



The *Drosophila* MLR COMPASS-like complex regulates *bantam* miRNA expression differentially in the context of cell fate

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ABSTRACT

The conserved MLR COMPASS-like complexes are histone modifiers that are recruited by a variety of transcription factors to enhancer regions where they act as necessary epigenetic tools for enhancer establishment and function. A critical *in vivo* target of the *Drosophila* MLR complex is the *bantam* miRNA that regulates cell survival and functions in feedback regulation of cellular signaling pathways during development. We determine that loss of *Drosophila* MLR complex function in developing wing and eye imaginal discs results in growth and patterning defects that are sensitive to *bantam* levels. Consistent with an essential regulatory role in modulating *bantam* transcription, the MLR complex binds to tissue-specific *bantam* enhancers and contributes to fine-tuning expression levels during larval tissue development. In wing imaginal discs, the MLR complex attenuates *bantam* enhancer activity by negatively regulating expression; whereas, in differentiating eye discs, the complex exerts either positive or negative regulatory activity on *bantam* transcription depending on cell fate. Furthermore, while the MLR complex is not required to control *bantam* levels in undifferentiated eye cells anterior to the morphogenetic furrow, it serves to prepare critical enhancer control of *bantam* transcription for later regulation upon differentiation. Our investigation into the transcriptional regulation of a single target in a developmental context has provided novel insights as to how the MLR complex contributes to the precise timing of gene expression, and how the complex functions to help orchestrate the regulatory output of conserved signaling pathways during animal development.

1. Introduction

COMPASS-like complexes (Complex of Proteins Associated with Set1) are highly-conserved chromatin modifiers responsible for methylation of the lysine 4 residue of histone 3 (H3K4), an epigenetic modification associated with active chromatin (Shilatifard, 2012). While yeast contains a single COMPASS complex, multicellular eukaryotes harbor multiple orthologous complexes specialized for specific genomic targets and methylation activity. Those containing methyltransferases KMT2C (MLL3) or KMT2D (MLL2/4) in humans and Trr in *Drosophila* are part of a branch of COMPASS-like complexes that we refer to as MLR (MLL/Trr) complexes (Fagan and Dingwall, 2019). MLR complexes are recruited to transcription enhancer regions by a variety of binding partners where they catalyze the deposition of H3K4me1, contribute to the removal of H3K27me3, and are required for recruitment of the histone acetyltransferase p300/CBP (Herz et al., 2012; Hu et al., 2013; Issaeva et al., 2007; Lai et al., 2017; Wang et al.,

2016). While canonically associated with activation, recent work from our group has identified an additional role for an MLR complex in suppressing enhancers to prevent premature activation (Zraly et al., 2020). Enhancers, also known as cis-regulatory elements, propagate transcription factor signals and control gene expression in part by forming contacts with target promoters and dramatically increasing transcription efficiency (Levine, 2010; Rickels and Shilatifard, 2018). This mechanism allows for the intricate patterns of spatiotemporal control of gene expression necessary for the normal development of most eukaryotes. Consequently, MLR complex regulation of enhancer activity is necessary for proper lineage determination, cellular differentiation, tissue patterning, and organismal development (Ang et al., 2016; Chauhan et al., 2013; Ford and Dingwall, 2015; Lee et al., 2013; Wang et al., 2016), as well as embryonic stem cell differentiation (Wang et al., 2016). The loss of MLR complex functions are causally associated with developmental disorders or lethality in multiple animal species (Andersen and Horvitz, 2007; Chauhan et al., 2012; Sedkov et al.,

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1999; Van Laarhoven et al., 2015). For example, germline mutations of both *KMT2C* and *KMT2D* are foundational to Kleefstra and Kabuki developmental disorders, respectively (Kleefstra et al., 2012; Ng et al., 2010); *KMT2C* and *KMT2D* are also two of the most frequently somatically mutated genes in a wide variety of human cancers, and identified as drivers of malignancy in some tumor types (Fagan and Dingwall, 2019; Ford and Dingwall, 2015). Previous work by our lab and others has determined that reduction of MLR complex activity in *Drosophila* larval imaginal discs results in adult organ malformation affecting tissue size and patterning. Notably, MLR complexes have been demonstrated to interact with and be necessary for proper elaboration of multiple developmental signaling pathways, and evidence suggests that alterations of Dpp/TGF- β , Hippo, and Notch signaling underlie these phenotypes (Chauhan et al., 2012, 2013; Kanda et al., 2013; Qing et al., 2014; Sedkov et al., 2003).

We sought to better understand how the MLR complex regulates critical gene enhancers and how alteration of this activity leads to disease states by examining its function in a developmental context. In *Drosophila*, the MLR complex contributes to the positive regulation of the expression of Tgf- β paracrine signaling molecule Dpp during wing development (Chauhan et al., 2013), and Trp physically interacts with Spn/SHARP for coactivating activity on Notch signaling targets (Oswald et al., 2016). The MLR complex also associates with the Hippo coregulator HCF and signaling effector Yorkie for proper Hippo pathway target gene activation (Nan et al., 2019; Oh et al., 2014). In the fly, these developmental signaling pathways are all linked by the miRNA *bantam*, which is a direct transcriptional target as well as a feedback regulator of these three pathways (Kane et al., 2018; Oh and Irvine, 2011; Shen et al., 2015; Wu et al., 2017). Defects resulting from alteration of these pathways can be enhanced or rescued through modulation of *bantam* levels, and *bantam* modulation alone can phenocopy these effects (Becam et al., 2011; Brennecke et al., 2003; Herranz et al., 2012; Hipfner et al., 2002; Nolo et al., 2006; Peng et al., 2009).

The *bantam* miRNA is generated from a ~12 kb non-coding precursor RNA (CR43334) and the *bantam* locus spans nearly 40 kb that includes multiple tissue-specific enhancers responsible for regulating proper expression levels (Oh et al., 2013; Slattery et al., 2013). As a miRNA, *bantam* operates through translational inhibition of multiple mRNAs (Brennecke et al., 2003). It is used to control cellular function during development via regulation of targets involved in cell survival, proliferation, migration, and organ growth and patterning (Becam et al., 2011; Gerlach et al., 2019; Herranz et al., 2010; Jordan-Alvarez et al., 2017; Ma et al., 2017; Qu et al., 2017; Weng and Cohen, 2015). The best-characterized role of *bantam* is inhibition of the proapoptotic gene *hid*, regulated through Hippo signaling (Brennecke et al., 2003). The Hippo effector Yorkie (Yki) transcription factor forms heterodimers with either Scalloped (Sd) or Homothorax (Hth) to directly regulate tissue-specific *bantam* expression through multiple enhancer elements that reside up to 20 kb upstream from the CR43334 transcription unit (Slattery et al., 2013). Yki recruits the MLR complex to enhancers to activate transcription of Hippo targets, linking the MLR complex to Hippo signaling and the control of cell proliferation (Oh et al., 2014; Qing et al., 2014). We therefore reasoned that the MLR complex would likely be necessary for proper *bantam* expression and that alteration of *bantam* levels might contribute to the MLR complex loss of function phenotypes.

An ancestral genetic split of the full-length MLR methyltransferase generated separate genes in the schizophora dipterans. The *Drosophila* Cmi (also known as Lpt) gene is homologous to the N-terminal portion and encodes the highly conserved zinc-finger plant homeodomains (PHD) and a high mobility group (HMG) domain (Chauhan et al., 2012). Trithorax-related (Trr) contains the SET domain associated with methyltransferase activity (Sedkov et al., 2003). Despite their split into distinct genes, *Cmi* and *ttr* are essential and both encoded proteins are core components of the *Drosophila* MLR complex (Chauhan et al., 2012). In this study we modulated the levels of *Cmi* and *Trr* in wing and eye precursor tissues, investigated *bantam*'s role in the resulting phenotypes, and assayed the function of the MLR complex on *bantam* regulation. We found that the MLR complex is bound to tissue-specific *bantam* enhancers and the complex has important functions in regulating miRNA

expression, with both positive and negative effects on *bantam* transcription based on developmental context including tissue type, stage of differentiation, and cell fate. We further demonstrate that the MLR complex has critical enhancer regulatory functions within undifferentiated eye tissue cells that are required for proper *bantam* expression upon differentiation. Our results suggest that the role of MLR complexes during organismal development is more multifaceted and nuanced than previously reported.

2. Materials and methods

2.1. *Drosophila* husbandry and stocks

All stocks and genetic crosses were reared on standard cornmeal/dextrose medium (13% dextrose, 6% yellow cornmeal, 3% yeast extract, 1% agar, 0.3% methylparaben) at 25 °C. Experimental crosses were performed at 28 °C. *UAS-Cmi-IR*, *UAS-ttr-IR*, *UAS-Cmi*, and *cmi¹/CyO* lines were previously described (Chauhan et al., 2012). The *UAS-ban-sponge* line was obtained from S. Cohen (Becam et al., 2011). The *bwe-LacZ*, *bee-LacZ*, and *bansensGFP* lines were obtained from R. Mann (Slattery et al., 2013). Fly strains obtained from the Bloomington *Drosophila* Stock Center (BDSC) and used in this report include *Ey-Gal4* (#8220), *GawB69B-Gal4* (#1774), *DE-Gal4/TM6B* (#78371), *GMR-Gal4* (#1104), *C765-Gal4* (#36523), *en-Gal4* (#30564), *UAS-bantam* (#60671), *FRT19A;ey-FLP* (#5579), *FRT19A;hs-FLP* (#5132), *hs-FLP*, *FRT42B* (#5131), and *ttr^B;FRT19A/FM7c* (#57138). These strains are described in Flybase (<http://flybase.bio.indian.a.edu>).

2.2. Imaginal disc preparation, immunofluorescence, and imaging

Wandering third instar larvae were collected and imaginal discs dissected in ice-cold PBS, fixed in 4% formaldehyde in PBS for 15–20 min, then washed in PBST (PBS + 0.1% Triton X-100) three times for 5 min each. Washed tissues were then blocked in PBSTB (PBST + 0.1% BSA) for 2 h at room temperature followed by incubation in the primary antibody diluted in PBSTB overnight at 4 °C. After incubation, tissues were washed twice in PBSTB for 5 min each, once in PBSTB + 2% NGS for 30 min, and then twice more in PBSTB for 15 min each, followed by incubation in secondary antibody diluted in PBSTB in the dark at room temperature. Tissues were washed three times in PBST for 5 min each, then mounted in ProLong Gold antifade reagent with DAPI (Invitrogen).

Primary antibodies included mouse α - β -Gal (JIE7) and mouse α -Elav (9F8A9) (Developmental Studies Hybridoma Bank/Univ. of Iowa), rabbit α -GFP (GenScript) and rabbit α -Dcp-1 (Asp216) (Cell Signaling Technologies). Guinea pig α -Cmi was generated as previously described (Chauhan et al., 2012). Primary antibodies were used at 1:1000 concentration, except α -Dcp-1 was used at 1:250 concentration. Secondary antibodies were used at 1:1000 concentration and included α -Mouse, α -Rabbit, and α -Guinea Pig IgG (H + L) conjugated to Alexafluor 488 or 568 fluorophores (Life Technologies). Compound microscopy images were captured using an Olympus BX53 microscope with a Hamamatsu ORCA Flash 4.0 LT camera. Confocal microscopy images were captured using a Zeiss LSM 880 Airyscan and processed using Zeiss Zen® software. Quantification of GFP signal mean fluorescence intensity was assayed using Fiji ImageJ software to measure fluorescence intensity as mean gray value of selected areas, subtracting background (Schindelin et al., 2012).

2.3. Generation of somatic clones

Somatic clones in eye discs were generated using either *eyeless-FLP* or *hs-FLP* as a source of FLP recombinase. Clones of a *ttr^B* null allele (Haelterman et al., 2014) were examined using progeny from a *ttr^B; FRT19A and FRT19A;ey-FLP* mating. Eye discs were dissected from wandering third instar larvae and prepared for staining as described above. Clones of either *ttr^B* or *cmi¹* (Chauhan et al., 2012) null alleles were also generated by a heat shock-inducible FLP recombinase. First instar larvae of the genotypes *ttr^B*,

FRT19A and *FRT19A,hs-FLP* or *cmi¹*, *FRT42B* and *hs-FLP;FRT42B* were heat-shocked for 60 min at 37 °C to induce recombination and allowed to continue development at 25 °C until the third instar larval stage. Eye discs were dissected and prepared for staining as described above. Quantification and threshold identification of Dcp-1⁺ or Cmi⁺ areas were performed using the colocalization module of the Zeiss Zen® software. The enrichment ratio of Dcp-1 staining in *Cmi* or *trr* null tissue was calculated by dividing the ratio of Dcp-1⁺,Cmi⁻/Dcp-1⁺,Cmi⁺ area within the eye pouch by the total Cmi⁻/Cmi⁺ area.

2.4. TUNEL staining

TUNEL (terminal deoxynucleotidyl transferase dUTP nick end labeling) staining accomplished by collecting fixed imaginal discs and following manufacturers protocol using the In Situ Cell Death Detection Kit, Fluorescein (Roche Diagnostics). In short, fixed tissues were incubated in TUNEL solution (90% fluorescein-dUTP label solution, 10% TdT enzyme solution) for 90 min at 37 °C in a dark humidity chamber, then washed three times in PBST for 5 min each and mounted for imaging.

2.5. Scanning electron microscopy

Adult eyes were prepared for scanning electron microscopy using critical point drying as previously described (Wolff, 2011). SEM photography was taken at 1500X magnification using a Hitachi SU3500 microscope.

2.6. Adult Wing Preparation, Mounting, and Imaging

Wings were dissected from adult animals and dehydrated in isopropyl alcohol for 20 min. After dehydration, wings were mounted in DPX mountant (Fluka). Images were captured using a Leica MZ16 microscope with Leica DFC480 camera.

2.7. ChIP-seq

Chromatin from whole animals was collected, prepared, and analyzed according to (Zraly et al., 2020).

2.8. CPRG β -galactosidase assay

Ten wing or eye imaginal discs from wandering third instar larvae of the appropriate genotype were dissected and homogenized in 100 μ L CPRG assay buffer (50 mM Na-Phosphate buffer, pH 7.2 + 1 mM MgCl₂). Stepwise dilutions of homogenate were transferred to a 96-well plate, and 40 μ L 1 mM CPRG was added to each well. Plate was incubated at 37 °C overnight. Absorbance was measured at 562 nm on a BMG Labtech POLARstar® Omega plate reader. After linearity of dilutions was confirmed, the slope of each curve from regression analysis was used to determine β -Gal activity compared to control.

2.9. miRNA extraction, cDNA synthesis and qRT-PCR

Wing discs from 50 *Drosophila* third instar larvae of the appropriate genotype were dissected and miRNA was prepared using the miRVana isolation kit (Life technologies) to enrich for small RNAs, according to manufacturer's protocols. RNA (10 ng) was reverse transcribed using Multiscribe reverse transcriptase (ThermoFisher Scientific) and TaqMan small RNA assay RT primers specific for *bantam* (Assay ID: 000331) and 2S RNA (Assay ID: 001766; control), according to manufacturer's protocols. For qRT-PCR of miRNAs, TaqMan Universal PCR Master Mix II (No UNG, ThermoFisher Scientific) and miRNA-specific TaqMan microRNA Assay was used. To enrich for long RNAs, total RNA was also isolated using the miRVana kit according to manufacturer's protocols. The RNA was DNase digested, and reverse transcribed, as previously described (Chauhan et al., 2013). The primer sequences used to amplify *bantam* precursor, CR43334 are as follows: Forward primer: 5'-GCGATGTATGCGTGTAGTTAAAG-3';

Reverse primer: 5'-CCACTTTGTCGATCGTTTCATG-3'. Real time qPCR was performed on a QuantStudio6 Flex Real Time PCR System (BioRad). The 2^{- $\Delta\Delta$ CT} method was used for quantification.

2.10. Statistical analysis

Significant difference of eye phenotype severity between genetic populations was measured using Pearson's Chi-Squared Test. Significant difference of mean fluorescence intensity in eye and wing discs was measured using Student's T-test. Significant difference of β -Gal activity as determined by CPRG assay was also measured using Student's T-test.

3. Results

3.1. The MLR complex is Necessary for suppressing *bantam* expression in the developing wing

We previously reported that modulation of Cmi levels during development leads to a variety of defects, including wing vein pattern disruptions, small and rough eyes, decreased expression of ecdysone hormone regulated genes and effects on organismal growth (Chauhan et al., 2012). In the developing wing, reduced *Cmi* function leads to retraction of the L2 and L5 longitudinal veins and posterior crossvein, while overexpression causes smaller wings with bifurcation of distal veins, as well as ectopic vein formation. These phenotypes result in part from alteration of Dpp/Tgf- β signaling (Chauhan et al., 2013). *Cmi* or *trr* knockdown in larval eye discs causes reduction in eye size, while overgrowth is observed at low penetrance with *Cmi* overexpression (Chauhan et al., 2012; Sedkov et al., 2003). These phenotypes likely reflect disruptions in gene regulatory networks that control development in response to multiple signaling pathways due to improper enhancer regulation. Indeed, the MLR complex has been implicated in regulating aspects of the Hippo and Notch pathways as well (Kanda et al., 2013; Oh et al., 2014; Qing et al., 2014).

The Tgf- β , Hippo and Notch pathways share a common transcriptional target, the *bantam* miRNA, and regulation of *bantam* expression by these pathways is necessary for proper imaginal disc formation (Attisano and Wrana, 2013; Becam et al., 2011; Boulan et al., 2013; Doumpas et al., 2013; Herranz et al., 2012; Kane et al., 2018; Li and Padgett, 2012; Martin et al., 2004; Oh and Irvine, 2011; Oh et al., 2014; Qing et al., 2014; Wu et al., 2017; Zhang et al., 2013). Given the requirement for the MLR complex in regulating targets of these pathways, we hypothesized that the *bantam* miRNA is a potential direct target of the complex. To test this, we turned to the wing imaginal disc, which broadly expresses *bantam* in a stable pattern (Brennecke et al., 2003). To reduce MLR complex activity, we used the Gal4-UAS system (Brand et al., 1994) to knockdown expression of either *Cmi* or *trr* via RNAi constructs previously validated by our group (Chauhan et al., 2012, 2013; Zraly et al., 2020). *Cmi* and *Trr* were chosen because they are core subunits necessary for function of the MLR complex (Shilatfard, 2012; Zraly et al., 2020). If *bantam* is a regulatory target of the complex, then *bantam* expression should be sensitive to a reduction in MLR activity. Unexpectedly, we observed increased transcript levels of both *bantam* miRNA and its precursor transcript lncRNA:CR43334 upon knockdown of *Cmi* or *trr* in the wing disc (*C765-Gal4 > Cmi-IR* and *C765-Gal4 > trr-IR*) compared to wild type (Fig. 1A), suggesting that *bantam* expression was negatively regulated or repressed by the MLR complex in this tissue. We sought to verify these results using a *bantam* sensor GFP (*bansensGFP*) construct that functions as a constitutively expressed inverse reporter of *bantam* activity, such that the higher the expression of *bantam* in a cell, the lower the levels of GFP (Brennecke et al., 2003). Compared to control wing discs and wild type anterior tissue, knockdown of *Cmi* or *trr* within the posterior wing compartment using the *en-Gal4* driver results in decreased GFP reporter expression, reflecting increases in *bantam* activity in the disc (Fig. 1B). These data demonstrate that the MLR complex plays a necessary role in attenuating *bantam* expression in the developing wing.

As *bantam* acts as a negative feedback regulator of Dpp signaling in the wing disc (Kane et al., 2018) and we have previously suggested that

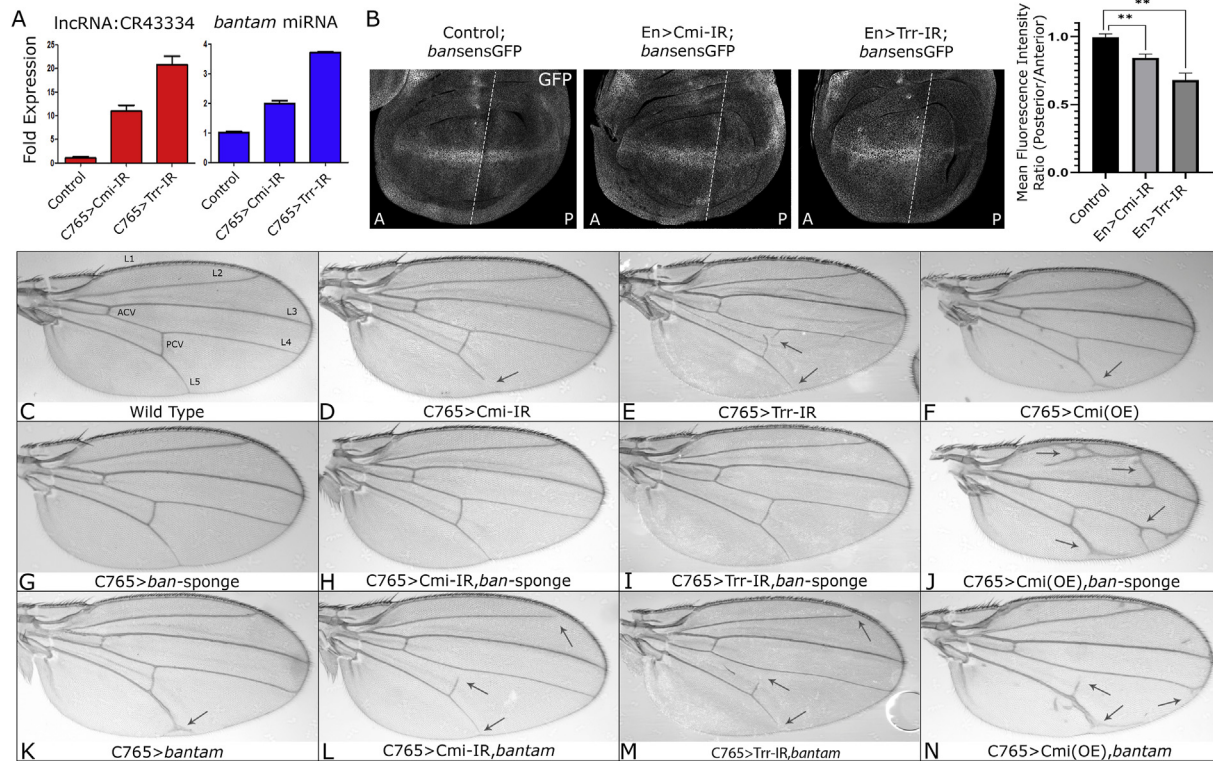


Fig. 1. The MLR complex genetically interacts with *bantam* during wing development. (A) RT-qPCR was performed on whole wing discs. Both the *bantam* precursor *IncRNA:CR43334* and processed *bantam* miRNA are upregulated upon *Cmi* or *trr* knockdown. (B) *en-Gal4* was used to knockdown *Cmi* or *trr* in the posterior wing disc; a *bantam*-GFP inverse sensor construct was used to compare *bantam* miRNA levels between posterior (P) and anterior (A) of organ. (Left) Tissue in the posterior of *en > Cmi-IR* and *en > trr-IR* organs demonstrate increased *bantam* activity (lower GFP). (Right) posterior/anterior ratio of *bantam* sensor signal quantified; ** = $p \leq 0.01$. (C–K) *C765-Gal4* was used to drive expression of genetic constructs modulating levels of *Cmi* or *bantam* in the wing disc. Compared to wild type wings (C), knockdown of *Cmi* (*Cmi-IR*) (D) or *trr* (*Trr-IR*) (E) causes retraction of wing veins, most commonly L2, L5, or PCV. (F) *Cmi* overexpression results in smaller wings and ectopic distal veins. (G) Reduction of *bantam* activity via expression of a *bantam*-specific miRNA sponge (UAS-*ban-sponge*) does not alter adult wing phenotype. (H–I) *bantam* reduction in the background of *Cmi* or *trr* knockdown rescues the wild type phenotype. (J) *bantam* reduction alongside *Cmi* overexpression enhances the phenotype, causing ectopic vein formation. (K) Overexpressing *bantam* during wing formation causes end vein forking of L5. (L–M) *bantam* overexpression in parallel with *Cmi* or *trr* knockdown enhances the *Cmi* loss-of-function phenotype. (N) Overexpression of both *Cmi* and *bantam* results in enhanced vein forking, crossvein retraction, and rescues wing size. Black arrows highlight vein formation defects.

disregulation of *Dpp* signaling is causal to wing vein phenotypes associated with modulated MLR activity (Chauhan et al., 2013), we investigated whether these phenotypes were also sensitive to *bantam* levels. Knockdown of either *Cmi* or *trr* in the developing wing results in distal vein retraction, while overexpression of *Cmi* causes vein end splitting (Chauhan et al., 2012, 2013) (Fig. 1D–F, Table 1). Genetic interaction tests between *Cmi* or *trr* and *bantam* were performed: *Cmi* or *Trr* levels were modulated in the entire wing disc via expression of transgenic shRNAi or *Cmi* overexpression constructs; in parallel, *bantam* activity was either increased through *ban* overexpression or depleted using a *bantam*-specific

miRNA sponge (*ban-sponge*) (Herranz et al., 2012) (Fig. 1C–N, Table 1). While expression of *bantam* or the *ban-sponge* in the developing wing with different *Gal4* drivers (*En-Gal4* or *MS1096*) causes alteration of adult wing size (Hipfner et al., 2002; Herranz et al., 2012), modulation of *bantam* activity driven by *C765-Gal4* has no effect on wing size (Fig. 1G,K). Simultaneous knockdown of *Cmi* or *trr* and reduction of *bantam* activity results in phenotypically wild type wings, demonstrating suppression of the short vein phenotype associated with reduced MLR complex function (Fig. 1H–I, Table 1). Conversely, overexpression of *bantam* combined with *Cmi/trr* knockdown results in greater retraction of L2 and L5 veins, thus

Table 1
Quantification of wing phenotypes.

Genotypes	N Value	WT	Distal Vein Retraction			Distal Vein Branching			Ectopic PCV	PCV Loss
			L2	L5	PCV	L4	L5	PCV		
Wild Type	121	100	0	0	0	0	0	0	0	
<i>C765 > Cmi-IR</i>	51	90	0	4	10	0	0	0	0	
<i>C765 > Cmi-IR, bantam</i>	284	86	5	6	12	0	0	0	0	
<i>C765 > Cmi-IR, bansponge</i>	133	100	0	0	0	0	0	0	0	
<i>C765 > trr-IR</i>	160	43	15	11	38	0	0	0	0	
<i>C765 > trr-IR, bantam</i>	247	19	5	12	67	0	0	0	0	
<i>C765 > trr-IR, bansponge</i>	210	59	0	0	0	9	91	5	0	
<i>C765 > Cmi(OE)</i>	128	0	0	0	0	9	91	5	0	
<i>C765 > Cmi(OE), bantam</i>	132	0	0	0	0	100	100	0	49	
<i>C765 > Cmi(OE), bansponge</i>	502	6	0	0	0	54	94	4	0	
<i>C765 > bantam</i>	80	0	0	0	0	0	100	0	0	
<i>C765 > bansponge</i>	601	72	0	0	0	0	0	28	0	

enhancing the severity of the MLR complex loss of function phenotype (Fig. 1L–M, Table 1). In the background of *Cmi* overexpression ($C765-Gal4 > Cmi(OE)$), depletion of *bantam* activity enhances the severity of vein bifurcation and ectopic vein formation (Fig. 1J, Table 1). While concurrent overexpression of *Cmi* and *bantam* does not appear to suppress the *Cmi* gain-of-function vein phenotype, wild type wing size is rescued (Fig. 1N, Table 1). The inverse phenotypic relationship between MLR activity and *bantam* expression suggests that the role of MLR in suppressing *bantam* levels may be mechanistically involved in the dose dependent *Cmi* wing phenotypes.

3.2. The MLR complex directly modulates tissue-specific *bantam* enhancers

The reduction of *bantam* expression in wing discs in response to *Cmi* or *trr* knockdown may be the result of direct or indirect regulatory mechanisms. If the MLR complex directly regulates the expression of *bantam*, it would do so through control of *bantam*-specific enhancers. Such regulatory regions have been identified as necessary and sufficient for transcription factor control of *bantam* locus expression (Oh and Irvine, 2011; Slattery et al., 2013). A *bantam* wing enhancer locus was previously identified as a cis-regulatory element residing approximately 20 kb upstream of the lncRNA:CR43334 transcription start site (Fig. 2A); an enhancer reporter of this region recapitulates the expression pattern of the

bantam miRNA in the wing imaginal discs, demonstrating tissue-specific regulatory activity (Slattery et al., 2013). To identify MLR complex-bound regions of the genome, we recently performed *Cmi* ChIP-seq at various developmental stages (Zraly et al., 2020). Examination of *Cmi* enrichments near *bantam* revealed localization throughout the locus of the precursor RNA as well as the upstream regulatory region, including peaks at the identified wing enhancer (Fig. 2A). Although *Cmi* is broadly enriched at the *bantam* locus at multiple stages of development it is only present at the wing enhancer at the wandering third instar and prepupal stages. To determine if these binding events are associated with regulatory activity by the MLR complex, we took advantage of a LacZ reporter construct created to verify the activity patterns of these enhancers (Slattery et al., 2013). Knockdown of either *Cmi* or *trr* in the wing lead to increased activity of the *bantam* wing enhancer in the cells normally expressing the miRNA (Fig. 2B,D), corresponding with the previous genetic and biochemical data in the wing (Fig. 1) and demonstrating that the MLR complex is critical for attenuating *bantam* expression during wing development through restraints on enhancer activity.

Our ChIP-seq data also revealed developmentally-associated *Cmi* enrichment at an eye disc-specific *bantam* enhancer (Fig. 2A) (Slattery et al., 2013), suggesting that the MLR complex also plays a role in *bantam* regulation in the developing eye as well. An enhancer-LacZ reporter of this region demonstrates activity at the very anterior margin of the eye

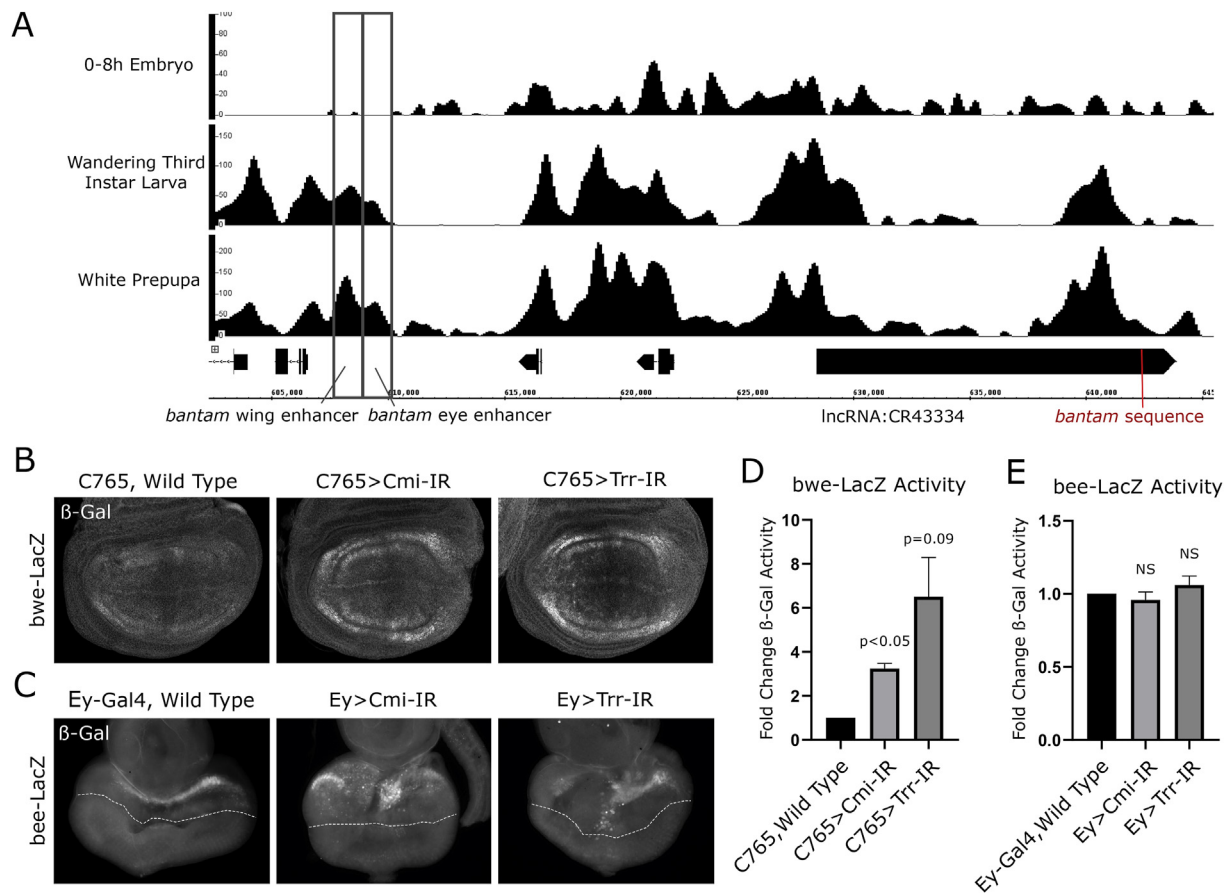


Fig. 2. The MLR complex localizes to and regulates tissue-specific *bantam* enhancers. (A) ChIP-seq was used to assay binding of MLR subunit *Cmi* across the *bantam*-containing lncRNA locus as well as at upstream tissue-specific enhancer regions. Binding profiles across different developmental timepoints including early embryo (E0-8), wandering third-instar larva (W3L), and white prepupa (WPP), demonstrate consistent localization of the complex to the lncRNA locus as well as previously identified eye- and wing-specific enhancer regions (Slattery et al., 2013) during imaginal disc development. (B–C) A *bantam* wing enhancer- β -Gal reporter (*bwe-LacZ*) or *bantam* eye enhancer- β -Gal reporter (*bee-LacZ*) was used to compare enhancer activity levels. (B) *Cmi* or *trr* was knocked down throughout the wing disc using $C765-Gal4$. $C765 > Cmi-IR$ and $C765 > trr-IR$ discs demonstrate increased *bantam* wing enhancer activity levels compared to control. (C) *Cmi* or *trr* was knocked down throughout the eye pouch of the eye disc using *ey-Gal4*. The morphogenetic furrow is marked by a dotted white line. *ey-Gal4 > Cmi-IR* and *ey-Gal4 > trr-IR* organs demonstrated disrupted *bantam* eye enhancer activity patterns compared to control. (D–E) β -Gal activity was assayed in imaginal disc homogenates using a CPRG assay. (D) $C765 > Cmi-IR$ and $C765 > trr-IR$ discs demonstrate increased *bantam* wing enhancer activity level compared to control. (E) *ey-Gal4 > Cmi-IR* and *ey-Gal4 > trr-IR* organs demonstrated similar eye enhancer activity level as control.

pouch bordering the antennal section (Fig. 2C). Unlike in the wing disc, reduction of MLR complex activity in the eye disc does not result in a clear increase or decrease of enhancer activity, but instead causes pattern disruption with sporadic sections lacking enhancer activity and apparent ectopic activation in other sections (Fig. 2C,E). These data support a hypothesis that the MLR complex also directly regulates the *bantam* eye enhancer, but its role may be in modulating enhancer activity and restricting activation to certain cells.

3.3. The MLR complex interacts with *bantam* during eye development

Enrichment at the *bantam* eye enhancer suggests that the MLR complex may be required to regulate *bantam* expression in the developing eye. To investigate this, we first turned to previously characterized eye phenotypes. Reduced MLR complex function in the *Drosophila* eye imaginal disc results in rough and shrunken adult eyes, including disorganized ommatidia (Chauhan et al., 2012; Kanda et al., 2013; Sedkov et al., 2003). If this effect is sensitive to *bantam* levels in the eye, it would suggest mechanistic interaction between the miRNA and the complex during eye development. *eyeless-Gal4* (*ey-Gal4*) was used to express *Cmi*- or *trr*-specific shRNAi constructs in the entire eye pouch of the eye-antennal imaginal disc (*ey-Gal4* > *Cmi-IR* and *ey-Gal4* > *trr-IR*), resulting in phenotypes ranging in severity from small eyes with slight roughness on the posterior margin to near-complete loss of eye tissue (Fig. 3A). SEM imaging of adult eyes of both knockdown phenotypes revealed disruptions in ommatidial patterning accompanied by bristle loss and duplication, as well as lens fusion (Fig. 3D and E). Phenotypes associated with *trr* knockdown demonstrated higher penetrance and expressivity compared to *Cmi* knockdown (Fig. 3B,D-E). This difference is consistent with the fact that the MLR complex is able to form in the absence of *Cmi*, whereas loss of *Trr* causes instability of the entire complex (Zraly et al., 2020). The variable expressivity of the *Cmi* knockdown phenotype is more sensitive, and therefore allows us to more clearly observe changes in phenotype severity due to modulation of *bantam* levels. *ey-Gal4* > *bantam* overexpression in the eye disc results in similar rough and shrunken eyes (Fig. 3B,F), and *bantam* overexpression concurrent with *Cmi* or *trr* knockdown significantly enhances the rough and shrunken phenotype (Fig. 3B,G-H). Previous studies in which *bantam* was overexpressed exclusively within the differentiating eye have reported similar disruptions of ommatidial patterning due to excess interommatidial cells (Nolo et al., 2006; Thompson and Cohen, 2006). Reduction of *bantam* activity via *ban*-sponge expression suppresses the *Cmi* knockdown-associated defects despite the fact that *bantam* depletion alone does not produce a phenotype (Fig. 3B,I-K). These data reinforce that the MLR complex has a role in controlling *bantam* levels during eye development.

3.4. The MLR complex regulates *bantam* expression in a tissue- and differentiation-specific context

The phenotypes caused by MLR subunit knockdown both in the eye and in the wing are similarly sensitive to *bantam* levels, and the MLR complex directly regulates both the eye- and wing-specific *bantam* enhancers, suggesting that the MLR complex attenuates *bantam* expression in the eye imaginal disc as it does in the wing disc. To investigate this, the *bansensGFP* construct was used as an inverse assay of *bantam* activity in the eye disc. The eye disc is a useful developmental model as, unlike the wing, it contains cells at multiple stages of differentiation and development. A mobile boundary known as the morphogenetic furrow induces differentiation as it migrates from the posterior to the anterior margin of the eye pouch, marking the induction of differentiation into separate lineages of proneuronal and interommatidial cells (Fig. 4A). Therefore, tissue anterior to the furrow remains undifferentiated while that posterior to the furrow has begun synchronized differentiation, eventually giving rise to the multiple cell types that will make up the adult compound eye (Cagan, 2009). In wild type eye discs, *bantam* levels remain

high in undifferentiated tissue and vary by cell type once differentiation commences (Fig. 4B) (Tanaka-Matakatsu et al., 2009). Knockdown of *Cmi* or *trr* driven by *Ey-Gal4* occurs within both the undifferentiated and differentiating regions of eye tissue (Fig. 4D), and its effects of *bantam* expression appear to be region-specific: *bantam* activity in cells anterior to the morphogenetic furrow is unchanged compared to control, yet posterior to the furrow *bansensGFP* signal significantly increases, representing a decrease in *bantam* activity (Fig. 4J and K). These effects are confirmed using two additional Gal4 drivers: knockdown driven by either *GawB69B-Gal4* (ubiquitous throughout the entire disc (Fig. 4E)) or *DE-Gal4* (within the dorsal half of the eye pouch, leaving the ventral half as an internal wild type control (Fig. 4F)) has no effect on *bantam* activity anterior to the furrow but causes significant increase in *bansensGFP* signal posterior (Fig. 4L–O,R). These results suggest that the MLR complex is not required for maintaining *bantam* transcription in undifferentiated eye disc cells but may have critical functions in epigenetic reprogramming of cell-type specific *bantam* enhancers during differentiation. It is unclear at what stage of this process the MLR complex is necessary for reprogramming. This was investigated using *GMR-Gal4*, which drives knockdown only in cells posterior to the furrow (Fig. 4G). If MLR complex activity is required in differentiating cells for proper reprogramming of *bantam* expression, then depletion of *Cmi* or *trr* in this region will match the other Gal4 drivers. Unexpectedly, *GMR-Gal4* driven *Cmi/trr* knockdown had no effect, resulting in eye discs demonstrating wild type expression of *bantam* (Fig. 4P–Q,R). Taken together, these data suggest that the MLR complex has regulatory function in undifferentiated cells necessary for proper *bantam* transcription upon differentiation but is dispensable for maintaining *bantam* expression programming once differentiation has commenced.

The reduction in *bantam* activity observed in differentiating cells does not appear to be uniform, but rather sporadic (Fig. 3H-M). These cells have begun cell fate decisions that separate them into patterns of specific lineages (Fig. 4A), suggesting that perhaps cells of only particular lineages are affected by MLR subunit knockdown. In wild type eye discs, *bantam* is downregulated in proneuronal cells at the center of each developing ommatidia and upregulated in the interommatidial cells bordering the compound eye units (Fig. 4S) (Tanaka-Matakatsu et al., 2009). Differentiating cells in *Cmi* or *trr* knockdown discs were examined via confocal microscopy, revealing two separate effects: *bantam* is simultaneously upregulated in the differentiating proneuronal ommatidial cells (marked by Elav staining) and downregulated in interommatidial cells. These data reveal not only that the MLR complex has an important and unexpected role in establishing *bantam* expression levels prospectively in undifferentiated eye cells, but also that this regulatory activity is necessary for proper cell type-specific *bantam* expression in subsequent cell generations in response to different differentiation decisions.

3.5. The MLR complex regulates apoptosis in the developing eye

The reduced eye size and disrupted ommatidia phenotype caused by knockdown of *Cmi* or *trr* may reflect elevated apoptosis during organ development, and *trr* has previously been characterized as protective against programmed cell death in the eye (Kanda et al., 2013), implicating a role for the MLR complex in cell survival. A well-characterized function of *bantam* in the eye disc is the inhibition of apoptosis through translational blocking of proapoptotic *hid* (Brennecke et al., 2003; Grether et al., 1995), and the MLR activity is required in the eye to properly regulate *bantam* expression during differentiation (Fig. 4). Increased *bantam* function is associated with the rough and shrunken phenotype (Fig. 3), suggesting that *bantam* dysregulation may cause altered developmental apoptosis, leading to the malformed adult eye. To address whether *Cmi/trr* knockdown is associated with altered apoptosis, we stained eye discs for activation of effector caspase Dcp-1 (Song et al., 1997). Eye discs undergo sporadic apoptosis in undifferentiated tissue and regulated pruning of cells once the morphogenetic furrow passes and synchronized differentiation commences (Fig. 5A) (Brachmann and

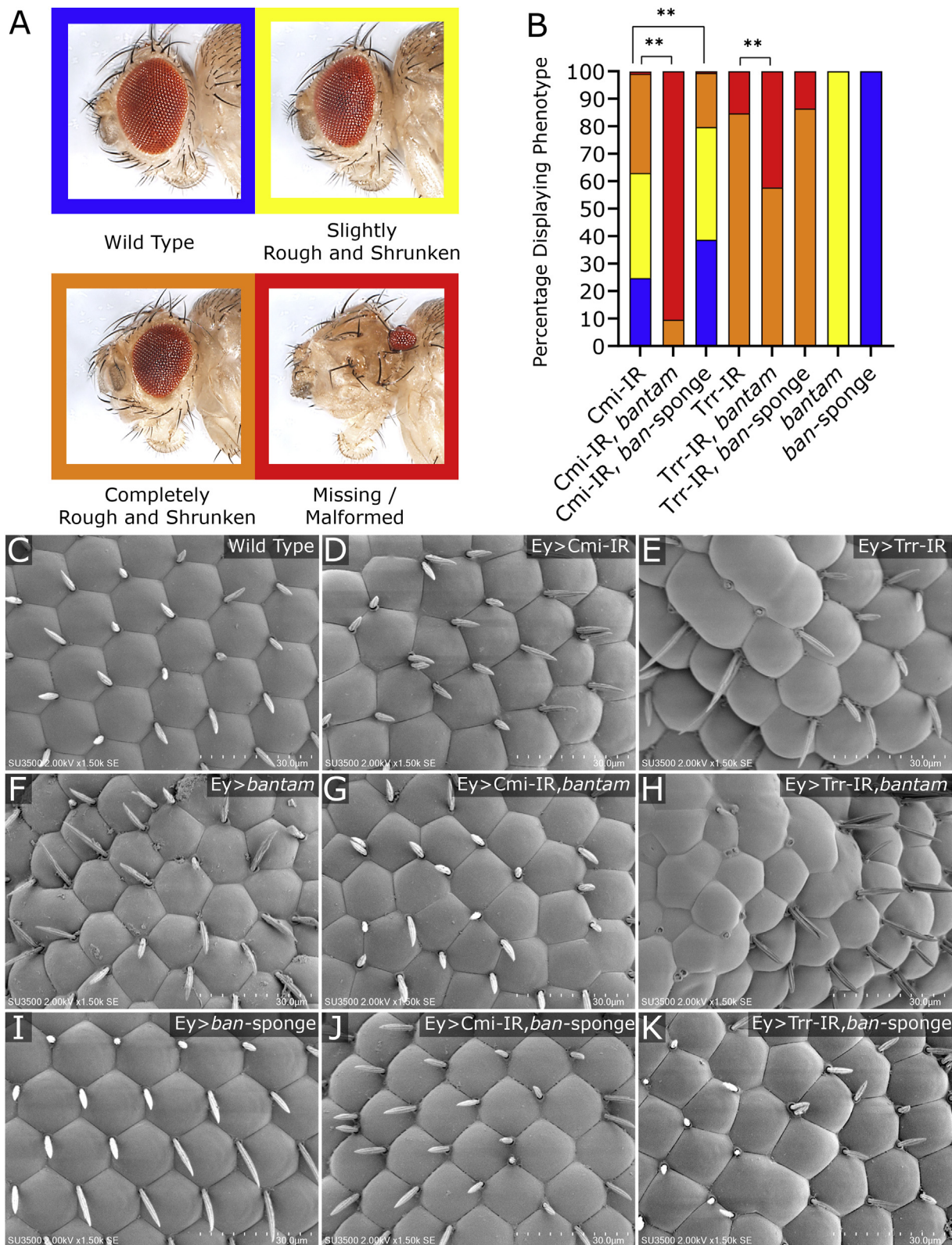


Fig. 3. The *MLR* complex genetically interacts with *bantam* during eye development. *ey-Gal4* was used to drive expression of genetic constructs modulating levels of *Cmi*, *Trr*, or *bantam* in the eye disc. (A) Knockdown of *Cmi* or *trr* during eye development results in rough and shrunken eyes that range in penetrance and expressivity. (B) Adults of the listed genotypes were scored according to eye phenotype severity; $N > 50$ for all genotypes; $** = p \leq 0.01$. Overexpression of *bantam* significantly enhances the *Cmi* or *trr* knockdown phenotypes, while reduction of *bantam* activity using the *ban-sponge* suppresses the *Cmi* knockdown phenotype. Overexpression of *bantam* alone causes slightly rough and shrunken eyes; expression of the *ban-sponge* alone has no phenotypic effect. (C–K) SEM images of adult compound eyes demonstrate that roughness is due to ommatidial patterning defects. As compared to wild type ommatidia (C), *Cmi* knockdown (D) and *trr* knockdown (E) eyes display ommatidial crowding, lens fusion, and bristle loss and duplication. This is also seen when *bantam* is overexpressed alone (F), in the *Cmi-IR* background (G), and the *Trr-IR* background (H). (I) Reduction of *bantam* activity has no effect on ommatidial patterning and appears to suppress the effect of *Cmi* (J) or *trr* (K) knockdown.

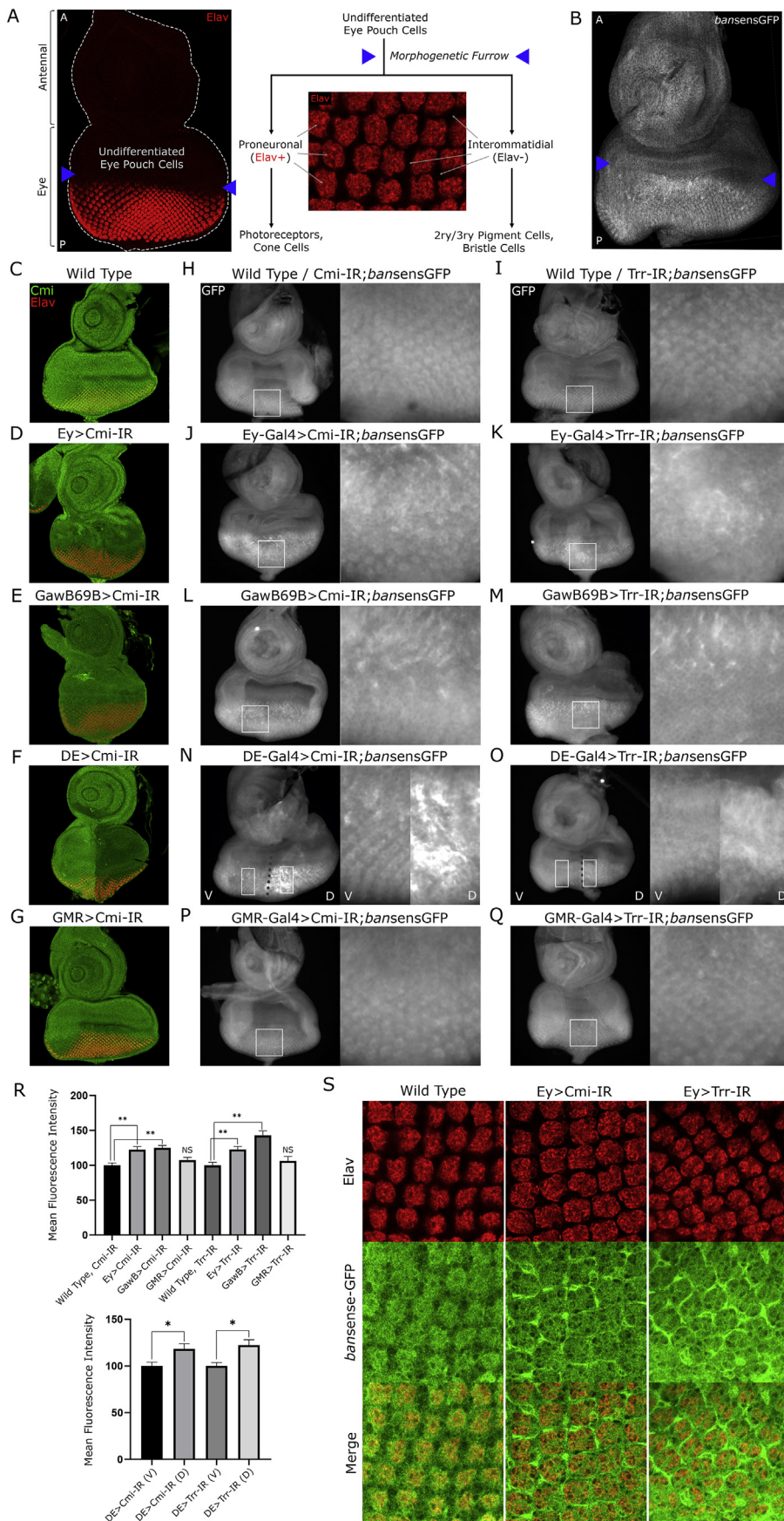


Fig. 4. The MLR complex regulates *bantam* expression. (A) Diagram of compound eye differentiation in the W3L eye disc. To the left, the eye disc is outlined in a thin dotted line. It is comprised of an anterior (A) antennal pouch and a posterior (P) eye pouch. The eye pouch is separated into anterior undifferentiated cells and posterior differentiating cells by the morphogenetic furrow (marked by blue arrow). After passage of the furrow, a cell fate decision is made between proneuronal lineage, positive for Elav (red), and interommatidial lineage. During metamorphic compound eye development, the proneuronal lineage gives rise to photoreceptors and cone cells while the interommatidial lineage gives rise to secondary and tertiary pigment cells as well as bristle cells. (B) Example of *bantamGFP* expression in a wild type eye disc. Undifferentiated eye cells anterior to the morphogenetic furrow (marked by blue arrow) have relatively high *bantam* activity (low *bantamGFP*), and differentiating cells posterior to the furrow have relatively high *bantam* activity (high *bantamGFP*). (C–G) *Cmi* was knocked down using various Gal4 drivers to visualize driver expression pattern and knockdown efficiency. Expression pattern of these drivers is visualized by immunostaining *Cmi* (green) and Elav (red), which labels proneuronal cells posterior to the morphogenetic furrow. (D) *Ey-Gal4* drives knockdown within the eye pouch. (E) *GawB69B-Gal4* drives ubiquitously throughout the organ. (F) *DE-Gal4* drives only within the dorsal half of the eye pouch. (G) *GMR-Gal4* drives only posterior to the morphogenetic furrow. (H–Q) The *bantam-GFP* inverse sensor construct (*bantamGFP*) was used to assay *bantam* miRNA levels. Magnified views of developing ommatidia posterior to the furrow are displayed to the right of each eye disc. *bantam* activity in the undifferentiated anterior tissue anterior to the furrow remain unchanged in all genotypes. (H–I) In control discs, *bantam* activity is relatively high anterior to the furrow (low GFP) and lower posterior (high GFP). (J–O) When *Cmi* or *trr* is knocked down both anterior and posterior to the furrow (*ey >*, *GawB69B >*, and *DE >*), *bantam* activity appears to decrease in the differentiating posterior tissue. (P–Q) If *Cmi/Trr* are lost only within this differentiating tissue (*GMR >*), *bantam* activity remains unchanged compared to control. (R) Mean fluorescence intensity anterior to the furrow was quantified from cohorts of each genotype; $N \geq 10$ for all genotypes; * = $p \leq 0.05$, ** = $p \leq 0.01$, NS = not significant. (S) Developing ommatidia in eye discs were stained for pro-neuronal marker Elav (red) and *bantamGFP* (green). Control organs demonstrate colocalization of GFP and Elav. Upon knockdown of either *Cmi* or *trr* (*Ey >*), changes in *bantam* expression vary by cell fate. In proneuronal cells *bantam* activity is increased (lower GFP), and in interommatidial cells *bantam* activity is decreased (higher GFP).

Cagan, 2003), yet knockdown of either *Cmi* or *trr* in the eye pouch results in increased Dcp-1 activation in the anterior undifferentiated section of the eye pouch, with a significant concentration of these cells on the dorsal-ventral midline (Fig. 5B and C), the same location previously identified in *trr* mutant clones (Kanda et al., 2013). TUNEL staining as a second apoptotic marker confirmed that these cells are undergoing programmed cell death (Supp. Fig. 1A), and knockdown of *Cmi* or *trr* exclusively in the dorsal eye region via the DE-Gal4 driver suggests that this effect is cell autonomous (Supp. Fig. 1B). These results suggest that reduction of MLR activity in undifferentiated eye cells has a deleterious

effect on the survival of those cells. Notably, *Cmi* or *trr* knockdown driven by *Ey*-Gal4 has no effect on *bantam* activity in that region (Fig. 4J and K); thus, dysregulation of the miRNA does not appear likely to be the mechanism underlying the apoptotic effect. However, if the apoptotic phenotype is sensitive to *bantam* levels, this would suggest involvement of *bantam* regulatory targets. To investigate this, Dcp-1 activation was examined in discs overexpressing *bantam* or the *ban*-sponge in combination with *Cmi* or *trr* knockdown (Fig. 5D–I). While the role of *bantam* and the results of its overexpression within differentiating eye cells posterior to the morphogenetic furrow has been previously described (Thompson and

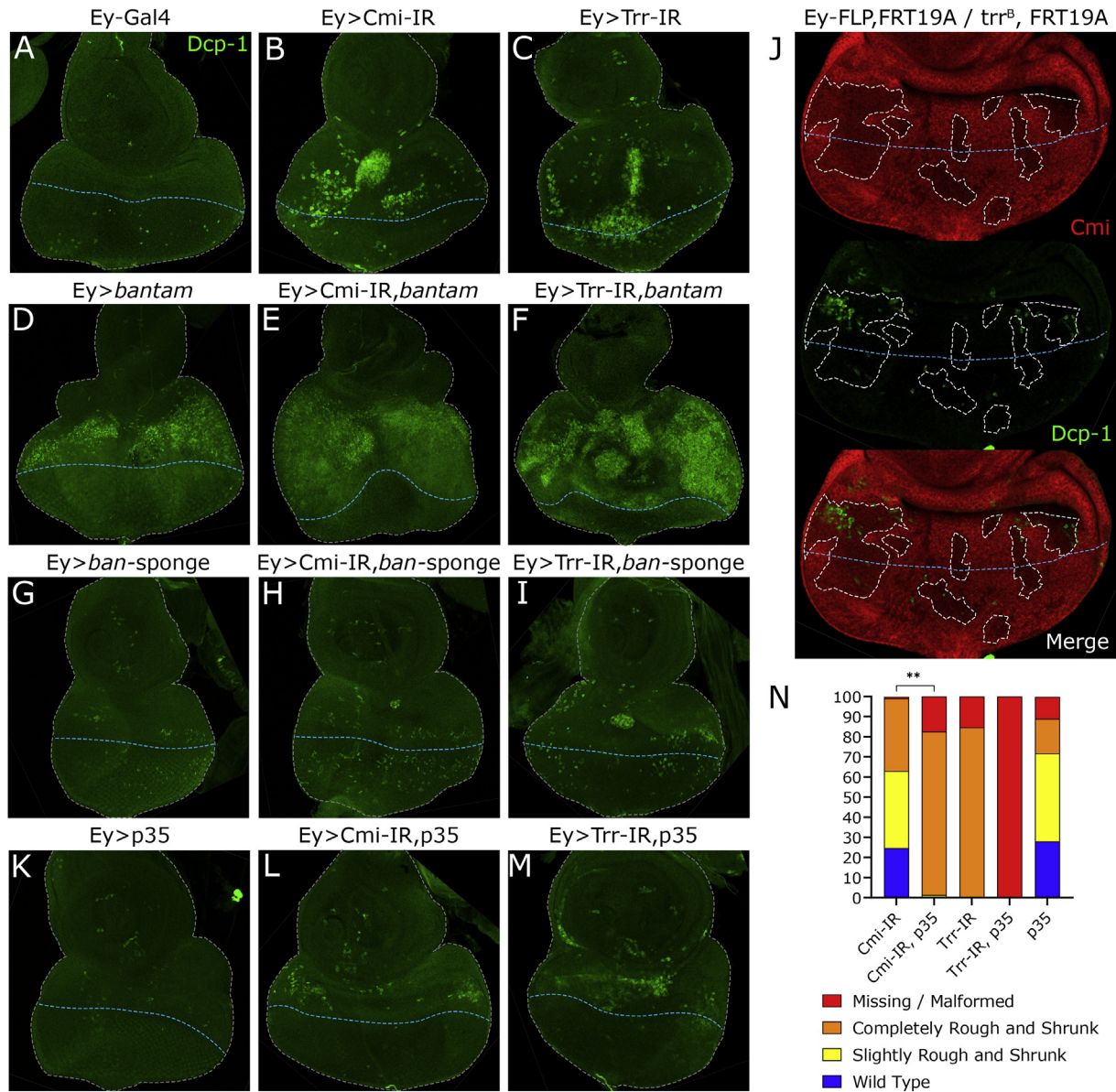


Fig. 5. The MLR complex suppresses caspase activation in the undifferentiated eye. Eye discs were dissected from wandering third instar larvae and assayed for apoptosis; the morphogenetic furrow is marked by a blue dotted line and the discs are outlined by a gray dotted line. (A–I, K–M) *Ey*-Gal4 was used to drive expression of genetic constructs modulating levels of *Cmi*, *Trr*, *bantam*, or *p35* in the eye disc. Apoptosis in the eye disc was assayed by staining for cleaved effector caspase Dcp-1 (green). (A–C) Knockdown of *Cmi* or *trr* results in effector caspase activation concentrated in a cluster of apoptotic cells on the dorsal-ventral midline anterior to the morphogenetic furrow. (D) Overexpression of *bantam* causes eye pouch overgrowth and caspase activation in undifferentiated tissue. (E–F) Increased *bantam* expression in the background of *Cmi*/*trr* knockdown enhances caspase activation. While reduction of *bantam* activity through expression of the *ban*-sponge has no effect on caspase activation (G), it suppresses the apoptotic induction of *Cmi*/*trr* knockdown organs (H–I). (J) *trr*-null somatic clones were generated using the FLP-FRT recombinant system driven by *ey*-FLP. Null clonal patches are marked by absence of *Cmi* staining (red) and outlined in gray dashed line. *trr*-null clones display increased caspase activation (green) compared to neighboring control tissue. (K) Caspase inhibitor *p35* represses caspase activation. (L–M) *p35* expression alongside *Cmi*/*trr* knockdown suppresses the caspase phenotype. (N) Adult eye phenotype was scored and quantified. *p35* expression phenocopies the rough and shrunken phenotype of *Cmi*/*trr* knockdown and enhances the phenotype in a *Cmi*/*trr* knockdown background; ** = $p \leq 0.01$. Simultaneous *trr* knockdown and *p35* expression results in synthetic lethality; therefore, statistical significance between *Trr*-IR and *Trr*-IR, *p35* not achieved due to low number of surviving adults.

Cohen, 2006), the effects of *bantam* modulation in undifferentiated eye tissue is uncertain. Surprisingly, overexpression of the miRNA within the entire eye pouch induces widespread caspase activation in undifferentiated cells, causes overgrowth of the eye pouch, and enhances the Dcp-1 activation phenotype in a *Cmi* or *trr* knockdown background (Fig. 5D–F). Reduction in *bantam* activity through the *ban*-sponge suppresses this phenotype (Fig. 5G–I). These results suggest that the MLR complex plays a role in protecting undifferentiated eye cells against apoptotic death, likely through a mechanism involving target(s) of *bantam*.

The increase in caspase activation in undifferentiated eye tissue upon knockdown of *Cmi* or *trr*, particularly the midline concentration of apoptotic cells, may be due to a variety of cell intrinsic and/or extrinsic effects. To investigate this, *trr*-null somatic clones were generated in the developing eye pouch. These mosaic clonal patches also lack *Cmi*, as the MLR complex is unstable in the absence of *Trr* (Zraly et al., 2020). While undifferentiated eye tissue deficient in *Cmi* or *Trr* displays increased sporadic levels of caspase activation compared to neighboring control tissue, the aggregation of apoptotic cells at the dorsal-ventral midline is absent, even in patches of null clones at this location (Fig. 5J, Supp. Fig. 1C and D). These results suggest that while deficiency in MLR complex activity promotes programmed cell death cell-autonomously in undifferentiated eye tissue, the centralized concentration of apoptotic cells seen in knockdown discs requires MLR complex depletion in a widespread area of eye tissue.

Given the fact that caspase activation in undifferentiated tissue as well as rough and shrunken adult eyes are both associated with reduced MLR complex activity and are similarly sensitive to *bantam* levels, we hypothesized that the aberrant cell death in the undifferentiated eye is causal to smaller adult organs with patterning defects. To verify this, caspase activation in the eye disc was suppressed by expression of p35, a baculovirus substrate inhibitor of caspases including Dcp-1 (Song et al., 2000; Zoog et al., 1999). If increased cell death in undifferentiated eye tissue is mechanistically linked to the rough and shrunken adult eyes, then suppression of apoptosis by p35 should rescue the adult phenotype. Intriguingly, while p35 successfully reduces apoptotic activation by *Cmi* or *trr* knockdown (Fig. 5K–M), it significantly enhances the adult phenotypes (Fig. 5N). From these data, we conclude that the adult eye phenotype is not a result of the induction of apoptosis in the undifferentiated eye disc. Rather, these are two independent effects both resulting from the loss of MLR complex activity.

4. Discussion

The highly conserved MLR COMPASS complexes serve to commission, bookmark and maintain epigenetic *cis*-regulatory controls on transcription enhancers, thus helping to fine tune signaling pathways that are critical for animal development and cellular homeostasis (Fagan and Dingwall, 2019; Rickels et al., 2017; Wang et al., 2018; Zraly et al., 2020). We recently showed that the complex displayed developmental context-dependent chromatin enrichments that linked regulatory functions of the complex to potential *in vivo* target genes, including those important for developmental pattern formation and growth control (Zraly et al., 2020). We therefore sought to explore *in vivo* mechanisms of target gene regulation by the Drosophila MLR complex and discovered two key features: 1) We have identified the miRNA *bantam* as an important regulatory target of the complex during tissue development with roles in both activating and repressing *bantam* expression depending on the context of cell fate (Fig. 6). To our knowledge, this is the first demonstration that the complex has a role in both positive and negative regulation of a single transcriptional target in the same tissue. 2) The MLR complex has important functions in enhancer maintenance and sustaining cell survival in undifferentiated cells.

We found that the MLR complex localizes to tissue-specific *bantam* enhancers during organ development, regulating the activity of those regions and the expression levels of the *bantam* miRNA. This regulatory

activity is consequential for the development of these organs. We previously showed that the complex was vital for proper Dpp signaling during wing vein formation (Chauhan et al., 2013). In this report, we show that the complex is necessary for attenuating *bantam* expression in the wing via repression of a *cis*-acting distal *bantam* wing enhancer. *bantam* acts as a negative regulator of Dpp signaling through translation inhibition of Mad, a downstream effector of Dpp (Kane et al., 2018); thus, the vein patterning defects resulting from altered Dpp signaling upon reduction of MLR complex activity may be exacerbated by elevated *bantam* levels. These results reveal that proper expression control of *bantam* by the MLR complex is necessary for correct tissue patterning during wing development, likely through regulation of Dpp/Tgf- β signaling.

The RNAi knockdown of *Cmi* and *trr* in the developing eye tissue produces a complex phenotype, including reduction of eye tissue and pattern disruptions in the remaining ommatidia. This likely is the result of multiple altered transcriptional targets at various stages of compound eye development. The rough and shrunken phenotype is reminiscent of eye malformations caused by increased cell death or altered developmental signaling (Brennecke et al., 2003; Chao et al., 2004; Nolo et al., 2006), and *bantam* is recognized as both an inhibitor of apoptosis as well as a feedback regulator of multiple signaling pathways (Brennecke et al., 2003; Kane et al., 2018; Oh and Irvine, 2011; Wu et al., 2017; Zhang et al., 2013). While *bantam* overexpression exclusively in the differentiating ommatidia suppresses apoptosis and results in eye tissue overgrowth (Thompson and Cohen, 2006), we demonstrate here that overexpression of the miRNA throughout the eye pouch induces caspase activation in undifferentiated cells through unknown mechanisms and phenocopies the rough and shrunken eyes resulting from reduced functions of the MLR complex. Massive cell death during development can result in reduced organ size, yet we demonstrate not only that the adult phenotype is not caused by increased apoptosis during development, but that caspase inhibition enhances the severity of the phenotype, potentially through suppression of non-apoptotic developmental caspase function (Kuranaga and Miura, 2007; Lamkanfi et al., 2007; Nakajima and Kuranaga, 2017). The fact that undifferentiated cell loss due to apoptosis does not affect the adult organ can be explained by

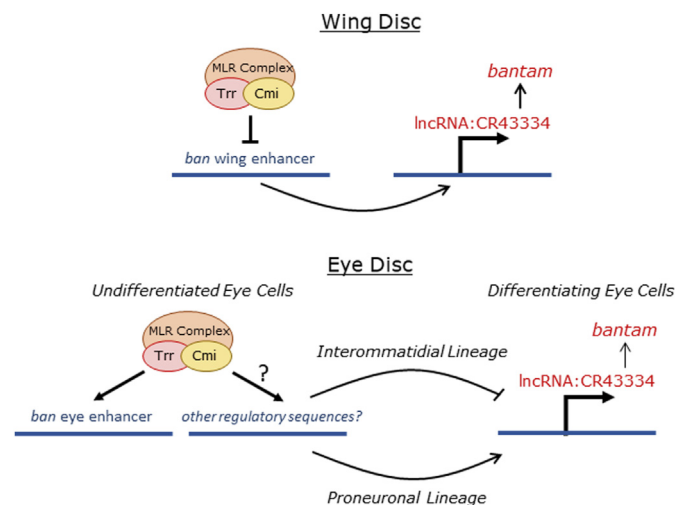


Fig. 6. Model of MLR functions in *bantam* enhancer control. The Drosophila MLR COMPASS-like complex may either have positive or negative regulatory activity on the miRNA *bantam* depending on the context of cell fate. Tissue-specific *bantam* enhancers active in undifferentiated wing or eye tissue are bound and regulated by MLR. In the wing disc, MLR represses the activity of the *bantam* wing enhancer and downregulates *bantam* miRNA. In the eye disc, MLR modulates the expression pattern of the *bantam* eye enhancer in undifferentiated cells. After passage of the morphogenetic furrow and cell fate choice between proneuronal or interommatidial lineages, the previous regulatory activity of MLR leads to increased *bantam* expression in the proneuronal cells and decreased expression in the interommatidial.

compensatory proliferation of surviving neighbor cells to replace those lost or other caspase-related growth signals (Fan and Bergmann, 2008; James and Bryant, 1981) (Shinoda et al., 2019; Song et al., 1997). These data not only further elucidate *bantam*'s roles in eye development, but also favor the hypothesis that alteration of developmental signaling rather than growth regulation underlies the adult eye defects we observe.

The importance of the MLR complex in contributing to either positive or negative regulation of a single transcriptional target depending on cell fate context is a novel observation with critical developmental consequence. This regulatory decision is likely not due to activity inherent to the complex itself, but rather influenced by the transcription factors that recruit the complex to specific developmental enhancers. Transcriptional control of *bantam* is orchestrated via complex signaling factors that regulate *bantam* enhancers (Peng et al., 2009; Slattery et al., 2013). For example, while Hippo effector Yki binds to and activates both the *bantam* eye and wing enhancers, binding partner Hth is necessary for regulating the eye enhancer and controlling *bantam* expression in the eye while separate partner Sd is necessary for regulating *bantam* expression in the wing through the wing enhancer (Peng et al., 2009; Slattery et al., 2013; Zhang et al., 2008). Multiple regulatory inputs are necessary for guiding proper spatiotemporal expression of the *bantam* miRNA in imaginal tissue, and our data suggests that the MLR complex plays a critical role in translating these inputs into regulatory decisions.

Negative regulation of a transcriptional target is a relatively novel role of an MLR complex. We have previously detailed how the Drosophila MLR complex is required to suppress ecdysone-response elements (EcREs) to prevent premature activation; loss of MLR subunits results in both upregulation of the regulatory targets as well as inability to properly respond to future stimulation (Zraly et al., 2020). While the mechanisms of repression that require the MLR complex at the *bantam* wing enhancer or at the elements controlling *bantam* in interommatidial-lineage eye cells are unknown, it is probable that the complex plays a similar role at these as at EcREs: inhibiting regulatory elements poised for activation. Much remains to be determined concerning these newly elucidated aspects of MLR complex function.

Through depletion of MLR complex activity in different regions of the eye imaginal disc, we determined that the complex is required in undifferentiated eye cells for proper regulation of *bantam* expression that occurs after furrow progression and the onset of differentiation into ommatidia, developmental stages separated by at least two rounds of mitosis (Baker, 2001; Cagan, 2009). Our eye disc studies correlate with *in vitro* data that the Kmt2D MLR complex is dispensable for maintaining gene transcription in murine ESCs, but is required for the transcriptional reprogramming that occurs during differentiation and dedifferentiation (Wang et al., 2016). Thus, our data not only verifies this role of MLR complexes through an *in vivo* developmental model but further demonstrates that this necessary preparatory activity may occur cell generations prior to the reprogramming event. We did not observe widespread changes to *bantam* activity in undifferentiated eye tissue upon *Cmi/trr* knockdown, suggesting that MLR complex activity is not required for maintaining widespread *bantam* expression after the initial activation event and establishment of enhancer functions that likely occurred during embryogenesis (Brennecke et al., 2003; Rickels et al., 2017; Rickels and Shilatifard, 2018; Zraly et al., 2020). Thus, the MLR complex appears to be dispensable for the maintenance of previously active enhancers that drive widespread activity during development, but essential for the *de novo* programming of tissue-specific enhancers that occurs at critical developmental transitions, perhaps similar to the enhancer priming functions of Kmt2D in murine ESCs (Wang et al., 2016).

We found that the MLR complex plays an unanticipated role in cell survival separate from *bantam* regulation. Knockdown of *Cmi* or *trr* in undifferentiated cells of the eye disc results in cell-autonomous elevation of executioner caspase (Dcp-1) signaling and positive TUNEL staining, indicative of caspase cascade and apoptotic cell death. Previous investigations in the Drosophila eye disc as well as human germinal B cells have reported that deletion of MLR complex methyltransferases both induces apoptosis and

provides a proliferative advantage in undifferentiated cells, possibly through inhibition of differentiation (Kanda et al., 2013; Ortega-Molina et al., 2015). These results and ours suggest that MLR complexes play critical roles in multipotent cell maintenance and preparation for differentiation during cell fate transitions. While the apoptotic phenotype caused by *Cmi/trr* knockdown in the larval eye disc is not directly caused by dysregulated *bantam* expression, its sensitivity to *bantam* levels suggests the likely involvement of signaling pathway(s) active in that area and regulated by both the MLR complex and *bantam*. While many developmental pathways are necessary for proper eye development, the Notch signaling pathway is activated at the dorsal-ventral midline to promote proliferation and survival (Chao et al., 2004), and Notch signaling is dependent on both the MLR complex and *bantam* regulation (Becam et al., 2011; Dhar et al., 2018; Giaimo et al., 2017; Oswald et al., 2016; Wu et al., 2017).

Clearer comprehension of MLR complexes' roles in enhancer regulation during normal development paves the way to better determine the mechanisms of disease associated with loss of complex function. The ability to regulate a single transcriptional target in opposite directions depending on tissue type and cell fate choice is critical for normal development. The miRNA *bantam* has previously been identified as tumorigenic in *Drosophila* cancer models (Sander and Herranz, 2019). Potential human counterparts of *bantam* have been identified, including *mir-450b* that contains close sequence similarity (Ibanez-Ventoso et al., 2008) and has been described as suppressing cancer cell proliferation and inducing protective differentiation (Sun et al., 2014; Zhao et al., 2014). The human *mir-130a* can act as either oncogene or tumor suppressor, impacts drug resistance (Zhang et al., 2017), and is functionally orthologous to *bantam* in its feedback regulation of Hippo pathway signaling (Shen et al., 2015). The discovery of the importance of miRNAs in both normal development and cancer together with emerging evidence for the fine-tuning of miRNA expression through MLR-dependent enhancer regulation, provides a strong rationale for the expansion of efforts to define the role of epigenetic gene control in the context of cell fate decisions.

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Appendix A. Supplementary data

Supplementary data to this article can be found online at <https://doi.org/10.1016/j.ydbio.2020.09.007>.

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