to amplify the region depicted in B. (B) A 1.5 kb region amplified from genomic DNA isolated wild-type flies, whereas d59 flies harbor a 1.1 kb deletion, also confirmed by sequencing. (C-F) d59/d59 mutant testes display an increased number of cyst cells expressing the CySCs/early cyst cell marker Zfh1 (white arrowheads in F) compared with wild-type testis (white arrows in D). Some of these cells also express low levels of the differentiation marker Eya (yellow arrowheads in F). Note that cyst cells expressing both markers are also present in wild-type testes (yellow arrowheads in D). The abnormal germ cell clusters are engulfed by two or more cyst cells that are Zfh1+ Eya− (red arrowheads in F) or Zfh1− Eya+ (white arrowheads in F). (G,H) d59/d59 testes show little, if any, expression of the differentiation marker Bam (white arrowheads in H) compared with wild-type testes (G). Note that the level of Bam staining in D59 testis (H) is similar to that observed in early germ cell progenitors in WT (white arrowheads in G). (I-K) Quantification (K) of control (I) and miR-312 sensor (J) testes described in Fig. 4D-E. (L,M) Merged images of Zfh1 expression in wild-type (shown in Fig. 4H) (L) and d59/d59 testes (shown in Fig. 4I) (M). (N,O) Merged images of EdU staining in wild-type (N) and d59/d59 (O) testes.
Fig. S3. Genetic interaction studies of d59/d59 mutant testes. Related to Fig. 5. (A-C’) Testes expressing either Axin-GFP (A,A’) or RNAi lines directed against arm (B,B’) or E-cad (C,C’) under the C587-GAL4 promoter do not show any discernable phenotypes on their own. (D-D’’) Genetic removal of a copy of each of the fz and fz2 genes in the absence of miR-310/13 and the consequent reduction of Arm levels leads to rescue of germ cell clustering and the differentiation phenotype (D’) and overall fertility (as shown in E). Note that the phenotype related to somatic cell expansion was not rescued with fz, fz2 removal (D’’). (E) Rescue of fertility in d59/d59 mutants lacking a copy of fz2. The hatching rate of embryos laid by WT flies (black), d59/d59 males crossed to WT females (white), fz/fz2+ males crossed to WT females (red) and d59/d59, fz/fz2− males crossed to WT females (yellow). All males were aged 24 hours after eclosion and the hatching rate was measured 24 hours after egg deposition. (F-K″) Rescue of germ and somatic cell accumulation phenotype in d59/d59 flies upon C587-GAL4-driven (soma-specific) expression of Axin (F-F″) or shRNA against arm (G-G″). RNAi depletion of E-cad in the soma resulted in a partial rescue (J-J″). Similar observations were made in the germline using the germline-specific nos-GAL4 driver to express Axin (H-H″), or shRNAs against arm (I-I″) or E-cad (K-K″). (L,L’) EdU+ germ cell clusters in d59/d59 mutants are associated with two or more Tj+somatic cells (note three somatic nuclei indicated by white arrowheads in L, and orthogonal view in L’). (M,M’) RNAi knockdown of Pan in the somatic lineage using C587-GAL4 in d59/d59 testes fails to significantly rescue the germ cell clustering/differentiation phenotype as evidenced by the presence of large germ cell clusters (M) and the increased number of Tj+somatic cell accumulation phenotype (M’). Note that Tj+somatic cell accumulation is not affected by Pan knockdown in the germ lineage (N’).
Fig. S4. Transmission electron microscopy images of WT (Ore\textsuperscript{R}) and d59/d59 testes. Related to Fig. 6. (A) WT testis at 3400× magnification. Germ cells are ensheathed by somatic cyst cells. Separation between individual germ cells is visible (yellow arrowheads). (B) d59/d59 testis germ cell accumulation at 4400×. Note the very tight arrangement of germ cells; separation between germ cells is not discernable. (C) Boundary between two cyst cells (labeled Soma) adjacent to a germ cell (labeled Germ) at 40,000×. Note the presence of adhesive structure between the somatic cells (black arrowhead). The boundary between germ and somatic cells is indicated by yellow arrowheads. (D) Boundary between two cyst cells (Soma) and two germ cells (Germ) in d59/d59 testes at 40,000×. Note the elongated and extensive electron-dense structure between somatic cells (black arrowheads). (E) WT germ cells at 31,000×. Separation between germ cells (Germ) is clearly visible (Y-shaped gap between three germ cells). (F) d59/d59 mutant germ cells at 19,000×. Note the clearly demarcated, electron-dense boundaries between germ cells.
Fig. S5. Phosphorylated Mad (p-Mad) levels as a readout for TGFβ signaling. p-MAD staining in WT (A,A’) and d59 mutant (B,B’) testes. d59 mutant testes show an increased number of both p-MAD+ early germ cells (white arrowheads in B) and cyst cells (blue arrowheads in B). p-MAD immunostaining is absent or undetectable in the large abnormal germ cell clusters (yellow arrowheads in B,B’; n=20).

Table S1. Primary screen data analysis of Clone 8 cells. Normalized log scores are shown for each miRNA, as assayed in four replicate experiments.

Download Table S1

Table S2. Primary screen data analysis of S2R+ cells. Normalized log scores are shown for each miRNA, as assayed in four replicate experiments.

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