Dally is not essential for Dpp spreading or internalization but for Dpp stability by antagonizing Tkv-mediated Dpp internalization

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Abstract

Dpp/BMP acts as a morphogen to provide positional information in the Drosophila wing disc. Key cell-surface molecules to control Dpp morphogen gradient and signaling are heparan sulfate proteoglycans (HSPGs). In the wing discs, two HSPGs, the glypicans Division abnormally delayed (Dally) and Dally-like (Dip) have been suggested to act redundantly to control these processes through direct interaction of their heparan sulfate (HS) chains with Dpp. Based on this assumption, a number of models on how glypicans control Dpp gradient formation and signaling have been proposed, including facilitating or hindering Dpp spreading, stabilizing Dpp on the cell surface, or recycling Dpp. However, how distinct HSPGs act remains largely unknown. Here we generate genome-engineering platforms for the two HSPGs and find that only Dally is critical for Dpp gradient formation and signaling through interaction of its core protein with Dpp. However, we find that this interaction is not sufficient and the HS chains of Dally are essential for Dpp gradient formation and signaling, but surprisingly, largely without interacting with Dpp. We provide evidence that the HS chains of Dally are not essential for spreading or recycling of Dpp but for stabilizing Dpp on the cell surface by antagonizing Dpp internalization through Tkv. These results provide new insights into how distinct HSPGs control morphogen gradient formation and signaling during development.

Introduction

Morphogens are signaling molecules secreted from localized source cells and control cell fates in a concentration dependent manner during development (Ashe and Briscoe, 2006; Rogers and Schier, 2011; Stapornwongkul and Vincent, 2021; Tabata, 2001). Amongst them, Decapentaplegic (Dpp), the vertebrate BMP2/4 homolog is the first validated secreted morphogen identified in Drosophila. Since its discovery, the role of Dpp signaling in the Drosophila wing imaginal disc (the larval precursor of the adult wing) has served as an excellent model to investigate how morphogen gradients form and act (Affolter and Basler, 2007; Kicheva and Gonzalez-Gaitan, 2008; Matsuda et al., 2016). In the wing disc, Dpp is expressed in an anterior stripe of cells along the anterior-posterior compartment boundary to control nested target gene expression and growth (Fig. 1A). Dpp binds to the type I receptor Tkv and the type II receptor Punt to phosphorylate the transcription factor Mad. Phosphorylated Mad (pMad) forms a complex with Co-Smad (Medea) to accumulate into the nucleus and activate or inhibit target gene transcription. Interestingly, the majority of Dpp target genes are repressed by a transcriptional repressor Brinker (Brk) while pMad directly represses brk expression (Campbell and Tomlinson, 1999; Jazwinska et al., 1999; Minami et al., 1999). Thus, the pMad gradient regulates nested target gene expression mainly through generating an inverse gradient of Brk (Muller et al., 2003) (Fig. 1A). The nested target gene expression patterns are thought to position the future adult wing veins such as L2 and L5 (Cook et al., 2004; Sturtevant et al., 1997) (Fig. 1A). In addition to patterning, Dpp signaling controls growth of the wing through repressing Brk as the strong proliferation defects of dpp mutant wing discs can be reversed by removing brk (Campbell and Tomlinson, 1999; Schwank et al., 2008).

While endogenous Dpp is indeed distributed in a graded manner (Matsuda et al., 2021), how the gradient arises is highly controversial (Akiyama and Gibson, 2015; Kicheva and Gonzalez-Gaitan, 2008; Matsuda et al., 2016). A class of molecules that control gradient formation of various morphogens are glypicans, GPI-anchored heparan sulfate proteoglycans (HSPGs) consisting of a core protein and covalently attached heparan sulfate (HS) chains (Lin and Perrimon, 2002; Nybakken and Perrimon, 2002) (Yan and Lin, 2009). In Drosophila, two glypicans, Division abnormally delayed (Dally) (Nakato et al., 1995) and Dally-like (Dip) (Khare and Baumgartner, 2000), are thought to redundantly act as co-receptors (Kuo et al., 2010) to regulate Dpp morphogen gradient formation (Belenkaya et al., 2004; Fujise et al., 2003) and control Dpp gradient scaling to tissue size (Romanova-Michaelides et al., 2015; Zhu et al., 2020).

The role of glypicans in Dpp signaling is dependent on HS chains. A variety of genes required for HS chain biosynthesis such as sulfateless (sfl) and a family of Ext genes (sister of tout-velu (soTV), brother of tout-velu (botv), and tout-velu (ttv)) (Bornemann et al., 2004; Han et al., 2004;
Takei et al., 2004) have been shown to be involved in Dpp distribution and signaling. Given that Dpp binds to heparin (Akiyama et al., 2008), the interaction of the HS chains of glypicans with Dpp is thought to be essential for their function.

Several, not necessarily mutually exclusive, models have been proposed on how glypicans control Dpp morphogen gradient formation. First, glypicans are suggested to be essential for Dpp spreading by handing it over from one cell to the next cell in a “bucket brigade” manner (so-called restricted diffusion model) (Kerszberg and Wolpert, 1998) (Fig. 1B). Consistently, FRAP assays using overexpression of GFP-Dpp revealed a low diffusion coefficient of Dpp (D=0.1μm²/s) (Kicheva et al., 2007). Indeed, pMad signal and Dpp distribution have been shown to be reduced not only in glycan mutant clones but also in wild type cells located adjacent to the clone and distally to the Dpp source (so-called “shadow” effect), indicating that Dpp spreading requires glypicans (Belenkaya et al., 2004).

Second, and in contrast to the first model, glypicans have been proposed to interfere with Dpp spreading (Muller et al., 2013) (Fig. 1C). The “hindered diffusion” model postulates that Dpp spreads freely and transient interaction of Dpp with cell surface molecules such as glypicans hinders Dpp spreading (Muller et al., 2013).

Third, glypicans have been proposed to stabilize Dpp on the cell surface (Akiyama et al., 2008) (Fig. 1D). In a previous study, Dpp lacking the N-terminal HS binding site was less stable than wild type Dpp (Akiyama et al., 2008), indicating that Dally regulates Dpp stability through direct interactions of HS chains with Dpp. Genetic analyses indicated that Dally competes with Tkv for Dpp binding to antagonize Tkv-mediated Dpp internalization (Akiyama et al., 2008).

Fourth, glypicans are proposed to control recycling and re-secretion of Dpp, which is critical for extracellular Dpp gradient scaling (Romanova-Michaelides et al., 2022), a phenomenon in which Dpp gradient expands to match the growing wing disc size during development (Fig. 1E). Surprisingly, although receptor-mediated endocytosis is thought to be a mechanism to clear morphogens from the extracellular space (Fig. 1D), the same study showed that Dpp internalization was not affected in tkv mutant clones, indicating that Dally, but not Tkv, mediates Dpp internalization (Romanova-Michaelides et al., 2022).

Finally, a recent attempt to create a synthetic morphogen system using free diffusing GFP as a morphogen raises alternative possibilities (Stapornwongkul et al., 2020). When dpp was replaced with GFP, and anti-GFP nanobodies fused to Tkv and Punt that can activate pMad signal upon GFP binding were introduced, rescue of severe dpp mutant phenotypes was not perfect but improved by further introducing GPI-anchored non-signaling receptors (corresponding to glypicans). Supported by mathematical modeling, this study suggests that glypicans may contribute by blocking morphogen leakage and assisting morphogen spreading likely via their GPI-anchors (Stapornwongkul et al., 2020).

Previously, we generated a platform to introduce epitope tags and/or mutations in the dpp locus and protein binder tools to directly manipulate endogenous Dpp spreading (Matsuda et al., 2021). By visualizing and manipulating the endogenous Dpp morphogen gradient, we found that Dpp spreading is critical for posterior patterning and growth but largely dispensable for anterior patterning and growth (Matsuda et al., 2021). Although the requirements for Dpp spreading were thus less important than previously thought, endogenous Dpp gradient was easily visualized in the wing discs and Dpp spreading was still critical for posterior patterning and growth (Matsuda et al., 2021). Thus, wing imaginal discs serve as an excellent model to study Dpp morphogen gradient formation at physiological conditions.

In this study, to better understand how glypicans are involved in Dpp morphogen gradient formation and signaling, we generated genome-engineering platforms to manipulate both Drosophila glypicans, Dally and Dlp. Although the two glypicans are thought to act redundantly, we first find that only Dally is critical for Dpp gradient formation through interaction of its core
protein with Dpp. Surprisingly, we found that, although the HS chains of Dally are essential for Dally's function, dpp mutants that lack the interaction with HS chains display minor phenotypes, indicating that the interaction of HS chains of Dally with Dpp is largely dispensable for Dpp gradient formation and signaling. We provide evidence that the HS chains of Dally are not essential for Dpp internalization or spreading but rather stabilize Dpp on the cell surface by antagonizing Tkv-mediated internalization of Dpp. These results provide new insights into how glypicans control dispersal and signaling of distinct morphogens during development.

**Results**

**Dally but not Dlp interacts with Dpp**

 Previous genetic studies suggested that both Dally and Dlp are involved in extracellular Dpp distribution and signaling through the interaction of Dpp with HS chains (Belenkaya et al., 2004). However, the relative contribution of each glypican remained unclear. To address this, we first expressed *dally* or *dlp* in the dorsal compartment of the wing disc using ap-Gal4 to compare their ability to interact with endogenous Dpp and activate pMad signal *in vivo*. To visualize endogenous extracellular Dpp distribution, we utilized a functional *Ollas-dpp* allele (Bauer Milena, 2022). Since Dpp distribution and pMad signal are similar between the dorsal and ventral compartments in discs expressing the *Ollas-dpp* allele (Fig. S1), the ventral compartment serves as an internal control in this assay. We found that expression of *dally* in the dorsal compartment increased HS chain levels, and expanded the extracellular Dpp distribution and pMad gradient compared to the control ventral compartment (Fig. 2A, B, E, F). We found that, in contrast to the broad extracellular Ollas-Dpp accumulation, the pMad signal dropped to background levels at the peripheral region (Fig. 2E, F), indicating that Dpp signal activation by Dally is inefficient likely due to sequestration of Dpp by overexpressed Dally. In contrast, expression of *dlp* using ap-Gal4 increased HS chain levels but rather reduced extracellular Ollas-Dpp distribution and pMad gradient in the dorsal compartment compared to the control ventral compartment (Fig. 2C, D, G, H), likely because ectopic Dlp expression sequestered HS (Fig. 2C) and blocked HS modification of Dally. Consistent with a previous report (Han et al., 2005), Dlp expression, in the dorsal compartment efficiently expanded extracellular Wg distribution compared to Dally expression (Fig. S2). These results suggest that, although both Dally and Dlp are modified by HS chains, only Dally interacts with Dpp.

**Dally, but not Dlp, is required for the Dpp activity gradient**

To compare the requirement of Dally and Dlp for Dpp signaling gradient formation, we then generated null mutants for *dally* and *dlp* (*dally*KO and *dlp*KO respectively) by replacing the first exon of each gene by an attP cassette via genome engineering (Fig. S3). Although there is no good anti-Dally antibody, defects in pMad signal in *dally*KO wing discs were similar to that in *dally*Mh32 (a *dally* null allele (Franch-Marro et al., 2005)) wing discs (Fig. 3A-C, I) and *dally*KO adult wings showed similar phenotypes to *dally*Mh32 adult wings (Fig. 3E-G, J). Immunostaining for Dlp revealed that Dlp expression is completely lost in *dlp*KO wing disc (Fig. S4). Thus, we confirmed that both *dally*KO and *dlp*KO are null alleles. These null alleles also serve as a platform to insert epitope tags or modified versions of *dally* and *dlp* in the endogenous genomic locus via the attP cassette.

We then compared the two null alleles and found that, while severely shrunk in *dally*KO wing discs, the pMad signal was not affected in *dlp*KO wing discs (Fig. 3C, D, I). In rare-occurring mutant adult wings, patterning defects linked to defects in Dpp signaling (truncated L5) were seen only in *dally*KO mutants (Fig. 3G). In contrast, the distance of the distal tips of L3 and L4 veins was reduced in *dlp*KO mutants, which is indicative of Hh signaling defects (see arrows in Fig. 3H). Although both *dally*KO and *dlp*KO adult wings are smaller than controls (Fig. 3J), only *dally*KO adult wings are narrower along the A-P axis, consistent with the role of Dpp along this axis (Fig. 3K).

It has been shown that *dally* and *dlp* display distinct expression patterns in the wing disc since *dally* expression is controlled by Dpp signaling (Fujise et al., 2003) and *dlp* expression is controlled by Wg signaling (Han et al., 2005). Thus, the differential expression pattern of Dally...
and Dlp may underlie the differential requirements of glypicans in Dpp signaling. To address this, we inserted dlp in dally locus using the attP site in dally\textsuperscript{KO} allele to express dlp in the dally expression pattern. As a control, YFP-dally inserted in dally\textsuperscript{KO} rescued pMad defects (Fig. 3L-N, V) and adult wing defects of dally\textsuperscript{KO} mutants (Fig. 3P-R, W). In contrast, 3xHA-dlp insertion into dally\textsuperscript{KO} failed to rescue pMad defects (Fig. 3M, O, V) or adult wing defects in dally\textsuperscript{KO} mutants (Fig. 3Q, S, W), although 3xHA-dlp insertion into dlp\textsuperscript{KO} restored adult wing growth in dlp\textsuperscript{KO} mutants (Fig. 3T, U, W). Taken together, these results indicate that dally, but not dlp, is required for Dpp signaling gradient formation.

**Interaction of the core protein of Dally with Dpp**

How can only Dally be involved in Dpp signaling although both HSPGs are modified with HS chains (Fig. 2A, C)? Dally has been shown to interact with Dpp in vitro not only through HS chains but also through the core protein (Kirkpatrick et al., 2006). Thus, we wondered whether the core protein of Dally can interact with Dpp in the wing disc to provide ligand specificity. To test this, we expressed dally or dally\textsuperscript{ΔHS}, a dally mutant lacking all HS chain attachment sites (Kirkpatrick et al., 2006) using ap-Gal4 to compare the ability of Dally and Dally\textsuperscript{ΔHS} to interact with Dpp and activate pMad signal. Upon dally expression in the dorsal compartment, HS chains, extracellular Dpp and pMad signal increased as compared to the control, ventral compartment (Fig. 4A-C, G, J). In contrast, upon dally\textsuperscript{ΔHS} expression, HS chains rather decreased (Fig. 4D), probably because the core protein of Dally sequesters the factors required for synthesis of HS chains. Under this condition, extracellular Dpp still increased but pMad signal rather decreased (Fig. 4E, F, H, K). These results indicate that the core protein of Dally can interact with Dpp; however, this interaction is not sufficient to activate pMad signal but requires the presence of the HS chains.

To address the relative contribution of the interaction of core protein and HS chains of Dally with Dpp, we measured the relative Dpp accumulation against Dally or Dally\textsuperscript{ΔHS} expression (Fig. 4I), since Dally\textsuperscript{ΔHS} expression was higher than Dally expression (Fig. 4B, E'). We found that the relative Dpp accumulation upon dally\textsuperscript{ΔHS} expression is only slightly reduced compared to that upon dally expression (Fig. 4I), indicating a major contribution of the core protein in the interaction between Dally and Dpp in the wing disc.

**HS chains of Dally are critical for Dpp distribution and signaling**

The relatively minor contribution of HS chains of Dally for Dpp interaction raises questions on how HS chains of Dally are involved in Dpp distribution and signaling. To address this, we inserted YFP-dally\textsuperscript{ΔHS}, a dally mutant allele lacking HS chain modification (Kirkpatrick et al., 2006), in the attP site of the dally\textsuperscript{KO} allele. We found that both pMad signaling (Fig. 4L-O) and extracellular Dpp distribution (Fig. 4P-S) were severely affected in the YFP-dally\textsuperscript{ΔHS} wing discs resembling the phenotypes of dally\textsuperscript{KO} mutants. Consistently, the YFP-dally\textsuperscript{ΔHS} adult wings mimicked patterning and growth defects of dally\textsuperscript{KO} adult wings (Fig. 4T-V). These results show that the interaction of the core protein of Dally with Dpp is not sufficient and that the HS chains of Dally are critical for Dpp distribution and signaling under physiological conditions.

**HS chains of Dally control Dpp distribution and signaling largely independent of interaction with Dpp**

How do HS chains of Dally control Dpp distribution and signaling gradient? We first asked whether a direct interaction of the HS chains of Dally and Dpp is critical. A previous study showed that 7 basic amino acids at the N-terminus of the Dpp mature domain are crucial for the interaction with heparin but dispensable for interaction with receptors to activate downstream signaling (Akiyama et al., 2008). To test the importance of this interaction in vivo, we generated dpp\textsuperscript{ΔN}, a dpp mutant allele lacking the basic amino acid stretch. We found that the dpp\textsuperscript{ΔN} allele is embryonic lethal, likely because the basic amino acids overlap with a collagen binding site and the Dpp-Collagen IV interaction is important for Dpp gradient formation in the early embryo (Wang et al., 2008). To bypass this embryonic lethality without affecting wing disc development, we utilized a previously reported transgene called JAX, which contains the genomic region of dpp (JAX) critical for the early embryogenesis (Hoffmann and...
Goodman, 1987) but does not rescue the wing phenotypes of dpp mutants (Fig. S5). Indeed, the early lethality of homozygous HA-dppΔN was greatly rescued in the presence of this transgene (JAX; HA-dppΔN), thus allowing to investigate the requirement of the interaction of Dpp with the HS chains of Dally in later stages. Surprisingly, and in contrast to dallyKO or dallyΔHS flies (Fig. 3, 4), we found that JAX; HA-dppΔN allele showed only minor defects in Dpp distribution and signaling in wing discs (Fig. 5A-D) and in patterning and growth in adult wings (Fig. 5E-G). Indeed, while dallyKO and dallyΔHS adult wings are 25-30% smaller than control wings (Fig. 3J, W, 4V), JAX; HA-dppΔN adult wings are only 6.4% smaller than control wings (Fig.5G).

The slight but consistent reduction of relative Dpp accumulation upon DallyΔHS expression compared to that upon Dally expression indicates minor interaction of HS chains of Dally with Dpp (Fig. 4I). In contrast, the relative DppΔN accumulation upon DallyΔHS expression in JAX; dppΔN was comparable to that upon DallyΔHS expression in JAX; dppΔN (Fig. 5H-J). This indicates that, consistent with a previous report (Akiyama et al., 2008), DppΔN lacks the ability to interact with HS chains of Dally.

Using the same experimental setup, we then compared the ability of Dally and DallyΔHS to activate Dpp signal in JAX; dppΔN, in which HS chains-Dpp interaction is lost (Fig. 5J). We found that Dally expression expanded Dpp signaling in JAX; dppΔN (Fig. 5H*, K), indicating that Dally can activate Dpp signaling independent of interaction of HS chains of Dally with Dpp. In contrast, DallyΔHS expression failed to activate Dpp signaling in JAX; dppΔN (Fig. 5I*, L), indicating that co-receptor function of HS chains of Dally does not require direct interaction with Dpp. Taken together, these results suggest that, although HS chains of Dally are essential for Dpp distribution and signaling, a direct interaction of HS chains of Dally with Dpp is largely dispensable.

HS chains of Dally stabilize Dpp by antagonizing Tkv-mediated Dpp internalization

So far, all the models on how glypicans control Dpp distribution and signaling are based on direct interaction of the HS chains with Dpp (Fig. 1). How can the HS chains of Dally control Dpp distribution and signaling without direct interaction with Dpp? Among the models, we wondered whether Tkv is a factor through which HS chains of Dally can act (Fig 1D), although HS chains are thought to act through Dpp in the original model (Akiyama et al., 2008).

We first re-investigated the role of Tkv. According to the model (Akiyama et al., 2008) (Fig. 1D), Tkv is the Dpp receptor on the cell surface that internalizes and degrades Dpp in addition to activating pMad signal. In contrast, a recent study proposed that it is Dally, but not Tkv, that internalizes Dpp (Romanova-Michaelides et al., 2022). To address this discrepancy, we first depleted tkv by RNAi using ap-Gal4 and visualized endogenous extracellular Dpp distribution. If Tkv is required for Dpp internalization, extracellular Dpp distribution should increase by removing Tkv. Indeed, we found that extracellular Dpp increased and expanded in the dorsal compartment (Fig. 6A, C), indicating that Tkv is required to internalize Dpp.

To address whether Dally antagonizes Tkv-mediated Dpp internalization, we asked if the defects in Dpp distribution in daily mutants can be reduced by RNAi-mediated depletion of tkv. We found that the extracellular Dpp gradient was expanded in the dorsal compartment (Fig. 6B, D), indicating that Dally antagonizes Tkv-mediated Dpp internalization. Expanded Dpp distribution in daily mutants upon blocking Dpp internalization also suggest that Dally is largely dispensable for Dpp spreading, and that there are factors other than Tkv and Dally that bind to Dpp on the cell surface. We then asked if Dally contributes to internalization of Dpp by comparing extracellular Dpp levels in the dorsal compartment of ap>tkv RNAi/wing discs (Fig. 6A) and that of ap>tkv RNAi, dallyKO wing discs (Fig. 6B), since extracellular Dpp is removed by Tkv in daily mutants. We found that extracellular Dpp levels do not increase (Fig.6E), indicating that Dally is not essential to internalize Dpp.
Interestingly, when tkv was depleted in the dorsal compartment, pMad was not only lost from the dorsal compartment but was also upregulated in ventral cells along the dorsal-ventral compartment boundary (arrow, Fig. 6F). This observation is consistent with the idea that, upon blocking internalization, Dpp from the dorsal compartment can reach to the ventral compartment to activate pMad through Tkv binding. In contrast, this phenotype was not seen upon depletion of daily by RNAi, although depletion of daily in the dorsal compartment mimicked the pMad defects of daily mutants (Fig 6G).

Cumulatively, these results suggest that Tkv, but not Dally, is essential to internalize Dpp, and support a model, in which Dally stabilizes Dpp on the cell surface by blocking Tkv-mediated Dpp internalization (Akiyama et al., 2008). While HS chains of Dally are thought to compete with Tkv for binding to Dpp in the original model, the minor phenotypes of dppΔHS (Fig. 4, 5) suggest that HS chains act through a different mechanism. To test if the HS chains of Dally act through Tkv, we depleted tkv by RNAi using ap-Gal4 in dailyΔHS mutants. We found that extracellular Dpp gradient expanded in the dorsal compartment (Fig. 6H, I), indicating that the HS chains of Dally antagonize Tkv-mediated Dpp internalization through Tkv.

**Discussion**

HSPGs are essential for Dpp/BMP morphogen gradient formation and signaling in the Drosophila wing disc. Although a variety of models have been proposed, it remains largely unknown how distinct HSPGs control Dpp gradient formation and signaling. In this study, we generated and utilized genome engineering platforms for two HSPGs, daily and dlp, and dpp (Matsuda et al., 2021) to address this question.

**Mechanism of Dally function**

It has been thought that Dally and Dlp act redundantly to control Dpp signaling. However, we demonstrate here that Dally, but not Dlp, is critical for Dpp distribution and signaling through interaction of the core protein of Dally with Dpp (Fig. 2-4). We do not know if the interaction is direct but previous surface plasmon resonance (SPR) and co-immunoprecipitation experiments suggest that this is the case (Kirkpatrick et al., 2006). It has recently been shown that a specific structure, present in the core protein of Dlp but absent in Dally, shields the lipid moiety of Wg and is important for its spreading (McGough et al., 2020). Thus, each glypican appears to acquire ligand specificity through its core protein. In contrast to Wg, Dpp has no lipid modification and is diffusible by nature. Indeed, the expanded Dpp distribution in daily mutants upon blocking Dpp internalization (Fig. 6) does not support the idea that Dally is required for Dpp spreading itself (Belenkaya et al., 2004; Schwank et al., 2011; Stapornwongkul et al., 2020) (Fig. 1B) or that Dally and Tkv are essential for blocking Dpp leakage from the wing epithelia (Stapornwongkul et al., 2020). These results also suggest that there are other cell surface molecules that bind to Dpp on the cell surface. Furthermore, the expanded extracellular Dpp distribution upon daily expression (Fig. 2) suggests that the ability of Dally to hinder Dpp spreading is, if any, mild (Fig. 1C).

Our results thus suggest that Dally is not critical for Dpp spreading and that interaction of Dally with Dpp occurs downstream of Dpp spreading. Despite the interaction of the core protein of Dally with Dpp upon Dally overexpression, this interaction is not sufficient to control Dpp distribution and Dpp signaling at its physiological level, while HS chains of Dally are absolutely essential for both processes (Fig. 4). Since increase of HS chains was not sufficient for interaction with Dpp (Fig. 2C, D), we speculate that HS chains of Dally act downstream of the interaction of the core protein of Dally with Dpp.

Among the proposed models (Fig. 1), our results support a model, in which HS chains of Dally stabilize Dpp on the cell surface by antagonizing Tkv-mediated internalization of Dpp (Akiyama et al., 2008) (Fig. 1D, 7A). Consistently, pulse-chase experiments showed that Dpp is less stable in daily mutants (Akiyama et al., 2008). However, while the original model suggests that HS chains compete with Tkv for Dpp binding (Akiyama et al., 2008) (Fig.1D, 7A), our results suggest that HS chains act largely through Tkv based on two observations.
(Fig. 7B). First, although the dally\textsuperscript{AHs} allele showed severe defects similar to the dally\textsuperscript{XO} allele, the defects of the dpp\textsuperscript{AN} allele were surprisingly mild despite the lack of interaction with HS chains (Fig. 4, 5), indicating that HS chains act largely through other molecules. Second, defects in Dpp distribution in dally\textsuperscript{AHs} were rescued by removing tkv (Fig. 6). These results indicate that HS chains act through Tkv rather than through Dpp to stabilize Dpp on the cell surface (Fig. 7C-E).

It remains unknown how HS chains of Daily stabilize Dpp through Tkv. Since heparin has been shown to bind to BMP receptors (Kanzaki et al., 2008), the HS chains of Daily may directly interact with Tkv to stabilize Dpp on the cell surface. Interestingly, we found that the HS chains of Daily control Dpp signaling also largely independent of interaction with Dpp (Fig. 5). The simplest possibility is that the HS chains of Daily control both Dpp stability and signaling through Tkv (Fig. 7), but it is equally possible that the HS chains of Daily control Dpp signaling through other factors. In the future, it will be also important to address how the interaction of the core protein of Daily with Dpp and functions of HS chains through Tkv are coordinated.

**Mechanisms of Dpp internalization**

Our proposed model is based on the result that Dpp is internalized by Tkv (Fig. 6), consistent with a widely accepted idea that extracellular morphogens are removed by their receptors. However, this result contradicts a recent report proposing that Dally, but not Tkv, internalizes and recycles Dpp (Romanova-Michaelides et al., 2015). How can we explain the discrepancy?

If Tkv is involved in internalization of Dpp, internalized Dpp should decrease upon loss of tkv. Romanova-Michaelides et al. applied a nanobody internalization assay to visualize internalized Dpp (Romanova-Michaelides et al., 2015). In this assay, extracellular GFP-Dpp was first labeled using fluorophore conjugated anti-GFP nanobodies under endocytosis-restricted condition. After allowing internalization of the labelled GFP-Dpp, extracellular anti-GFP nanobodies were then stripped away by acid wash to follow only the internalized Dpp. Using this assay, Romanova-Michaelides et al. surprisingly found that internalization of GFP-Dpp was not affected in tkv mutant clones. However, the experiment was performed under overexpression of GFP-Dpp from the anterior stripe of cells, in which the authors estimate 400 times higher expression than the physiological level. Under such unphysiological conditions, Dpp internalization through other Dpp binding factors can easily mask the impact of Tkv on Dpp internalization. Consistently, we found that Dpp binds to the cell surface without Tkv and Dally (Fig. 6).

If Tkv is involved in Dpp internalization, extracellular Dpp should increase. Romanova-Michaelides et al. used the observation that extracellular GFP-Dpp did not increase in tkv mutant clones upon GFP-Dpp expression from the anterior stripe of cells (Schwank et al., 2011) to further support Tkv-independent Dpp internalization (Romanova-Michaelides et al., 2015). However, under 400 times higher expression levels of GFP-Dpp than the physiological level, it is not surprising that an increase of extracellular Dpp can be missed. Indeed, we find that endogenous extracellular Dpp is increased upon tkv depletion, indicating that Dpp is internalized through Tkv (Fig. 6). Notably, even when GFP-Dpp was overexpressed, accumulation of GFP-Dpp on the surfaces of cells lacking Tkv have been previously reported (Entchev et al., 2000).

Romanova-Michaelides et al. proposed that, instead of Tkv, Dally is required for internalization of Dpp based on reduced internalization of endogenous GFP-Dpp upon blocking HS chain synthesis or upon cleaving HSPGs using the nanobody internalization assay (Romanova-Michaelides et al., 2015). In contrast, our results suggest that Dally rather antagonizes internalization of Dpp by blocking Tkv-mediated internalization of Dpp (Fig. 6). How can Dpp internalization be reduced in dally mutants if Dally antagonizes internalization of Dpp? We found that extracellular Dpp distribution is severely reduced in dally mutants due to Tkv-mediated internalization of Dpp (Fig. 6), indicating that the amount of extracellular Dpp
available for labeling with nanobody is reduced in *dally* mutants. Thus, we speculate that the reduced internalization of Dpp in *dally* mutants revealed by the nanobody internalization assay reflects the reduced extracellular Dpp rather than reduced ability to internalize Dpp.

Our results do not thus support Dally-mediated internalization and recycling of Dpp as a mechanism of Dpp gradient scaling, (Romanova-Michaelides *et al.*, 2015), although recycling of Dpp through other factors may contribute to Dpp gradient scaling. It would be interesting to test if the Dally-mediated Dpp stabilization on the cell surface is involved in the Dpp gradient scaling. Consistent with this, decrease of Dpp degradation rates has previously been proposed as a mechanism for Dpp gradient scaling (Wartlick *et al.*, 2011). Furthermore, previous pulse-chase experiments showed that, similar to Dally, Pentagone (Pent), a secreted feedback factor required for Dpp gradient scaling, is required for Dpp stability (Vuilleumier *et al.*, 2010). Interestingly, Pent has been shown to interact with HS chains of Dally (Vuilleumier *et al.*, 2010) (Norman *et al.*, 2016), raising a possibility that interaction of HS chains of Dally with Pent is critical for Dpp stability, thereby Dpp gradient scaling.

**Conclusion**

To recognize and activate the specific signal, the affinity of a morphogen to its receptors is high. However, the high affinity of the morphogen-receptor interaction could at the same time compromise the range of morphogens through receptor-mediated removal of morphogens from the extracellular space. Thus, there must exist mechanisms ensuring long-range action of the morphogen despite its high receptor affinity. Our results suggest that Dally counteracts the internalization of Dpp through Tkv to increase the chance that Dpp can reach further despite the high affinity of Dpp to Tkv.
Fig. 1: Distinct models on the roles of HSPGs on Dpp/BMP gradient formation

(A) Schematic view of the wing pouch (future wing tissue) and adult wing tissue. Dpp spreads from the anterior stripe of cells along the A-P compartment boundary to generate pMad gradient and an inverse Brk gradient to specify adult wing veins such as L2 and L5. (B) HSPGs are proposed to transport Dpp via repetitive interaction of HS chains with Dpp (Restricted diffusion model.) (C) HSPGs are proposed to hinder Dpp spreading via reversible interaction of HS chains with Dpp (Hindered diffusion model). (D) HSPGs are proposed to stabilize Dpp on the cell surface via reversible interaction of HS chains with Dpp that antagonizes Tkv-mediated Dpp internalization. (E) HSPGs are proposed to internalize and recycle Dpp.
Fig. 2. Dally, but not Dlp, interacts with Dpp
(A-D) α-HS (3G10) (A, C), extracellular α-Ollas (B, D), and α-pMad (B', D') staining of Ollas-dpp+/+, ap>dally disc (A-B) and Ollas-dpp+/+, ap>dlp disc (C-D). (E-H) Average fluorescence intensity profile of (B, B', D, D') respectively. Data are presented as mean +/- SD. Scale bar: 50µm.
Fig. 3. Dally, but not Dlp, is required for Dpp signaling gradient
(A–D) α-pMad staining of yw (A), daily MH32 (B), daily KO (C), and dlpx KO (D) wing disc. (E–H) Adult wings of yw (E), daily MH32 (F), daily KO (G), and dlpx KO (H). (I) Average fluorescence intensity profile of (A–D). Data are presented as mean +/- SD. (J) Comparison of adult wing size of yw (n=15), daily MH32 (n=20), daily KO (n=13), and dlpx KO (n=13). Data are presented as mean +/- SD. Two-sided unpaired Student’s t test with unequal variance was used for all the comparison. ****p<0.0001. n.s; not significant. (K) Comparison of normalized A-P length against D-V length of adult wings of yw (n=15), daily MH32 (n=20), daily KO (n=13), and dlpx KO (n=13). Data are presented as mean +/- SD. Two-sided unpaired Student’s t test with unequal variance was used for comparison under black lines. Two-sided Mann–Whitney test was used for comparison under red lines. ****p<0.0001, **p<0.01. (L–O) α-pMad staining of yw (L), daily KO (M), YFP-daily (N), and 3xHA-dlp in daily KO (O) wing disc. (P–U) Adult wings of yw (P), daily KO (Q), YFP-daily (R), 3xHA-dlp in daily KO (S), dlpx KO (T), and 3xHA-dlp (U). (V) Average fluorescence intensity profile of (L–O). Data are presented as mean +/- SD. (W) Comparison of adult wing size of yw (n=10), daily KO (n=8), YFP-daily (n=13), 3xHA-dlp in daily KO (n=13), dlpx KO (n=13), and 3xHA-dlp (n=13). Data are presented as mean +/- SD. Two-sided unpaired Student’s t test with unequal variance was used for all the comparison. ****p<0.0001. Scale bar: 50 µm.
Fig. 4. Interaction of core protein of Dally with Dpp

(A-F) α-HS (3G10) (A, D), extracellular α-Ollas (B, E), GFP (B′, E′), α-pMad (C, F) staining of Ollas-dpp/+; ap>dally disc (A-C) and Ollas-dpp/+; ap>dallyΔHS disc (D-F). (G) Average fluorescence intensity profile of (B). (H) Average fluorescence intensity profile of (E). (I) Average fluorescence intensity profile of (B, E) normalized against (B′, E′) respectively. (J) Average fluorescence intensity profile of (C). (K) Average fluorescence intensity profile of (F). (G-K) Data are presented as mean ± SD. (L-N) α-pMad staining of YFP-dally (L), dailyKO (M), and YFP-dailyΔHS (N) wing disc. (O) Average fluorescence intensity profile of (L-N). Data are presented as mean ± SD. (P-R) Extracellular α-HA staining of HA-dpp (P), HA-dpp:dailyKO (Q), and HA-dpp;YFP-dailyΔHS (R) wing disc. (S) Average fluorescence intensity profile of (P-R). Data are presented as mean ± SD. (T-U) Adult wings of YFP-dally (T) and YFP-dailyΔHS (U). (V) Comparison of adult wing size of YFP-dally (n=12) and YFP-dailyΔHS (n=20). Data are presented as mean ± SD. Two-sided unpaired Student’s t test with unequal variance was used for the comparison. ****p < 0.0001. Scale bar: 50 µm.
Fig. 5 HS chains of Dally act largely independent of interaction with Dpp  
(A-B) α-pMad (A, B) and extracellular α-HA (A', B') staining of JAX;HA-dpp disc (A, A') and JAX;HA-dpp ΔHS disc (B, B'). (C) Average fluorescence intensity profile of (A, B). Data are presented as mean +/- SD. (D) Average fluorescence intensity profile of (A', B'). Data are presented as mean +/- SD. (E-F) Adult wings of JAX;HA-dpp (E) and JAX;HA-dpp ΔHS (F). (G) Comparison of adult wing size of JAX;HA-dpp (n=19) and JAX;HA-dpp ΔHS (n=18). Data are presented as mean +/- SD. Two-sided unpaired Student's t test with unequal variance was used for the comparison. ****p < 0.0001. (H-I) extracellular α-Ollas (H, I), YFP (H', I'), and α-pMad (H'', I'') staining of JAX;HA-dpp ΔHS, ap>dally disc (H-H'') and JAX;HA-dpp ΔHS, ap>dallyΔHS disc (I-I''). (J) Average fluorescence intensity profile of (H, I) normalized against (H', I') respectively. (K) Average fluorescence intensity profile of (H''). Data are presented as mean +/- SD. (L) Average fluorescence intensity profile of (I''). Data are presented as mean +/- SD.
Fig. 6 HS chains of Dally stabilize Dpp by antagonizing Tkv-mediated Dpp internalization

(A, B) Extracellular α-Ollas staining of Ollas-dpp/+, ap>tkv RNAi wing disc (A) and dailyKO, Ollas-dpp/+, ap>tkv RNAi wing disc (B). (C, D) Average fluorescence intensity profile of (A, B) respectively. Data are presented as mean +/- SD. (E) Average fluorescence intensity profile of the dorsal compartment of (A, B). Data are presented as mean +/- SD. (F, G) α-pMad staining of Ollas-dpp/+, ap>tkv RNAi wing disc (F), and Ollas-dpp/+, ap>dally RNAi wing disc (G). (H) Extracellular α-Ollas staining of dallyΔHS, Ollas-dpp/+, ap>tkv RNAi wing disc. (I) Average fluorescence intensity profile of (H). Data are presented as mean +/- SD. Scale bar: 50 µm.
Fig. 7 A revised model on the roles of Dally on Dpp/BMP gradient formation and signaling

(A) A previous model. Dally and Dlp compete with Tkv for Dpp binding to antagonize Tkv-mediated Dpp internalization. (B) A revised model. Dally, but not Dlp, reversibly interacts with Dpp partially through its HS chains and mainly through its core protein. The former interaction is not essential and the latter interaction is not sufficient to antagonize Tkv-mediated Dpp internalization under physiological conditions. Upon interaction of Dpp with the core protein of Dally, HS chains of Dally antagonize Tkv-mediated Dpp internalization through Tkv to stabilize Dpp on the cell surface. HS chains of Dally control Dpp signaling also independent of interaction with Dpp likely through Tkv. It remains unknown whether and how Dpp signal activation by core protein bound Dpp and stability by HS chains are coordinated. (C) In dally^−^ mutants, Tkv-mediated Dpp internalization by Tkv is not antagonized. Dpp is irreversibly bound to Tkv and then removed by Tkv from the extracellular space and degraded. (D) In dally^ΔHS^ mutants, the core protein of Dally can interact with Dpp but this interaction is not sufficient to antagonize Tkv-mediated Dpp internalization. Without HS chains, Dpp is irreversibly bound to Tkv and then removed by Tkv from the extracellular space and degraded. (E) In dpp^ΔN^ mutants, Dpp can interact with core protein but not with HS chains of Dally. Without interaction of HS chains with Dpp, HS chains can antagonize Tkv-mediated Dpp internalization through Tkv to stabilize Dpp on the cell surface.
Fig. S1 Comparison of endogenous Dpp gradient and signaling between dorsal and ventral compartment

(A) Extracellular α-Ollas staining of Ollas-dpp/Ollas-dpp wing disc. (B) Average fluorescence intensity profile of (A). (C) α-pMad staining of Ollas-dpp/Ollas-dpp wing disc. (D) Average fluorescence intensity profile of (C).
Fig. S2 Preferential interaction of Dlp with Wg

(A) Extracellular α-Wg staining of Ollas-dpp/+ (A), Ollas-dpp/+, ap>dally (B), and Ollas-dpp/+, ap>dlp (C). (D) Average fluorescence intensity profile of (A-C).
Fig. S3 Generation of dailyKO and dlpKO

(A) Schematic view of manipulation of the glypican genomic loci exemplified with dlp. (B) the exact sequences at the lesion of dailyKO and dlpKO: color code as in (A) (blue: genomic sequences at the inner boundaries of homology arms for HDR, yellow: attP, green: loxP, black: “scars” (remnants of vector sequences including cloning sites).
Fig. S4 Validation of $dlp^{KO}$ allele

(A, B) $\alpha$-Dlp and DAPI staining of $dlp^{KO}/+$ (A) and $dlp^{KO}/dlp^{KO}$ (B).
Fig. S5 JAX does not rescue *dpp* disc allele

(A-D) α-pMad staining (A, C) and α-Brk staining (B, D) of *dpp*<sup>68</sup>/dpp<sup>d12</sup> (A, B) and JAX;*dpp*<sup>68</sup>/dpp<sup>d12</sup> (C, D). (E, F) Adult wings of *dpp*<sup>68</sup>/dpp<sup>d12</sup> (E) and JAX;*dpp*<sup>68</sup>/dpp<sup>d12</sup> (F).
Materials and Methods
Data reporting
No statistical methods were used to predetermine sample size. The experiments were not randomized, and investigators were not blinded to allocation during experiments and outcome assessment.

Fly stocks
Flies for experiments were kept at 25°C in standard fly vials containing polenta and yeast. The following fly lines were used: UAS-tkv-RNAi (Bloomington 40937), UAS-dally-RNAi (VDRC 14136), Ollas-dpp (Bauer Milena, 2022), UAS-GFP-HA-dally (Suzanne Eaton), UAS-dlp (Baeg et al., 2001), HA-dpp (Matsuda et al., 2021), HA-dppΔN (this study), dlp [ko;attB] (this study), dally[ko;attB] (this study), dally[YFP-dally;attP] (this study), dlp[3xHA-dlp;attP] (this study) dally[YFP-dallyΔHS;attP] (this study), dally[3xHA-dlp;attP] (this study), ap-Gal4 (Markus Affolter), JAX (Hoffmann and Goodman, 1987), Ollas-HA-dppΔN (this study), hh-Gal4 (Gift from Dr Manolo Calleja), dallyMH32 (Franch-Marro et al., 2005). yw, dppΔ8 and dppNT2 are described from Flybase.

Genotypes by figures

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Generation of transgenic flies and alleles

HA-dpp\(^{\Delta N}\) plasmid was constructed by GENEWIZ by removing 21bp encoding the 7 basic amino acids (RRPRTRRK) from HA-dpp plasmid (Matsuda et al., 2021).

Ollas-HA-dpp\(^{\Delta N}\) plasmid was constructed by using the HA-dpp\(^{\Delta N}\) plasmid and the Ollas-HA-dpp plasmid as templates. Both plasmids were digested using XhoI and BspEI as restriction enzymes. The small fragment from the HA-dpp\(^{\Delta N}\) plasmid and the large fragment from the Ollas-HA-dpp plasmid were used for ligation. Generation of endogenous dpp alleles using these plasmids were previously described (Matsuda et al., 2021).

Genomic manipulation of glypicans

The daily\(^{KO}\) and dlp\(^{KO}\) alleles were generated by replacing parts of the first exon of the genes with attP cassettes. In both cases, the deleted fragment comprise part of the 5’ UTR, and the complete coding sequence, including the start codon and the signal peptide encoding sequence. For the generation of daily\(^{KO}\) homology arms flanking the target region were amplified by genomic PCR using primer pairs daily\(_5’\)_for (ATAGCTAGGCTCTAAAGACTCAAATTAATTAATTCATTTCAGATTGCGC) and daily\(_5’\)_rev (ATACATATGTTTTTGATGGGTGATTTCATCCAGACACAGTG), or daily\(_3’\)_for (TATACTAGTGAAATGTTTCCATCCATCGTAGAGTTATAATATCG) and daily\(_3’\)_rev (ATAGGGCGCCCTTTTACTCAATTACAGAACAGATATATGGGTACATTCCG) and cloned into the 5’ MCS and 3’ MCS of vector pTV(Cherry) using restriction enzymes NheI and NdeI or SpeI and AscI, respectively (restriction sites underlined). The construct was subjected to P-element transgenesis and was subsequently mobilized and linearized by crossing transgenic flies to flies carrying hsFLP and hsSceI and applying heat-shock as described (Baena-Lopez et al., 2013).

The dlp\(^{KO}\) allele was generated by Cas9/CRISPR mediated homologous repair. Guide RNA sequences flanking the target region were GAAGCATGGCAGAATGCAGAATAAATTCATTTCAGATTGCCG and dlp\(_5’\)_rev (ATACATATGTTTTTGATGGGTGATTTCATCCAGACACAGTG), or dlp\(_3’\)_for (TATACTAGTGAAATGTTTCCATCCATCGTAGAGTTATAATATCG) and dlp\(_3’\)_rev (ATAGGGCGCCCTTTTACTCAATTACAGAACAGATATATGGGTACATTCCG) and cloned into vector pHD-DsRed-attP using restriction enzymes SphI and NotI or SpeI and XhoI, respectively. Homologous directed repair was achieved by co-injecting gRNA plasmids and the repair donor as described (Gratz et al., 2014).
For both dallyKO and dlpKO, successful genomic integrations were identified by the presence of eye markers (mini-white and dsRed, respectively) followed by Cre-mediated excision of marker cassettes. The integrity of the deletion and attP integration was verified by genomic PCR and subsequent sequencing. The exact sequences at the manipulated loci are depicted in Fig. S3. Dally and Dlp genomic constructs were generated in the reintegration vector pRIVwhite and incorporated into the attP site of the dallyKO and dlpKO alleles by phiC31/attB integration (Baena-Lopez et al., 2013). The YFP sequence in Dally constructs was inserted after lysine 90. DallyΔHS contains serine to alanine amino acid substitutions at positions 549, 569, 573 and 597. Three copies of the influenza hemagglutinin derived HA epitope were inserted after glycine 69 of Dlp.

Immunohistochemistry

Total staining
Wing discs from third instar larvae were dissected and stored temporarily in Phosphate Buffered Saline (PBS) (Gibco) on ice until enough samples were collected. The discs were then fixed in 4% Paraformaldehyde (PFA) in PBS for 30min on the shaker at room temperature (25°C). After fixation, the discs were rinsed three times quickly with PBS and three times for 15 min with PBS at 4°C. Wing discs were permeabilized in PBST (0.3% Triton-X in PBS) and then blocked in 5% normal goat serum (NGS) in PBST for at least 30min. Primary antibodies were added in 5% normal goat serum (NGS) in PBST for incubation over night at 4°C. The next day, the primary antibodies were carefully removed and the samples were rinsed three times quickly in PBST and three times 15 min at room temperature in PBST. Discs were incubated in secondary antibody for 2h at room temperature. Afterwards the samples were washed again three times quickly and three times 15 min in PBST at room temperature. After the final washing the PBST was rinsed with PBS, then the PBS was removed completely and the samples were mounted in VECTORSHIELD on glass slides. For HS stainings, wing discs were dissected as described above. After fixation, discs were washed in PBS and blocked in 5% NGS in PBST, then treated with Heparinase III (Sigma-Aldrich) for 1.5h at 37°C. Afterwards the discs were blocked again in 5% NGS in PBST and stained as described above.

Extracellular staining
Wing discs from third instar larvae were dissected and stored temporarily in Schneider's Drosophila medium (S2) on ice until enough samples were collected. The discs were then blocked in cold 5% NGS in S2 medium on ice for 10min. The blocking solution was removed carefully and the primary antibody was added for 1h on ice. During the 1h incubation period, the tubes were tapped carefully every 15min, to make sure the antibody is distributed evenly. After 1h incubation on ice, the antibody was removed and the samples were washed at least 6 times with cold S2 medium and another two times with cold PBS to remove excess primary antibody. Wing discs were then fixed with 4% PFA in PBS for 30min on the shaker at room temperature (25°C). After fixation the protocol continued as described in total stainings.

Antibodies
Primary antibodies: rabbit-anti-phospho-Smad1/5 (41D10, Cell Signaling, #9516; 1:200), mouse-anti-Wg (4D4, DSHB, University of Iowa; total staining: 1:120, extracellular staining: 1:120), mouse-anti-Ptc (DSHB, University of Iowa; total staining: 1:40), rat-anti-HA (3F10, Roche, 11867423001; total staining: 1:300, extracellular staining: 1:20), rat-anti-Ollas (L2, Novus Biologicals, NB1-06713; total staining: 1:300, extracellular staining: 1:20), mouse-anti-HS (F69-3G10, amsbio; 1:100)

The following secondary antibodies were used at 1:500 dilutions in this study. Goat anti-rabbit IgG (H+L) Alexa Fluor™ 488 (A11008 Thermo Fischer), goat anti-rabbit IgG (H+L) Alexa Fluor™ 568 (A11011 Thermo Fischer), goat anti-rabbit IgG (H+L) Alexa Fluor™ 680 (A21109 Thermo Fischer), goat anti-rat IgG (H+L) Alexa Fluor™ 568 (A11077 Thermo Fischer), goat anti-mouse IgG (H+L) Alexa Fluor™ 568 (A11004 Thermo Fischer), goat anti-rat IgG Fc 488 (ab97089 abcam), goat anti-mouse IgG Fc Alexa Fluor™ 680 (115625071 Jackson Immuno Research).
Imaging
Samples were imaged using a Leica SP5-II-MATRIX confocal microscope and Leica LAS AF and Images were analyzed using ImageJ. Figures were obtained using Omero and Adobe Illustrator.

Quantification of pMad and extracellular staining
From each z-stack image, signal intensity profile along A/P axis was extracted from average projection of three sequential images using ImageJ (v.2.0.0-rc69/1.52p). Each signal intensity profile collected in Excel (Ver. 16.51) was aligned along A/P compartment boundary (based on anti-Ptc staining or pMad staining) and average signal intensity profile from different samples was generated and plotted by the script (wing_disc-alignment.py). The average intensity profile from control and experimental samples was then compared by the script (wingdisc_comparison.py). Both scripts can be obtained from https://etiennees.github.io/Wing_disc-alignment/. The resulting signal intensity profiles (mean with SD) were generated by Prism (v.8.4.3(471)). Figures were prepared using Omero (ver5.9.1) and Illustrator (24.1.3).

Quantification of adult wing size
The size of each compartment was measured using ImageJ (v.2.0.0-rc69/1.52p) and collected in Excel (Ver. 16.51). Scatter dot plots (mean with SD) were generated by Prism (v.8.4.3(471)). Figures were prepared using Omero (ver5.9.1) and Illustrator (24.1.3).

Statistics
All images were obtained from multiple animals (n > 3). The experiments were repeated at least two times independently with similar results. Statistical significance was assessed by Prism (v.8.4.3(471)) based on the normality tests. The observed phenotypes were highly reproducible as indicated by the significance of p values obtained by statistical tests.

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References


