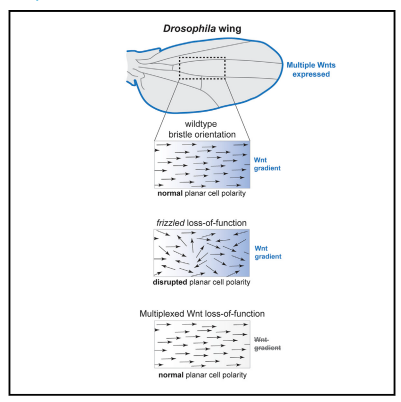
No Evidence that Wnt Ligands Are Required for Planar Cell Polarity in *Drosophila*

Graphical Abstract



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In Brief

Previous studies have come to differing conclusions on whether Wnt ligands provide a tissue-level orientation cue for the planar cell polarity pathway. Ewen-Campen et al. re-examine this question in *Drosophila* using multiplex *in vivo* CRISPR and double RNAi against Wnt ligands and find no evidence that Wnts are required for PCP patterning.

Highlights

- Previous studies disagree on whether Wnt ligands affect planar cell polarity (PCP)
- Multiplex in vivo CRISPR or RNAi against multiple Wnt ligands does not alter PCP
- CRISPR against wntless/evi does not affect PCP
- We find no evidence that Wnt ligands are required for PCP in Drosophila







Report

No Evidence that Wnt Ligands Are Required for Planar Cell Polarity in *Drosophila*

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https://doi.org/10.1016/j.celrep.2020.108121

SUMMARY

The *frizzled* (*fz*) and *dishevelled* (*dsh*) genes are highly conserved members of both the planar cell polarity (PCP) pathway and the Wnt signaling pathway. Given these dual functions, several studies have examined whether Wnt ligands provide a tissue-scale orientation cue for PCP establishment during development, and these studies have reached differing conclusions. Here, we re-examine this issue in the *Drosophila melanogaster* wing and notum using split-Gal4 co-expression analysis, multiplex somatic CRISPR, and double RNAi experiments. Pairwise loss-of-function experiments targeting *wg* together with other Wnt genes, via somatic CRISPR or RNAi, do not produce PCP defects in the wing or notum. In addition, somatic CRISPR against *evi* (aka *wntless*), which is required for the secretion of Wnt ligands, did not produce detectable PCP phenotypes. Altogether, our results do not support the hypothesis that Wnt ligands contribute to PCP signaling in the *Drosophila* wing or notum.

INTRODUCTION

Planar cell polarity (PCP) refers to the coherent orientation of cells in a sheet and is driven by a conserved molecular pathway that is required for the proper development and function of many animal tissues (Adler, 2012; Butler and Wallingford, 2017; Goodrich and Strutt, 2011; Yang and Mlodzik, 2015). Decades of functional analyses in the epithelia of *Drosophila* and other model organisms have led to the identification of six highly conserved "core" pathway members, whose asymmetric localization and function drives downstream manifestations of PCP, both autonomously and non-autonomously. In cells of the developing fruit fly wing, the transmembrane protein Frizzled (Fz) and the cytosolic proteins Dishevelled (Dsh) and Diego (Dgo) accumulate at the distal cortex. At the proximal cortex, an asymmetric signaling complex forms that includes the transmembrane protein Strabismus aka Van Gogh (Stbm/Vang) and the cytosolic protein Prickle (pk). The sixth core pathway member, Flamingo aka Starry night (Fmi/Stan), accumulates both proximally and distally (Adler, 2012; Butler and Wallingford, 2017; Goodrich and Strutt, 2011; Yang and Mlodzik, 2015). Loss of function of any core pathway members leads to PCP defects, which ultimately manifest in mispolarization of the actin-rich wing hairs (Gubb and García-Bellido, 1982; Wong and Adler, 1993). In addition to the core pathway, a second PCP pathway that includes Fat, Dachsous, and Four-jointed has been identified in Drosophila (Adler, 2012; Butler and Wallingford, 2017; Goodrich and Strutt, 2011; Yang and Mlodzik, 2015).

At relatively small spatial scales, feedback interactions among core PCP pathway components provide a working mo-

lecular model for PCP within cells and between neighboring cells. However, there remains active debate regarding the molecular mechanisms that allow core pathway members to correctly orient regarding "global" patterning at the level of whole tissues. Researchers have long hypothesized that there may be a secreted molecule (referred to as "Factor X" by Lawrence et al., 2002) that coordinates tissue-scale morphogenesis to the PCP pathway. Among the possible candidates for "Factor X," Wnt ligands have long been considered "obvious candidates for such factors" (Goodrich and Strutt, 2011) because both Fz and Dsh are highly conserved members of this signaling pathway: Fz and its paralog Fz2 function as co-receptors for Wnt ligands (Bhanot et al., 1999; Chen and Struhl, 1999; Müller et al., 1999), and Dsh is also a highly conserved component of the Wnt ligand-receptor signalosome (Perrimon and Mahowald, 1987; Sharma et al., 2018). However, a number of loss-of-function studies of Wnt ligands have failed to find any PCP phenotypes (Chen et al., 2008; Lawrence et al., 2002). Specifically, Lawrence et al. (2002) examined clones in the adult abdominal epithelium homozygous for a deletion covering four of the seven Drosophila Wnt ligands (wg, wnt4, wnt6, and wnt10), and found no PCP defects within or nearby such clones. Chen et al., (2008) generated quintuple clones in the wing lacking wg, wnt2, wnt4, wnt6, and wnt10, as well as clones lacking porcupine, a gene necessary for the correct lipid modification of Wnt ligands, and found no evidence of PCP defects in either case. In addition, homozygous mutants for evi (aka wntless), which are defective in Wnt secretion, display wild-type PCP (Bartscherer et al., 2006)





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Because Wg is required in early larval stages to specify the presumptive wing (Ng et al., 1996), the above studies relied on the analysis of wg mutant clones in an otherwise wild-type background and thus could not conclusively rule out a role for longrange Wnt signaling from surrounding cells. In contrast, some gain-of-function studies have demonstrated that overexpression of either Wg or Wnt4 (but not other Wnt ligands) in ectopic regions of the developing wing can reorient the polarity of surrounding wing hairs (Lim et al., 2005; Wu et al., 2013) and that this reorientation requires fz (Wu et al., 2013), although a study in the abdomen did not see re-polarization when overexpressing the seven Wnt ligands in a dachsous -/- background (Casal et al., 2006). In addition, a study that bypassed the early requirements for wg using a temperature-sensitive mutation indicated that double-mutant flies lacking wg and wnt4 displayed statistically significant PCP phenotypes in the pupal and adult wing (Wu et al., 2013). Furthermore, studies in vertebrates have implicated certain Wnt ligands in PCP signaling (Yang and Mlodzik, 2015). Taken together, the evidence on whether Wnts, either individually or in combination, influence PCP signaling remains inconclusive.

In this study, we re-examine whether Wnt ligands are required for PCP using systematic double and triple CRISPR via Gal4-UAS-driven Cas9 and double RNAi. We independently analyzed the expression of each Wnt using a collection of knockin split-Gal4 reporters and performed multiplex somatic CRISPR using short guide RNAs (sgRNAs) targeting wg in combination with each of the other six Wnt genes, as well as various combinations thereof. In addition, we used somatic CRISPR to target evi (wntless), a protein required for the secretion of Wnt genes except wntD (Herr and Basler, 2012). Last, we performed double RNAi experiments against wg, wnt4, and wnt6 in the wing and in the notum. We did not detect PCP phenotypes in any of these loss-of-function conditions, in either wing or notum. In contrast to previous reports, our results do not support the hypothesis that Wnt ligands impinge on PCP signaling. We note that our data are consistent with a study recently posted on bioRxiv (Yu et al., 2020), which we discuss below.

RESULTS

Split-Gal4 Knockin Reporters Confirm that Multiple Wnt Genes Are Co-expressed in the Larval Wing Disc

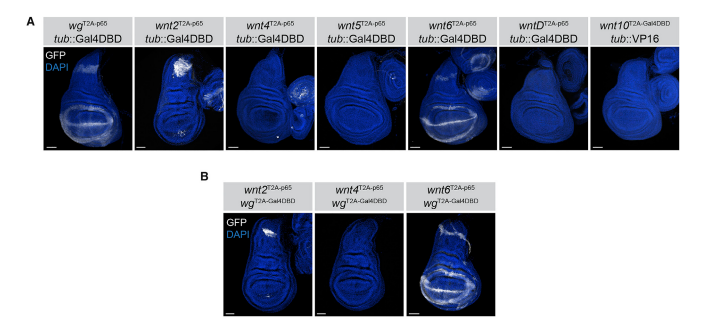
To characterize which Wnt genes are expressed in the wing disc during the period when PCP patterning is established in the developing wing, we systematically re-examined the expression and co-expression of the seven Drosophila Wnt family genes. In the wing disc, core PCP components progressively acquire asymmetrically localization during pupal stages, and there is also evidence that at least some polarity information exists during L3 larval stages in the form of coherent Fmi/Stan localization that is subsequently remodeled during pupal stages (Classen et al., 2005). To independently examine which Wnt ligands are expressed during wing development, we used CRISPR-based homology directed repair to knock in split-Gal4 reporter cassettes into an early exon of each Wnt gene (Figure 1A) (Gratz et al., 2014; Pfeiffer et al., 2010). These cassettes include an in-frame T2A self-cleaving peptide, followed by either the p65 activation domain or the Gal4 DNA binding domain (DBD), and thus allow intersectional labeling of cells expressing multiple Wnt genes. Pilot experiments with wg and wnt2 indicated that this approach recapitulates previously described endogenous gene expression of wg and wnt2, for both p65 and Gal4DBD knockins (Figure S1) (Chen et al., 2008; Kozopas and Nusse, 2002). As predicted, homozygous wg knockins were lethal, and wnt2 knockin homozygotes were male sterile and displayed characteristic wing posture abnormalities (Kozopas and Nusse, 2002; Kozopas et al., 1998). We ultimately created split-Gal4 knockin reagents for each of the seven Wnt genes. In support of the accuracy and utility of these split-Gal4 reporters, we find that they recapitulate known Wnt expression in the adult ovary and that they can be used to characterize Wnt-expressing cells various adult stem cell populations of the gut, and in the larval CNS, which we summarize below.

We crossed this collection of split-Gal4 reagents to a ubiquitously expressed cognate (Gal4DBD or VP16 driven by the tubulin promoter) and examined the expression in larval wing discs via a UAS-driven fluorophore (Figure 1A). We observed expression of wg, wnt2, and wnt6 in the L3 wing disc, consistent with previous reports (Baker, 1988; Janson et al., 2001; Kozopas and Nusse, 2002). Split-Gal4 intersection analysis confirmed that wg and wnt2 are co-expressed in the epithelium of the developing notum and a portion of the wing pouch, and that wg and wnt6 are extensively co-expressed throughout their entire expression domain at this stage (Janson et al., 2001) (Figure 1B). In the pupal wing margin, we observed overlapping expression of wg, wnt2, wnt4, and wnt6 (Figure 1C).

Unexpectedly, although we did observe wnt4 expression in the margin of the pupal wing (Figures 1D and S2), we did not detect wnt4 expression in the dorsoventral (DV) boundary at the L3 larval stage, as has been demonstrated via RNA in situ hybridization (Chen et al., 2008; Gieseler et al., 2001; Harris et al., 2016) and a GFP knockin (Yu et al., 2020). Instead, our wnt4 reporter drove UAS:GFP expression in a small cluster of cells at the anterior ventral boundary of the wing pouch and was not detected in the wing margin until early pupal stages (Figures 1B and S2). Given this discrepancy between our reporter line and previous studies, we examined the expression pattern of an existing knockin Gal4 reporter from the CRIMIC collection, which contains a "trojan exon" including a splice acceptor and T2A-Gal4, inserted in the first intron of wnt4 (Lee et al., 2018). The expression of wnt4 (CRIMIC was consistent with our split-Gal4 reporter in the L3 wing disc and largely consistent in the larval CNS (Figure S2). As an additional test, we generated a third knockin reporter (using T2A-Gal4) using an independently derived donor plasmid (Bosch et al., 2020), inserted into the final exon of wnt4. This independent Gal4 insertion (wnt4^{T2A-Gal4}) was consistent with the other two tested in this study and did not show detectable expression in the DV boundary at the L3 stage (Figure S2). To test whether wnt4 may be expressed in the wing disc during earlier stages, we crossed wnt4^{CRIMIC} line to G-TRACE, a lineage-tracing tool that reports both current and past Gal4 expression using two different fluorophores (Evans et al., 2009). This analysis did not reveal any past expression in the wing disc aside from the small cluster of cells on the border of the wing pouch. Using RNA in situ hybridization, we found that Gal4 is transcribed in the DV boundary at the L3







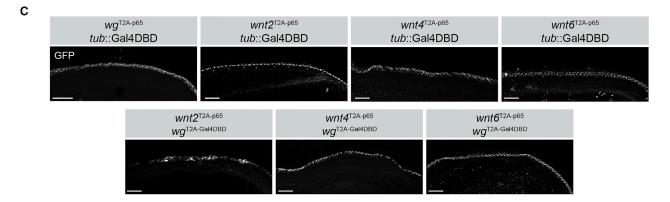


Figure 1. Analysis of Wnt Gene Expression and Co-expression Using Split-Gal4 Reporters in the Larval and Pupal Wing

(A) Expression pattern of each reporter construct, visualized by crossing to a ubiquitously expressed reciprocal split-Gal4 reporter, visualized with UAS:EGFP or UAS:EYFP (see STAR Methods).

- (B) Intersection of wg expression domain with wnt2, wnt4, and wnt6 in the L3 wing disc.
- (C) Co-expression of wg, wnt2, wnt4, and wnt6 in the pupal wing margin. Bottom row shows co-expression of wg with wnt2, wnt4, and wnt6, respectively. Portions of the anterior wing margin of \sim 45–60 h APF pupal wings are shown. Scale bars, 50 μm .

stage in wnt4-T2A-Gal4 flies, suggesting that the Gal4-UAS system introduces a temporal delay in GFP expression relative to endogenous wnt4 expression.

To provide additional validation of the split-Gal4 lines described in this study, we examined Wnt ligand expression in the adult ovary, larval CNS, and two stem cell populations in the adult gut (Figures S3 and S4). In the adult ovary, our split-Gal4 for wg, wnt2, wnt4, and wnt6 are co-expressed in a partially overlapping pattern in the female germarium, consistent with previous descriptions (Waghmare and Page-McCaw, 2018). In addition, we found that wnt5 is also expressed in the adult germarium, in a pattern similar to wnt6 (Figure S3A), which to our knowledge has not been previously reported.

In the adult gut, we find that wg, wnt2, wnt4, and wnt6 are coexpressed in and surrounding adult stem cells located in the cardia (located at the foregut-midgut junction) and, with the exception of wnt2, at the midgut-hindgut juncture (Figure S4) (Singh et al., 2011; Tian et al., 2016). These observations add to our understanding of which ligands are responsible for Wnt signaling observed in these regions of the gut (Tian et al., 2016, 2018) and support the observation that overlapping expression of multiple Wnt ligands is a common feature of different adult stem cell populations in Drosophila.

We also examined Wnt gene expression in the larval nervous system. wg, wnt4, and wnt5 are known to have roles in the developing nervous system (Inaki et al., 2007; Packard et al., 2002; Yoshikawa et al., 2003), and we find that in fact all Wnts are expressed in the larval nervous system. This includes wnt10 (Figure S4). Thus, these split-Gal4 lines may be useful for future studies of Wnt function and redundancy.





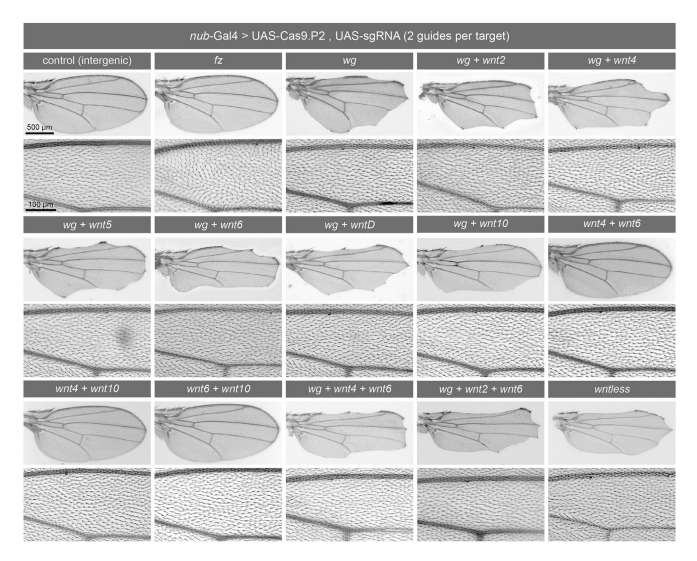


Figure 2. Multiplexed Somatic CRISPR Knockout of Wnt Genes or wntless Does Not Disrupt Planar Cell Polarity UAS:sgRNAs targeting the indicated genes (two guides per target gene) were co-expressed with UAS:Cas9 in the developing wing using nub-Gal4. An example of

a PCP phenotype is shown for fz. Overall wing morphology is shown in the top rows, and magnified views of wing hair orientation in the L3-L4 intervein region are shown in bottom rows; note that CRISPR against wg results in shorter L3-L4 intervein distance. Images are representative of approximately 20 wings analyzed per genotype.

Systematic Pairwise CRISPR against Wnt Genes Using Gal4-UAS:Cas9

Previous studies in Drosophila have demonstrated that ectopic expression of either Wg or Wnt4 can reorient PCP patterning in surrounding wing hairs (Lim et al., 2005; Wu et al., 2013), and Wu et al. (2013) reported that double mutants for wg and wnt4 display disrupted PCP signaling in the pupal and adult wing. We sought to systematically address the role of Wnt ligands in PCP signaling using multiplexed in vivo somatic CRISPR. For these experiments, we identified two effective sgRNAs to target each Wnt ligand (see STAR Methods) and created transgenic lines expressing sgRNAs targeting either one or two Wnt genes simultaneously (with either two or four sgRNAs total, respectively) under UAS control. We confirmed that these sgRNAs cleaved their intended

target in vivo both for single- and double-targeting experiments (Figure S5).

To avoid early roles for Wg in wing specification, we used the nubbin-Gal4 driver to express both Cas9 and sgRNAs. nubbin-Gal4 is expressed throughout the wing pouch beginning in late L2 larval stages (Zirin and Mann, 2007), and thus this approach allowed us to permanently remove target gene function throughout L3, pupal, and adult stages. Control experiments targeting fz confirmed that this approach produces strong PCP phenotypes in the adult wing (Figure 2; Figure S6) and that frizzled is unique among the four Fz-family genes in producing this phenotype, both in single and pairwise CRISPR experiments (Figure S6). Although we did not detect expression of reporters for wnt5, wntD, or wnt10 in the wing disc, we included them in our functional studies in order to rule

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out the possibility that one or more of these ligands is expressed at low levels.

We used somatic CRISPR to target wg together with wnt4, as well as each of the other five Wnt genes and screened for defects in both wing morphology and PCP. As expected, the wing margin was disrupted in all pairwise CRISPR experiments that included wg, and the intervein regions were reduced in size (Figure 2). However, in each case, wing hair orientation was indistinguishable from wild-type (compare with fz panel in Figure 1 and Figure S6 for examples of PCP phenotypes caused by disrupting fz). We next tested pairwise combinations of wnt4, wnt6, and wnt10, which are part of a genomic cluster on the second chromosome, as well as triple CRISPR against wg, wnt4, and wnt6, and against wg, wnt2, and wnt4. We did not observe disrupted PCP in any of these conditions (Figure 2). Last, we used this approach to target evi (wntless), which is required for the secretion of all Wnt genes except WntD (Bänziger et al., 2006; Bartscherer et al., 2006; Goodman et al., 2006; Herr and Basler, 2012). Knocking out evi led to loss of wing margin, essentially phenocopying wa loss of function, and did not produce PCP defects, consistent with observations from evi mutant pharate adults (Bartscherer et al., 2006) (Figure 2). Together, these results suggest that Wg and Wnt4 are not redundantly involved in global PCP patterning and provide no evidence that Wnt ligands direct PCP orientation in the Drosophila wing.

Pairwise RNAi against wg Together with wnt4 or wnt6 **Does Not Produce PCP Phenotypes in the Wing or Notum**

Somatic Gal4-UAS-based CRISPR has been shown to be a powerful and scalable technique for in vivo knockouts (Port and Bullock, 2016; Port et al., 2020). However, one inherent drawback of somatic CRISPR is that it creates mosaic tissues containing both mutant and wild-type cells and does so variably between individuals. This can be seen, for example, in the variable phenotypes and incomplete loss of margin tissue observed when targeting wg (Figure 2). In the case of morphogens such as Wnt ligands, it is conceivable that a small number of wild-type cells could still be sufficient to produce an instructive cue for PCP orientation.

To address this issue, we performed double RNAi experiments against wg, wnt4, and wnt6 in the developing wing using nub-Gal4, using transgenic double-stranded RNA (dsRNA) lines recently shown to produce strong loss-of-function phenotypes and co-expressed with UAS:dcr2 to strengthen RNAi-mediated knockdown (Barrio and Milán, 2020). Under these conditions, wg RNAi led to both a complete loss of the wing margin and a profound growth defect, similar in appearance to the effects of inducing apoptosis in wg-expressing cells by overexpressing the pro-apoptotic genes hid and rpr (Yu et al., 2020). However, even in these severely mispatterned wings, PCP appeared wild-type. Similarly, no PCP defects were observed upon pairwise RNAi against wg and wnt4 or wg and wnt6 (Figure 3). Incidentally, our results confirmed recent observations that wnt4 and wnt6 have opposing effects on modulating the effects of wg-dependent growth (Barrio and Milán, 2020).

Fz-dependent PCP phenotypes can also be studied in the adult notum, by the orientation of bristles (Figure S7A). We thus used pairwise RNAi to knock down wg and wnt4 or wg and wnt6 in the adult notum using pnr-Gal4. Consistent with our observations in the adult wing, we observed no discernable PCP phenotypes in the adult notum in any of these conditions (Figure S7).

DISCUSSION

In this study, we performed loss-of-function studies in the wing and notum using multiplexed somatic CRISPR and double RNAi to systematically target Wnt ligands, and in no combination did we observe PCP phenotypes. In contrast to a study that found a role for Wg and Wnt4 in global PCP establishment (Wu et al., 2013), our results provide no evidence that Wnt ligands, whether singly or redundantly with one another, necessary to provide an instructive global cue to the core PCP pathway. If there is a secreted "Factor X" molecule that provides tissuescale information, our results suggest that it is not a Wnt ligand or combination of two or three Wnt ligands. Previous experiments have suggested that this hypothetical factor is not a member of the Notch, EGF, FGF, or Dpp signaling pathways (Lawrence et al., 2002).

Our results do not rule out the possibility that Wnt ligands could play a redundant role with an additional global cue or cues. For example, a number of studies have suggested that signaling via Fat, Dachsous, and Four-jointed may provide large-scale positional information that is interpreted and refined by the activity of core PCP components within and between neighboring cells (Butler and Wallingford, 2017; Goodrich and Strutt, 2011). The mechanisms that couple local and global PCP thus remain incompletely understood.

Our results imply that the ability of ectopically expressed Wg and Wnt4 to reorient the PCP of surrounding cells is not reflective of such a role in wild-type flies. One possible explanation for these results is that feedback mechanisms between Wnt ligands and Fz proteins may account for the ability of ectopic Wnt ligands to disrupt wild-type Fz function (Cadigan et al., 1998; Chaudhary et al., 2019), irrespective of the physiological role for Wnts under normal conditions. We also note that a separate study found that overexpression of any of the seven UAS:Wnt ligands in clones in the adult abdomen did not cause bristle reorientation (Casal et al., 2006).

During the preparation of this manuscript, a preprint by Yu et al. (2020) reported related data and similar conclusions using several independent approaches. Specifically, Yu et al. (2020) found that normal PCP patterning occurs in the wings of double-mutant flies expressing only membrane-tethered Wg and homozygous for mutations for wnt4, as well as in quintuple mutant lacking wnt2, wnt4, wnt6, and wnt10 and expressing solely membrane-tethered Wg. They also report that evi/wntless loss of function does not interfere with PCP, nor does driving apoptosis in all wg-expressing cells in the developing disc by overexpressing hid and rpr.

One experimental difference between our studies and those reported by Yu et al. (2020) is that the latter use a membranetethered Wg mutant, rather than a knockout or knockdown approach, when addressing Wg function. These mutants lack secreted Wg yet, unlike wg loss-of-function mutants, display nearly normal patterning and development (Alexandre et al.,





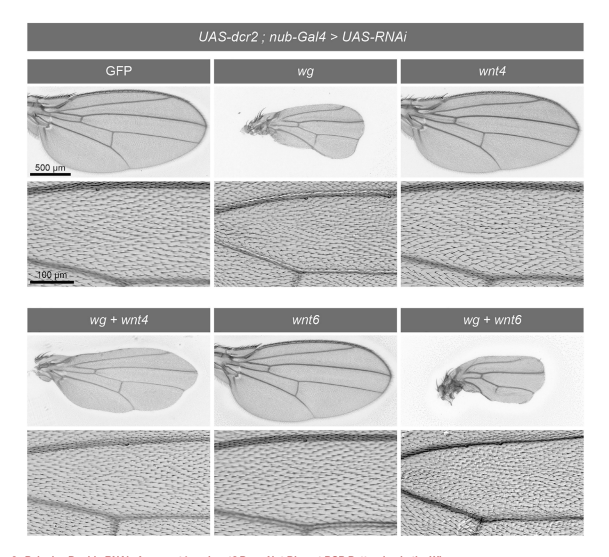


Figure 3. Pairwise Double RNAi of wg, wnt4, and wnt6 Does Not Disrupt PCP Patterning in the Wing UAS:RNAi constructs targeting the indicated genes were co-expressed with UAS:dcr2 in the developing wing using nub-Gal4. Overall wing morphology is shown in the top rows, wing hair orientation in the L3-L4 intervein region is shown in magnification in the bottom rows. Images are representative of approximately 20 wings analyzed per genotype.

2014). Subsequent studies have attributed this phenomenon to a combination of fz2-dependent feedback and perdurance of membrane-tethered Wg from earlier stages (Chaudhary et al., 2019). Thus, it is formally possible that membrane-tethered Wg could serve as a global PCP cue in these experiments, even if not secreted. In the present study, we use both somatic CRISPR and RNAi to greatly reduce or eliminate wg function altogether. We note, for example, that our wg RNAi phenotypes resemble those caused by inducing apoptosis in wg-expressing cells and thus likely represent very strong loss-of-function phenotypes, yet we also observed no PCP phenotypes. Taken together, the results presented in the present study are complementary and consistent with those presented by Yu et al. (2020): both studies fail to confirm the hypothesis that Wg and Wnt4 are redundantly required for global PCP orientation in the Drosophila wing, and both studies provide accumulate additional evidence that Wnt ligands in general, whether singly or jointly, are not

required for PCP orientation in the *Drosophila* wing in an otherwise wild-type background.

A Note on Somatic CRISPR versus RNAi

A number of studies have demonstrated that *in vivo* somatic CRISPR is an effective and scalable approach for loss-of-function studies (Poe et al., 2019; Port and Bullock, 2016; Port et al., 2020), and the results in the present study further demonstrate that this approach can also be applied to efficiently perform multiplexed somatic knockouts in a tissue-specific manner. Our results also illustrate a known caveat of the approach that may be especially relevant for secreted factors: somatic CRISPR leads to mosaic tissues that contain a mixture of wild-type and mutant cells. Although the proportion of mutant cells can be increased by using multiple sgRNAs (Port and Bullock, 2016), it is unlikely to ever reach 100%, and variability between individuals is likely to remain an issue. For secreted

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proteins such as Wg, even small numbers of remaining cells that secrete wild-type protein may affect the interpretation of results. In vivo RNAi, on the other hand, tends to be far more uniform within and between individuals, and in those cases in which particularly effective reagents are available, such as those used in the present study (compare wg loss-of-function phenotypes in Figures 2 and 3), this can result in stronger tissue-level phenotypes than somatic CRISPR. Thus, the advent of somatic CRISPR does not obviate the power of in vivo RNAi for functional studies.

STAR*METHODS

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SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j. celrep.2020.108121.

ACKNOWLEDGMENTS

We thank Fillip Port for providing pCFD6-evi flies; Marco Milan and Lara Barrios for providing combined RNAi lines targeting wg, wnt4, and wnt6; Rich Binari for assistance with fly work; Justin Bosch for assistance designing pCFD4FLPOUT; and Pedro Saavedra, Stephanie Mohr, Sudhir Tattikota, and members of the Perrimon lab for critical feedback. B.E.-C. received funding from the Charles A. King Postdoctoral Research Fellowship program, and the Perrimon lab received funding from NIH Office of Research Infrastructure Programs (ORIP) grant R24 OD26435 and the Howard Hughes Medical Institute (HHMI).

AUTHOR CONTRIBUTIONS

B.E.-C. designed and performed experiments, generated reagents, and wrote the manuscript. T.C. performed gut imaging experiments. E.V. generated reagents. N.P. designed experiments and edited the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: June 8, 2020 Revised: August 4, 2020 Accepted: August 18, 2020 Published: September 8, 2020

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STAR***METHODS**

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Rabbit anti-GFP, Alexa 488 conjugate	Molecular Probes	(Molecular Probes Cat# A-21311, RRID:AB_221477)
Anti-Digoxigenin-AP, Fab fragments	Sigma Aldrich	(Sigma-Aldrich Cat# 11093274910, RRID:AB_2734716)
Experimental Models: Cell Lines	<u>'</u>	
Drosophila melanogaster S2R+ cells pMK33/Cas9	Perrimon lab (Viswanatha et al., 2018)	pMK33/Cas9
Experimental Models: Organisms/Strains		
Drosophila melanogaster nub-Gal4 ; UAS:Cas9.P2	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 67086; RRID:BDSC_67086)
Drosophila melanogaster UAS:dcr2 ; nub-Gal4	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 25754; RRID:BDSC_25754)
Drosophila melanogaster pnr-Gal4 / TM3	Perrimon lab stock	N/A
Drosophila melanogaster elav-Gal4 ; UAS:Cas9.P2	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 67073; RRID:BDSC_67073)
Drosophila melanogaster tub::Gal4DBD, UAS:EGFP	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 60298; RRID:BDSC_60298)
Drosophila melanogaster tub::VP16[AD], UAS:EYFP	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 60294; RRID:BDSC_60294)
Drosophila melanogaster wnt4 CRIMIC	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 67449; RRID:BDSC_67449)
Drosophila melanogaster "GTRACE" GTRACE: UAS-RedStinger, UAS-Flp1.D, Ubi (FRT.Stop)Stinger/CyO	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 28280; RRID:BDSC_28280)
Drosophila melanogaster GFP RNAi	Perrimon lab stock	HMS00314
Drosophila melanogaster pCFD6-wntless2X	Fillip Port & Simon Bullock	N/A
Drosophila melanogaster wg RNAi	Vienna Drosophila Resource Center (VDRC)	VDRC #104579 (KK collection)
Drosophila melanogaster wnt4 RNAi	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 29442; RRID:BDSC_29442)
Drosophila melanogaster wnt6 RNAi	Vienna Drosophila Resource Center (VDRC)	VDRC #26669 (GD collection)
Drosophila melanogaster w ;; pCFD6-wg2X	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wg2x-wnt22x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wg2x-wnt42x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wg2x-wnt52x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wg2x-wnt62x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wg2x-wntD2x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wg2x-wnt102x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wnt42x-wnt62x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wnt42x-wnt102x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wnt62x-wnt102x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wnt42x-wnt62x ; pCFD6-wg2x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-wnt22x-wnt62x ; pCFD6-wg2x	This study (see Table S1)	N/A
Drosophila melanogaster w ; pCFD6-intergenic control	This study (see Table S1)	N/A
		(Continued on next page

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Continued		
REAGENT or RESOURCE	SOURCE	IDENTIFIER
Drosophila melanogaster w ; pCFD6-fz	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz2	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz3	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz4	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz-fz2	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz-fz3	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz-fz4	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz2-fz3	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz2-fz4	This study (see Table S1)	N/A
Drosophila melanogaster yv ; pCFD4FLPOUT-fz3-fz4	This study (see Table S1)	N/A
Drosophila melanogaster wgT2A-p65	This study (see Table S1)	N/A
Drosophila melanogaster wnt2T2A-p65	This study (see Table S1)	N/A
Drosophila melanogaster wnt4T2A-p65	This study (see Table S1)	N/A
Drosophila melanogaster y v wnt5T2A-p65	This study (see Table S1)	N/A
Drosophila melanogaster wnt6T2A-p65	This study (see Table S1)	N/A
Drosophila melanogaster wntDT2A-p65	This study (see Table S1)	N/A
Drosophila melanogaster wnt10T2A-Gal4DBD	This study (see Table S1)	N/A
Drosophila melanogaster wgT2A-Gal4DBD	This study (see Table S1)	N/A
Drosophila melanogaster UAS:2xEGFP	Bloomington Drosophila Stock Center (BDSC)	(BDSC Cat# 60293; RRID:BDSC_60293)
Drosophila melanogaster wnt4T2A-Gal4	This study (see Table S1)	N/A
Oligonucleotides		
PCR primers	See Table S2	
In situ RNA probes	See Table S2	
Recombinant DNA		
pCFD6	Addgene	73915
pBPp65ADZpUw (split-Gal4 p65)	Addgene	26234
pBPZpGAL4DBDUw (split-Gal4 Gal4DBD)	Addgene	26233
pHD-dsRed-attP	Addgene	51019
pCFD4FLPOUT	This study	N/A
Software and Algorithms		
Zen	www.zeiss.com	(ZEN Digital Imaging for Light Microscopy RRID:SCR_013672)
Adobe Photoshop vCS6	www.adobe.com	(Adobe Photoshop; RRID:SCR_014199)
Adobe Illustrator v23.0.4	www.adobe.com	Adobe Illustrator; RRID:SCR_010279
Helicon Focus	www.heliconsoft.com	(Helicon Focus; RRID:SCR_014462)

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Norbert Perrimon (perrimon@genetics.med.harvard.edu)

Materials Availability

All unique/stable reagents generated in this study are available from the Lead Contact without restriction.

Data and Code Availability

This study did not generate any unique datasets or code.

Report



EXPERIMENTAL MODEL AND SUBJECT DETAILS

Fly strains

Drosophila melanogaster stocks were maintained and crossed on standard laboratory cornmeal food, and all experiments were conducted at 25°C. The following previously described stocks were used: nub-Gal4; UAS:Cas9.P2 (BL67086), UAS:dcr2; nub-Gal4 (BL25754), pnr-Gal4 / TM3 (Perrimon lab stock), elav-Gal4; UAS:Cas9.P2 (BL67073), split-Gal4 "p65 tester line": tub::Gal4DBD, UAS:EGFP (BL60298), split-gal4 "Gal4DBD tester line": tub::VP16[AD], UAS:EYFP (BL60294), wnt4 CRIMIC "trojan Gal4" (BL67449), GTRACE: UAS-RedStinger, UAS-Flp1.D, Ubi (FRT.Stop)Stinger/CyO (BL28280), GFP RNAi: HMS00314 (Perrimon lab stock), pCFD6-wntless^{2X} – gift of F. Port (Port and Bullock, 2016), wg RNAi – dsRNA, VDRC (KK104579), wnt4 RNAi – dsRNA, VDRC (BL29442), wnt6 RNAi – dsRNA, VDRC (GD26669), UAS:2xEGFP (BL60293)

Drosophila cell lines

Cas9-expressing S2R+ cell lines, used to estimate sgRNA efficiency, are described in Viswanatha et al. (2018).

METHOD DETAILS

sgRNAs construction

For each Wnt ligand, five candidate sgRNAs were designed using the *Drosophila Resource Screening Center Find CRISPR v2* online tool (https://www.flyrnai.org/crispr2/). These five candidate sgRNAs were cloned into U6:3-expression vector pCFD3 (Port et al., 2014), and tested for cutting efficiency in S2R+ cells engineered to express Cas9 (Viswanatha et al., 2018), followed by T7 endonuclease activity three days after transfection. Cutting efficiency was estimated by quantifying band intensity via ImageJ, and the top two scoring sgRNAs were selected for usage in this study. *In vivo* guides were cloned into pCFD6, which expresses multiple sgRNAs under UASt control, each flanked by tRNA sequences (Port and Bullock, 2016). The sgRNA sequences used in this study are shown in Table S1. For single gene knock-out experiments, two sgRNAs were cloned into the pCFD6 backbone and inserted into the attP2 landing site on the third chromosome, and for double gene knock-out experiments, four sgRNAs were cloned into pCFD6, two sgRNAs per target gene, and inserted into the attP40 site on the second chromosome, using standard PhiC31 recombination. For triple-CRISPR experiments, single-target sgRNA constructs on the third chromosome (attP2) were combined with double-target sgRNA constructs on the second chromosome (attP40) using genetic crosses.

Construction of pCFD4FLPOUT vector (used in Figure S6)

For pairwise *frizzled* knock-out experiments, we created a modified pCFD4 variant (Port et al., 2014) conceptually based on the Co-inFLP approach described in Bosch et al. (2015). In this two-guide sgRNA vector, each sgRNA is flanked by a distinct fluorescent reporter, and interlocking FRT/FRT3 sites, for a final orientation of *FRT-3xP3-GFP-U6:1-sgRNA1-FRT3-FRT-U6:3-sgRNA2-3xP3-mCherry-FRT3*. This fragment was synthesized by GENEWIZ, Inc. (South Plainfield, NJ) and subcloned into the pCFD4 backbone using Gibson cloning.

We reasoned that this plasmid would allow us to create a single transgenic fly line expressing two guides, and to subsequently generate each of the two single sgRNA components by crossing this "parent" line to a source of *hsFlp* and screening the offspring for those that underwent each of the two possible recombination events (FRT versus FRT3), which would express either GFP and mCherry. We validated that this construct successfully expressed two sgRNAs targeting *ebony* and *forked*, which both produce visible adult phenotypes (not shown). We also validated that the FLP-FRT procedure could be used to successfully generate two stable single-sgRNA lines from one "parent" double-sgRNA line. This construct may be of interest for those wishing to screen large numbers of pairs of genes, and subsequently perform single gene studies on a smaller subset of those genes.

T7 endonuclease assay

To test for *in vivo* target cleavage of Wnt genes, pCFD6 males were crossed to unmated *elav-Gal4*; *UAS:Cas9.P2* females, and the adult heads of their F1 offspring were collected for genomic DNA. \sim 5-10 heads were collected on dry ice and genomic DNA was extracted by physical homogenization in DNA extraction buffer (10mM Tris-Cl pH 8.2, 1mM EDTA, 25mM NaCL, 200 μ g/mL Proteinase K), followed by incubation at 37°C for 20-30 minutes, then boiling at 98°C for \sim 90 s to inactivate Proteinase K. 1 μ L of genomic DNA was used as template in a 20 μ L PCR reaction using primers described in Table S2 to amplify regions surrounding the sgRNA target site. PCP products were used as template for T7 endonuclease assay (New England BioLabs), following manufacturer instructions, and digested products were visualized on a 2% agarose gel.

Generation of split-Gal4 knock-in reporter lines

To create split-Gal4 donor vectors (both p65 and Gal4DBD), p65 and Gal4DBD amplified from pBPp65ADZpUw (Addgene 26234) and pBPZpGAL4DBDUw (Addgene 26233), respectively, and subcloned into pHD-dsRed-attP (Addgene 51019) (Gratz et al., 2014; Pfeiffer et al., 2010). For each Wnt gene, homology arms of approximately 1 kb flanking the sgRNA were designed to create an in-frame fusion between the first exon of the target gene and the T2A sequence of the donor construct, and were cloned into donor vector using Gibson cloning.





Donor constructs and sgRNAs (in pCFD3) were co-injected into yv;; nanos: Cas9 flies. Injected offspring were backcrossed to balancer stocks on the X, II, or III chromosome depending on the target gene, and screened for RFP+ eyes. Multiple transformants were recovered for both p65 and Gal4DBD knock-ins of each Wnt gene, with the exception of wnt4 and wnt10, for which only p65 and Gal4DBD lines were recovered, respectively. Expression patterns were ascertained by crossing to ubiquitous split-Gal4 "tester lines" (BL60298 and BL60294), which ubiquitously express either VP16 or Gal4DBD together with a UAS-driven reporter. For every Wntfamily knock-in created, multiple transformants gave indistinguishable expression patterns, regardless of whether p65 or Gal4DBD was knocked-in. PCR analyses confirmed that each line was inserted in the predicted location, with the exception of wntD, which we were unable to conclusively validate via PCR. We note that all wntD knock-in lines, independently generated, and both with p65 and Gal4DBD, recapitulate indistinguishable expression patterns from one another.

Generation of additional Wnt4 reporter lines

An additional wnt4 reporter line was generated using a "universal donor" homology-independent knock-in strategy (Bosch et al., 2020). Briefly, a universal T2A-Gal4, 3xP3-dsRed donor was injected into yv;; nos::Cas9 embryos along with two sgRNAs: one to linearize the donor construct and one to target in the last exon of wnt4. These injected flies were crossed to a w; Sp/CyO; UAS-GFP balancer line and screened for RFP and GFP expression. Of 28 fertile injected flies, six were RFP+, of which five failed to drive UAS::GFPexpression, indicating they were likely out-of-frame and/or inserted at the wrong location. The remaining one RFP+ line drove UAS:GFP in an expression pattern indistinguishable from the two other wnt4 lines tested in this study.

Antibody staining and imaging

Tissues were dissected in cold PBS, fixed for 20-30 minutes in 4% paraformaldehyde in PBS, and then stained following standard immunhistochemical protocols. Tissues were stained with rabbit anti-GFP conjugated to Alexa 488 (Molecular Probes, 1:400) and DAPI (1:1000), and imaged using either a Zeiss LSM 780 or an Olympus IX83 confocal microscope, both part of the Microscopy Resources of the North Quad (MicRoN) facility at Harvard Medical School. Maximum intensity projections are shown for all tissues except the proventriculus, for which single z-slices are also shown. Adult wings were mounted in a 1:1 mixture of Permount (Fisher) and xylenes, and imaged using brightfield optics on a Zeiss Axioskop 2. Nota were imaged using a Zeiss AxioZoom, and z stacks were compiled into a single image using Helicon Focus software.

In situ hybridization

RNA probes against wnt4 and Gal4 were generated by amplifying T7-containing PCR fragments using primers shown in Table S2, which contain 5' and 3' linker sequences that allow directional T7 sites to be added in a second PCR reaction, for either antisense or sense probe synthesis. Digoxigenin-labeled RNA probes were synthesized and purified using T7 5X Megascript kit (Invitrogen). L3 larvae were bisected, inverted, and fixed overnight in 4% paraformaldehyde, then sequentially dehydrated into methanol, stored at -20°C for three days, washed twice with ethanol, treated with 1:1 ethanol:xylenes for one hour, then rehydrated, washed repeatedly with PBS + 0.3% Triton-X, and hybridized overnight at 58°C with RNA probes at 100 ng/μL in RNA hybridization buffer (50% formamide, 5X SSC, 0.1mg/mL heparin, 1.0% salmon sperm single stranded DNA, 0.1% Tween-20 in RNase-free water.) Larvae were then washed repeatedly in hybridization buffer, a series of increasingly dilute hybridization buffer, multiple washes of PBS + 0.3% Trixon-X, and incubated overnight with anti-digoxigenin-AP Fab fragments (Roche, 1:1000). Finally, larvae were washed repeatedly with PBS + 0.3% Triton-X, and signal was detected using 5-bromo-4-chloro-3-indolyl-phosphate/nitro blue tetrazolium (Promega) following manufacturer's protocol, then washed in ethanol, mounted in vectashield, and imaged using brightfield optics on a Zeiss Axioskop 2.

QUANTIFICATION AND STATISTICAL ANALYSIS

This study did not use quantification or statistical analysis.