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1 Chondroitin sulfate proteoglycan Windpipe modulates Hedgehog signaling in *Drosophila* 2

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12 Abstract

- 13
- 14 Proteoglycans, a class of carbohydrate-modified proteins, often modulate growth factor signaling on the
- 15 cell surface. However, the molecular mechanism by which proteoglycans regulate signal transduction is
- 16 largely unknown. In this study, using a recently-developed glycoproteomic method, we found that
- 17 Windpipe (Wdp) is a novel chondroitin sulfate proteoglycan (CSPG) in *Drosophila*. Wdp is a single-pass
- 18 transmembrane protein with leucin-rich repeat (LRR) motifs and bears three CS sugar chain attachment
- 19 sites in the extracellular domain. Here we show that Wdp modulates the Hedgehog (Hh) pathway.
- 20 Overexpression of *wdp* inhibits Hh signaling in the wing disc, which is dependent on its CS chains and
- 21 the LRR motifs. Conversely, loss of *wdp* leads to the upregulation of Hh signaling. Furthermore,
- 22 knockdown of *wdp* increase the cell surface accumulation of Smoothened (Smo), suggesting that Wdp
- 23 inhibits Hh signaling by regulating Smo stability. Our study demonstrates a novel role of CSPG in
- 24 regulating Hh signaling.
- 25

26 Introduction

27

28 Spatial and temporal regulation of growth factor signaling pathways is essential to proper development

- and disease prevention. Cell surface signaling events, such as ligand-receptor interactions, are often
- 30 modulated by proteoglycans (D. Xu & Esko, 2014). Proteoglycans are carbohydrate-modified proteins
- that are found on the cell surface and in the extracellular matrix. They are composed of a core protein and one or more glycosaminoglycans (GAGs) covalently attached to specific serine residues on the core
- 32 one of more grycosammogrycans (GAOs) covarently attached to specific serine residues on the core 33 protein. GAGs are long, unbranched, and highly sulfated polysaccharide chains consisting of a repeating
- disaccharide unit. Based on the composition of the disaccharide units, proteoglycans are classified into
- several types, including heparan sulfate proteoglycans (HSPGs) and chondroitin sulfate proteoglycans
- 36 (CSPGs).
- 37
- 38 HSPGs function as co-receptors by interacting with a wide variety of ligands and modulate signaling
- activities (Holt & Dickson, 2005; J.-S. Lee & Chien, 2004; Lindahl & Li, 2009; Poulain & Yost, 2015; D.
- 40 Xu & Esko, 2014). *Drosophila* offers a powerful model system to study the functions of HSPGs *in vivo*
- 41 because of its sophisticated molecular genetic tools and minimal genetic redundancy in genes encoding
- 42 core proteins and HS synthesizing/modifying enzymes (Lander & Selleck, 2000; Nakato & Li, 2016;
- 43 Perrimon & Bernfield, 2000; Takemura & Nakato, 2015). *In vivo* studies using the *Drosophila* model
- 44 have shown that HSPGs orchestrate information from multiple ligands in a complex extracellular milieu
- and sculpt the signal response landscape in a tissue {Nakato & Li, 2016}. However, the molecular
- 46 mechanisms of co-receptor activities of HSPGs still remain a fundamental question. Our previous studies
- 47 predict that there are unidentified molecules involved in the molecular recognition events on the cell
- 48 surface (Akiyama et al., 2008).
- 49

50 In addition to HS, *Drosophila* produces CS, another type of GAG (Toyoda, Kinoshita-Toyoda, & Selleck,

- 51 2000). CSPGs are well known as major structural components of the extracellular matrix. CSPGs have
- 52 also been shown to modulate signaling pathways, including Hedgehog (Hh), Wnt, and fibroblast growth
- factor signaling (Cortes, Baria, & Schwartz, 2009; Townley & Bülow, 2018). Given the structural
- 54 similarities between CS and HS, CSPGs may have modulatory, supportive and/or complementary
- 55 functions to HSPGs. However, the mechanisms by which CSPGs function as a co-receptor are unknown.
- 56 In contrast to a large number of studies on HSPGs, very few CSPGs have been identified and analyzed in
- 57 Drosophila (Momota, Naito, Ninomiya, & Ohtsuka, 2011). Unlike HSPGs, CSPG core proteins are not
- 58 well conserved between species (Olson, Bishop, Yates, Oegema, & Esko, 2006). Therefore, the
- 59 identification of CSPGs cannot rely on the sequence homology to mammalian counterparts.
- 60

61 Recently, we have developed a glycoproteomic method to identify novel proteoglycans (Noborn et al.,

- 62 2016; 2018; 2015). Briefly, this method includes trypsinization of protein samples, followed by
- 63 enrichment of glycopeptides using strong anion exchange (SAX) chromatography. After enzymatic
- 64 digestion of HS/CS chains, the glycopeptides bearing a linkage glycan structure common to HS and CS
- 65 chains are identified using nano-liquid chromatography-tandem mass spectrometry (nLC-MS/MS). This
- 66 method has successfully identified novel CSPGs in humans (Noborn et al., 2015) and *Caenorhabditis*
- 67 *elegans* (Noborn et al., 2018).
- 68
- To study the function of CSPGs in signaling, we applied the glycoproteomic method to identify
- 70 previously unrecognized CSPGs in *Drosophila*. We found that Windpipe (Wdp) is a novel CSPG and
- 71 affects Hh signaling. Overexpression of *wdp* inhibits Hh signaling in the wing disc. This inhibitory effect
- of Wdp on Hh signaling is dependent on its CS chains and LRR motifs. Consistent with the
- 73 overexpression analysis, loss of *wdp* increases Hh signaling. Loss of *wdp* also increases cell surface
- accumulation of Smoothened (Smo), the Hh signaling transducer. Therefore, we propose that Wdp
- 75 downregulates Hh signaling by disrupting cell surface accumulation of Smo.
- 76

77 Results

78

79 A glycoproteomic approach identified Wdp as a novel *Drosophila* CSPG

80

81 We investigated the potential presence of CSPGs in *Drosophila* using our recently-developed 82 glycoproteomic approach that identifies core proteins and its CS attachment sites. A general workflow for 83 the sample preparation, CS-glycopeptide enrichment, LC-MS/MS analysis and the subsequent data 84 analysis is shown in Fig. 1A. Brifely, Drosophila third-instar larvae were collected from two different genotypes (wild type [Oregon-R] and a loss-of-function mutant for tout-velu [ttv⁵²⁴]) and the material was 85 86 homogenized in ice-cold acetone. tty encodes a Drosophila HS polymerase, and tty mutants lack HS 87 chains (Toyoda et al., 2000). The samples were incubated with trypsin and then passed over an anion 88 exchange column equilibrated with a low-salt buffer. This procedure enriches for CS-attached 89 glycopeptides as the matrix retains anionic polysaccharides and their attached peptides, whereas neutral or 90 positively charged peptides flow through the column. The bound structures were eluted stepwise with 91 three buffers of increasing sodium chloride concentrations. The resulting fractions were treated with 92 chondroitinase ABC. This procedure reduces the lengths of the CS chains and generates a residual 93 hexasaccharide structure still attached to the core protein. The chondroitinase-treated fractions were 94 analyzed with positive mode nLC-MS/MS and an automated search strategy was used to identify CS 95 modified peptides in the generated data set (Noborn et al., 2015). 96

97 The analysis revealed the Windpipe (Wdp) protein as a novel CSPG, which was modified with three CS-

98 polysaccharides on two unique peptides (Fig. 1B and 1C). We detected Wdp glycopeptides from both 99 wild-type and *tty* mutant samples, further supporting that Wdp bears CS chains, not HS. One of the

99 wild-type and *ttv* mutant samples, further supporting that Wdp bears CS chains, not HS. One of the 100 identified precursor ions (m/z 983.38; 3+) equated to the mass of a peptide with a

101 SDOVEGSGDLSETNMELK sequence, derived from the middle part of the protein (amino acids 276–

102 293) (Fig. 1B). The peptide was modified with one hexasaccharide structure and one methionine

103 oxidation. The measured mass (2947.1186 Da) deviated - 3.27 ppm from the theoretical value. The other

104 identified precursor ion $(m/z \ 1276.76; 4+)$ equated to the mass of a peptide with a

105 EEHIVKDEDEDDEGSGSGGGLLIIPDPSK sequence, located in proximity to the previous peptide

106 (amino acids 320–348) (Fig. 1C). The peptide was found to be modified with two hexasaccharide

107 structures and where one of the hexasaccharides were modified with one phosphate modification. The

108 measured mass (5102.9389 Da) deviated +3.05 ppm from the theoretical value. Detailed inspection of the

109 spectra revealed several b- and y-ions as well as the prominent diagnostic oxonium ion at m/z 362.1,

110 corresponding to the disaccharide $[GlcAGalNAc-H_2O+H]^+$ (Fig. 1B and 1C). Furthermore, one of the

- 111 glycans in Fig 1C was found modified with one phosphate group at a xylose residue (peptide + xylose + phosphate, m/z 1625.70; 2+).
- 112

Wdp is a single-pass transmembrane protein containing four leucine rich repeat (LRR) motifs in the
extracellular domain (Huff, Kingsley, Miller, & Hoshizaki, 2002). The three CS attachment sites (S282,
S334, and S336) revealed by our glycoproteomic analysis are located in the extracellular domain (Fig.

3354, and 3556) revealed by our grycoproteonine analysis are located in the extracentular domain (Fig. 3A). Interestingly, a recent study reported that Wdp negatively regulates JAK–STAT signaling by

promoting internalization and lysosomal degradation of the Domeless (Dome) receptor (W. Ren et al.,

119 2015). We further investigated the role of Wdp, a novel CSPG, in signal transduction.

120

121

122 **Overexpression of** *wdp* **inhibits Hh signaling**

123

124 The growth and patterning of the *Drosophila* wing are controlled by multiple signaling pathways,

125 including Decapentaplegic (Dpp; the Drosophila BMP), Wingless (Wg; the Drosophila Wnt), and

- 126 Hedgehog (Hh) signaling (Baena-Lopez, Nojima, & Vincent, 2012; Tabata & Takei, 2004). To determine
- 127 the role of *wdp* in these developmental signaling pathways, we first asked whether overexpression of *wdp*

128 affects adult wing morphology. When *wdp* was overexpressed in the wing pouch using Bx^{MS1096} -GAL4

129 (Capdevila & Guerrero, 1994) ($Bx^{MS1096} > wdp$), the wing size was reduced compared to that of control

flies $(Bx^{MS1096}>)$ (Fig. 3C; compared to Fig. 3B). In addition, the distance between longitudinal wing veins

- 3 and 4 (L3 and L4) was aberrantly narrower. This decreased distance between L3 and L4 is indicative of
 reduced Hh signaling during wing development (Mullor, Calleja, Capdevila, & Guerrero, 1997; Strigini &
- 133 Cohen, 1997).
- 134

135 Hh is produced in the posterior compartment of the wing disc and spreads towards the anterior

- compartment where Hh signaling induces target genes expression in a concentration-dependent manner
 (Briscoe & Thérond, 2013; Gradilla & Guerrero, 2013; Hartl & Scott, 2014). Expression of high-
- 138 threshold target genes, such as Patched (Ptc; the Hh receptor) (Capdevila, Pariente, Sampedro, Alonso, &
- 139 Guerrero, 1994) and Engrailed (En) (Patel et al., 1989) are induced in anterior cells near the
- 140 anteroposterior compartment boundary by high levels of Hh signaling (Jia, Tong, Wang, Luo, & Jiang,
- 141 2004) (Fig. 2A and 2E). Low levels of Hh signaling induce the expression of *dpp* and the accumulation of
- 142 full-length Cubitus interruptus (Ci; the transcriptional factor of Hh signaling) in a broader region (more
- 143 distant away from the anteroposterior boundary) (Fig. 2A and 2C). To determine if Hh signaling is indeed
- 144 affected by wdp, we overexpressed wdp in the dorsal compartment of the wing disc using ap-GAL4
- 145 (Calleja, Moreno, Pelaz, & Morata, 1996; O'Keefe, Thor, & Thomas, 1998). We found that *wdp*
- 146 overexpression in the dorsal compartment reduced the expression domains of both "high-threshold" 147 targets (Ptc and En) and "low-threshold" targets ($dpp-lacZ^{10638}$, a reporter for dpp expression, and full-
- 147 targets (Ptc and En) and Tow-threshold targets (*app-tacz*), a reporter for *app* expression, and full-148 length Ci) compared to those in the ventral compartment (Fig. 2B, 2D, and 2F). Notably, overexpression
- 148 of *wdp* did not affect the pattern of a *hh* transcriptional reporter *hh*-lac Z^{P30} (J. J. Lee, Kessler, Parks, &
- Beachy, 1992) (Fig. 2F). Together, *wdp* acts as a negative regulator of Hh signaling without affecting *hh* transcription
- 151

153 On the other hand, Wdp does not appear to affect Dpp and Wg pathways. When *wdp* is overexpressed

- 154 using *ap-GAL4* or *hh-GAL4* (a posterior compartment-specific GAL4 driver) (Tanimoto, Itoh, Dijke, &
- 155 Tabata, 2000), we did not observe apparent defects in Dpp signaling activity, which was monitored by the
- 156 expression of phosphorylated Mad (pMad) and Spalt major (Salm) (readouts of Dpp signaling). Similarly,
- no changes in expression of Senseless (Sens) and Distal-less (Dll) (readouts of Wg signaling) were
- detected (Fig. S1). These results are consistent with a previous report (W. Ren et al., 2015).
- 159

160 We also found that overexpression of *wdp* induces massive apoptosis, as detected with anti-cleaved 161 Caspase-3 antibody (Fig. S2B). This likely contributed to the smaller adult wing phenotype observed in 162 Bx^{MS1096} >wdp flies. It was recently reported that Hh signaling is required for cell survival in wing disc

- 162 by wap mes. It was recently reported that fin signaling is required for een survival in wing disc 163 cells (Lu, Wang, & Shen, 2017). To determine whether reduced Hh signaling is responsible for the
- 164 observed apoptosis, we first asked if reduced Hh signaling results in apoptosis. We inhibited Hh signaling
- 165 either by expressing an RNAi construct targeting *smo* (TRiP.HMC03577) (Fig. S2E), or by
- 166 overexpressing *ptc* in the dorsal compartment using *ap-GAL4*. We found that neither treatment caused
- 167 massive apoptosis (Fig. S2F and S2G), indicating that reduced Hh signaling is not sufficient to induce
- 168 massive apoptosis in the wing disc. Furthermore, coexpression of a constitutively active form of Smo
- 169 with Wdp did not suppress apoptosis in the wing disc (Fig. S2H). Thus, these results suggest that
- 170 overexpression of *wdp* induces apoptosis, independent of reduced Hh signaling.
- 171
- 172

173 CS and LRR motifs are necessary for Wdp to inhibit Hh signaling

174

175 Next, we asked whether the CS chains of Wdp are required for its function. In a CSPG core-protein, CS is

- attached to specific serine residues in the consensus serine-glycine dipeptide surrounded by acidic amino
- acids (Esko & Zhang, 1996). We generated a UAS-wdp^{AGAG} construct in which all three serine residues
- 178 (S282, S334, and S336) are substituted with alanine residues so that CS cannot be attached to the core

- 179 protein (Fig. 3A). The UAS-wdp^{ΔGAG} construct was inserted in the same genomic location (ZH-86Fb;
- 180 (Bischof, Maeda, Hediger, Karch, & Basler, 2007)) as UAS-wdp using the phiC31 site-specific integration
- system (Groth, Fish, Nusse, & Calos, 2004) in order to ensure the same expression level of the UAS
 transgenes.
- 183
- We found that $Bx^{MS1096} > wdp^{AGAG}$ adult wings did not display the reduction in the distance between L3 and
- 185 L4 (Fig. 3B). Consistent with this, the expression of Ptc, En, Ci, and *dpp-lacZ* in the wing disc were not
- affected by $wdp^{\Delta GAG}$ overexpression in the dorsal compartment of the wing disc (Fig. 3E–G). These
- results indicate that CS chains are required for Wdp's activity to downregulate Hh signaling.
- 188

189 To determine whether the LRR motifs and/or the intracellular domain of Wdp are necessary for inhibiting 190 Hh signaling, we generated several Myc-tagged mutant constructs (Fig. S3) and examined their activities.

- 191 Consistent with the earlier result (Fig. 2B), expression of a Myc-tagged Wdp (Myc:Wdp) led to the
- 192 narrower Ptc expression domain (Fig. 3G). A mutant *wdp* construct lacking LRR motifs (Myc:Wdp^{Δ LRRs})
- failed to inhibit Hh signaling (Fig. 3H). On the other hand, a truncated construct lacking the intracellular
- domain (Myc:Wdp $^{\Delta ICD}$) retained the ability to inhibit Hh signaling (Fig. 3I). Thus, in addition to CS
- 195 chains, the LRR motifs of Wdp are required for inhibiting Hh signaling.
- 196 197

198 Wdp expression in the wing disc

199

To monitor Wdp expression, we generated transgenic flies (wdp^{KLHA} and $wdp^{KLOLLAS}$) expressing epitopetagged Wdp protein from its endogenous locus. We inserted a spaghetti monster GFP with 10 copies of HA or OLLAS tags (Nern, Pfeiffer, & Rubin, 2015; Viswanathan et al., 2015) near the C-terminus of Wdp (after Q652; Fig. 4A) using CRISPR–Cas9-mediated homology-directed repair (Gratz et al., 2014; X. Ren et al., 2014). The Wdp:HA expression was detected in the eye disc, adult midgut, and tracheal

- system (Fig. S3), consistent with previous reports (Huff et al., 2002; W. Ren et al., 2015).
- 206

In the wing disc, Wdp:HA is expressed in most of the wing disc cells with enrichment in the basal side, as
detected by anti-HA antibody (Fig. 4B and 4C). This result was confirmed by anti-OLLAS antibody
staining of the *wdp^{KLOLLAS}* wing discs (Fig. S3A and S3B). In the wing disc epithelium, mitotic nuclei
apically translocate, but the cells maintain contact with the basal lamina via actin-rich basal projection
(Ragkousi & Gibson, 2014). Interestingly, Wdp:HA is strongly enriched in such basal projections (Fig.
4A and 4D). However, physiological significance of this localization of Wdp in the basal projections of
mitotic cells is unknown.

- 214
- 215

216 Loss of *wdp* leads to higher levels of Hh signaling

217

218 To determine whether loss of wdp affects Hh signaling activity, we examined the effect of wdp RNAi knockdown in the wing disc. Expression of a wdp^{RNAi} construct (TRiP.HMC06302) using *ap-GAL4* in 219 220 wdp^{KLHA/+} flies led to the loss of Wdp:HA staining specifically in the dorsal compartment (Fig. 4E), 221 validating the efficacy of RNAi-mediated knockdown of wdp. We then examined the effect of wdp 222 knockdown on Hh signaling using the Ptc expression level as a readout of the Hh signaling activity. In 223 control wing discs (ap>FLP), the signal intensity of Ptc staining in the dorsal compartment is comparable to that in the ventral compartment (Fig. 5B). On the other hand, wdp^{RNAi} expression using ap-GAL4 224 225 increased the signal intensity of Ptc staining only in the dorsal compartment (Fig. 5A and 5C). In 226 addition, we observed that the *dpp-lacZ* expression domain was expanded anteriorly by *wdp* knockdown 227 (Fig. 5D). In the adult wing, knockdown of *wdp* slightly expanded the distance between wing vein L3 and 228 L4 near the distal tip (Fig. 5J, compared to Fig. 5I). Thus, wdp RNAi knockdown results in a moderate

229 increase in Hh signaling.

230

- 231 To confirm the *wdp* knockdown phenotypes, we generated a loss-of-function allele of *wdp* ($wdp^{KO.ACDS}$),
- in which most of the *wdp* coding sequence was removed using CRISPR–Cas9-mediated defined deletion
- 233 (Gratz et al., 2013) (Fig. 5E–G). $wdp^{KO, ACDS}$ homozygous mutant clones were induced in the wing pouch
- using the FLP–FRT system with *nubbin (nub)-GAL4 UAS-FLP* and their effect on Hh signaling was
- examined using anti-Ptc antibody. Consistent with the RNAi knockdown results, we observed a modest
- 236 increase of Ptc expression in cells mutant for *wdp* (Fig. 5H). Taken together, we conclude that *wdp*
- 237 negatively regulates Hh signaling in the *Drosophila* wing.
- 238
- 239

Wdp inhibits Smo cell surface accumulation

- 242 The seven-pass transmembrane protein Smo is a key transducer of Hh signaling. In the absence of Hh, Ptc
- 243 inhibits the phosphorylation of Smo, which is internalized and degraded (Zhu, Zheng, Suyama, & Scott,
- 244 2003). In the presence of Hh, restriction of Ptc on Smo is relieved, allowing Smo to accumulate on the
- cell surface and activate Hh signaling. Although *smo* transcription is ubiquitous, Smo protein expression
- 246 levels are high in the posterior compartment of the wing disc where Ptc is not expressed (Fig. 6A) (Denef,
- 247 Neubüser, Perez, & Cohen, 2000). We found that knockdown of *wdp* increases the cell surface
- 248 accumulation of Smo (Fig. 6B). This result suggests that Wdp downregulates Hh signaling either by
- 249 disrupting Smo translocation to the cell membrane or the stability of Smo on the cell surface.
- 250
- 251

252 Discussion

253

The molecular mechanism by which HSPG co-receptors regulate growth factor signaling remains a central question in cell biology. Dally, a *Drosophila* HSPG of the glypican type, potentiates Dpp signaling by stabilizing the ligand-receptor complex on the cell surface (Akiyama et al., 2008), suggesting that controlling the rate of receptor-mediated internalization of the signaling complex is the basis for co-

258 receptor activity. However, it is still unknown how HSPGs affect endocytosis and internalization. Since

259 glypicans do not have an intracellular domain, it is likely that these molecules cooperate with other

factors (e.g. membrane proteins) to exert co-receptor activity. Thus, it is clear that there are many more

unknown factors involved in molecular recognition events on the cell surface. To understand the

- 262 molecular basis for cell communications, it is critical to identify novel cell surface players.
- 263

264 We found that in the wing disc, Wdp negatively regulates Hh signaling in a CS- and LRR motif-

265 dependent manner. It has also been reported that Wdp negatively regulates JAK-STAT signaling and

controls adult midgut homeostasis and regeneration (W. Ren et al., 2015). The authors showed that Wdp

267 interacts with the Dome receptor and promotes its endocytosis and lysosomal degradation. Thus, it is

268 interesting to test if Wdp interacts with Dome via CS chains to modulate JAK–STAT signaling. We

269 observed that Wdp affects cell surface accumulation of Smo, suggesting its role in regulating the stability

270 of Smo protein. Thus, it is possible that Wdp modulates these pathways via a similar mechanism:

271 controlling the internalization of Dome and Smo on the cell membrane.

272

273 It is worth noting that both JAK–STAT and Hh signaling, the two pathways negatively controlled by

274 Wdp, are also regulated by HSPGs. Dally-like, a glypican family of HSPGs, positively regulates Hh

275 signaling by interacting with Hh and Ptc (Desbordes & Sanson, 2003; M.-S. Kim, Saunders, Hamaoka,

276 Beachy, & Leahy, 2011; Lum, Yao, et al., 2003a; Williams et al., 2010; Yan et al., 2010). In the

developing ovary, Dally upregulates the JAK–STAT pathway (Y. Hayashi et al., 2012). Given the

importance of precise dosage control of oncogenic pathways, such as JAK–STAT and Hh signaling, this

dual proteoglycan system could play an important role in fine-tuning of the signaling output in order to

280 prevent cancer formation. In vertebrates, HSPGs and CSPGs show opposing effects in neural systems.

For example, axon growth is typically promoted by HSPGs but inhibited by CSPGs (Bandtlow & Zimmermenn, 2000; Coles et al., 2011; Kantor et al., 2004; Matsurgets, Leis, Letters, Tassian Leis, Leis, Letters, Tassian Leis, Letters, Lette

Zimmermann, 2000; Coles et al., 2011; Kantor et al., 2004; Matsumoto, Irie, Inatani, Tessier-Lavigne, &
Yamaguchi, 2007; Silver & Miller, 2004; Van Vactor, Wall, & Johnson, 2006). Our findings suggest that
such competing effects of HSPGs and CSPGs may be a general mechanism for the precise control of
signaling cascades and pattern formation.

285

In addition to the functions in signaling, Wdp may play other roles. We found that overexpression of *wdp* results in massive apoptosis in the wing disc, independent of Hh signaling inhibition (Fig. S2). Since

results in massive apoptosis in the wing disc, independent of Hh signaling inhibition (Fig. S2). Since

289 CSPGs are well known for structural functions, an excess amount of Wdp may affect the epithelial

integrity of the wing disc, leading to subsequent apoptosis. Our observation that Wdp is enriched on the

basal side of the wing disc and adult midgut cells (Fig. 4B and S3F) suggests that Wdp may interact with

components of the basement membrane, which surrounds these organs.

293

In mice, sulfated CS is necessary for Indian hedgehog (Ihh) signaling in the developing growth plate
(Cortes et al., 2009). Ihh and Sonic hedgehog (Shh) bind to CS (Cortes et al., 2009; Whalen, Malinauskas,
Gilbert, & Siebold, 2013; F. Zhang, McLellan, Ayala, Leahy, & Linhardt, 2007). Thus, it will be

297 interesting to check if Wdp interacts with Hh via its CS chains.

298

299 Previous studies also reported that *wdp* is associated with aggressive behaviors in *Drosophila* species.

300 wdp is upregulated in the head of socially isolated male flies, which exhibit more aggressive behaviors

than males raised in groups (L. Wang, Dankert, Perona, & Anderson, 2008). Also, *wdp* expression is

302 slightly higher in the brain of *Drosophila prolongata*, which is more aggressive compared to its closely-

- 303 related species (Kudo et al., 2017). Since CSPGs are important in neuronal patterning (Saied-Santiago &
- Bülow, 2018), it is interesting to study the molecular mechanisms behind Wdp's effect on *Drosophila* behavior.
- 305 be 306
- 307 In mammals, there are a class of CSPG molecules with LRR motifs (small leucine-rich proteoglycans, or
- 308 SLRPs). A number of SLRP members are known as causative genes of human genetic disorders (Bech-
- 309 Hansen et al., 2000; Pusch et al., 2000; Schaefer & Iozzo, 2008). Although Wdp does not have cysteine-
- rich regions that are commonly found in mammalian SLRPs, MARRVEL (ver 1.1) (J. Wang et al., 2017)
- reports that *wdp* is a potential *Drosophila* ortholog of the human *NYX* gene (nyctalopin), a member of
- 312 SLRPs (DIPOT score 1 (Hu et al., 2011)). Mutations in *NYX* cause X-linked congenital stationary night
- 313 blindness (Bech-Hansen et al., 2000; Pusch et al., 2000). Further studies on Wdp will provide a novel
- 314 insight into the function of these disease-related human counterparts.
- 315 316

317 **Materials and Methods**

318

319 Preparation of glycosaminoglycan-glycopeptides and LC-MS/MS analysis

320

321 Glycosaminoglycan-glycopeptide samples were prepared from wild-type (Oregon-R) and ttv mutant

322 (ttv^{524}) third-instar larvae as previously described (Noborn et al., 2015; 2018). Briefly, 200-400 third 323 instar larvae (wet weight; 200-400 mg) were lyophilized and homogenized using a motor pestle in 1 ml of

324 ice-cold acetone. After extensive washes with acetone, the insoluble fraction was recovered by

325 centrifugation. After overnight desiccation, the pellet was dissolved in 1.5 ml 1% CHAPS lysis buffer and

boiled for 10 min at 96°C. The sample was adjusted to 2 mM MgCl₂ and incubated with 3 µl Benzonase 326

327 (MilliporeSigma, Burlington, MA) at 37°C for three hours. After heat-inactivation of Benzonase, the

328 sample was centrifuged and the supernatant was collected in a new tube.

329

An aliquot of the preparation (1 mg of protein) was further used. The sample was reduced and alkylated 330

in 1 ml 50 mM NH₄HCO₃, and trypsinized at 37°C overnight with 20 µg trypsin (Promega, Madison, WI). 331

332 The digested samples were applied onto DEAE (GE Healthcare, Chicago, IL) columns (600 µl) at 4°C.

333 The columns were washed with three different low-salt washing solutions at 4°C: 50 mM Tris-HCl, 100

334 mM NaCl, pH 8.0; 50 mM NaAc, 100 mM NaCl, pH 4.0; and 100 mM NaCl. The glycopeptides that

335 were bound to DEAE were eluted stepwise with three buffers with increasing sodium chloride

336 concentrations at 4°C: 4 ml 250 mM NaCl, 400 mM NaCl, 800 mM NaCl, and 3 ml 1500 mM NaCl.

337 Each fraction was desalted using PD10-columns (GE Healthcare).

338

339 All fractions were lyophilized and the salt-free samples were then individually treated with 1 mU of 340 chondroitinase ABC (Sigma-Aldrich, St. Louis, MO) for 3 h at 37°C. Prior to MS-analysis, the samples 341 were desalted using a C18 spin column (8 mg resin) according to the manufacturer's protocol (Thermo 342 Fisher Scientific, Waltham, MA). LC-MS/MS analysis was performed as previously described (Noborn et 343 al., 2015; 2018). In brief, the samples were analyzed on a Q Exactive mass spectrometer coupled to an 344 Easy-nLC 1000 system (Thermo Fisher Scientific). Briefly, glycopeptides (2-µl injection volume) were separated using an analytical column with Reprosil-Pur C18-AQ particles (Dr. Maisch GmbH, 345

346 Ammerbuch, Germany). The following gradient was run at 300 nl/min; from 7-35 % B-solvent

347 (acetonitrile in 0.2% formic acid) over 75 min, to 100 % B-solvent over 5 min, with a final hold at 100%

348 B-solvent for 10 min. The A-solvent was 0.2% formic acid. Spectra were recorded in positive ion mode

- 349 and MS scans were performed at 70,000 resolution with a mass range of m/z 600–1800. The MS/MS 350 analysis was performed in a data-dependent mode, with the top ten most abundant charged precursor ions
- in each MS scan selected for fragmentation (MS2) by higher energy collision dissociation with 351

352 normalized collision energy values of 30. The MS2 scans were performed at a resolution of 35,000 (at m/z

353 200). The data analyses were performed as previously described (Noborn et al., 2015) with some small

354 adjustments. In brief, the HCD raw spectra were converted to Mascot .mgf format using Mascot distiller

355 (version 2.3.2.0, Matrix Science, London, UK). The ions were presented as singly protonated in the

output Mascot file. Searches were performed using an in-house Mascot server (version 2.3.02) with the 356

357 enzyme specificity set to Trypsin, and then to Semitrypsin, allowing for one or two missed cleavages, in

358 subsequent searches on Drosophila sequences of the UniprotKB (42, 507, sequences, 2018-06-18). The

359 peptide tolerance was set to 10 parts per million (ppm) and fragment tolerance was set to 0.01 Da. The

360 searches were allowed to include variable modifications at serine residues of the residual hexasaccharide structure [GlcA(-H₂O)GalNAcGlcAGalGalXyI-O-] with 0 (C₃₇H₅₅NO₃₀, 993.2809 Da), 1 (C₃₇H₅₅NO₃₃S, 361

362 1073.2377 Da), or 2 (C₃₇H₅₅NO₃₆S₂, 1153.1945 Da) sulfate groups attached.

363 364

365 Fly husbandry and fly strains, and transgenic flies

366

The following fly strains were used in this study: 367

368

Oregon-R, w¹¹¹⁸ (Bloomington Drosophila Stock Center [BDSC] #5905), ttv⁵²⁴ (Takei, 2004), ap-GAL4 369 (O'Keefe et al., 1998), *hh-GAL4* (Tanimoto et al., 2000), *Bx*^{MS1096}-GAL4 (BDSC #8860) (Capdevila & 370 Guerrero, 1994), AB1-GAL4 (BDSC #1824) (Tavsanli et al., 2004), elav^{C155}>mCD8:GFP (BDSC #5146) 371 (Lin & Goodman, 1994), UAS-GFP (BDSC #1521), UAS-tdTomato (BDSC #36327 and #36328), UAS-372 373 FLP (BDSC #4539 and #4540), UAS-ptc (BDSC #44614), nub-GAL4 (BDSC #25754), FRT42D 2xUbi-374 GFP, UAS-smo:GFP (BDSC #44624), UAS-FLAG:smo^{Act} (BDSC #44621), UAS-wdp^{RNAi} (TRiP.HMC06302, BDSC #66004), UAS-wdp^{RNAi} (TRiP.HM05118, BDSC #28907), UAS-smo^{RNAi} 375 (TRiP.HMC03577, BDSC #53348), *hh-lacZ^{P30}* (a gift from Gary Struhl) (J. J. Lee et al., 1992), *dpp*-376 377 lacZ¹⁰⁶³⁸ (BDSC #12379) (Zecca, Basler, & Struhl, 1995), vas-Cas9 (BDSC #55821), esg-GAL4 (DGRC #113886) (S. Hayashi et al., 2002). The UAS-wdp, UAS-wdp^{ΔGAG}, UAS-Myc:wdp, UAS-Myc:wdp^{ΔLRRs}, 378 $UAS-Myc:wdp^{\Delta ICD}$, $wdp^{KO,\Delta CDS}$, $wdp^{KI,HA}$, $wdp^{KI,OLLAS}$ flies were generated in this study. A full list of 379 380 genotypes used in this study can be found in Table S1. 381 382 For constructing UAS-wdp, wdp CDS (corresponding to wdp-RA-E in FlyBase) was inserted into the 383 XhoI- and XbaI-digested pJFRC7 vector (a gift from Gerald Rubin; Addgene # 26220) (Pfeiffer et al., 384 2010) using NEBuilder HiFi DNA Assembly Master Mix (New England Biolabs [NEB], Ipswich, MA, E2621S). Similarly, wdp^{ΔGAG} (S282A, S334A, and S336A), Myc:wdp, Myc:wdp^{ΔLRRs}, and Myc:wd^{ΔLRRs}, and Myc:wd^{ΔLRRs}, and Myc:wd^{ΔLRRs}, and Myc:wd^{ΔLRRs}, and Myc:wd^{ΔLRRs}, and Myc:wd^{ΔLRRs}, and Myc^{ΔLRRs}, and Myc^{ΔLRRs}, and Myc^{ΔLRRs}, an 385 were inserted into the pJFRC7 vector. The UAS transgenic flies were generated using phiC31 integrase-386 387 mediated transgenesis at the ZH-86Fb attP (FBti0076525) integration site. Embryonic injection was 388 performed by BestGene Inc (Chino Hills, CA). Primers used in this study will be available upon request. 389 To generate the *wdp^{KO.ACDS}* allele, two sgRNAs (pU6-sgRNA-wdp-1 and pU6-sgRNA-wdp-2) were 390 391 introduced to delete the wdp CDS. To construct sgRNA plasmids, 5' -392 CTTCGACAGGGCCAACCAGGCGGTC - 3' and 5' - AAACGACCGCCTGGTTGGCCCTGTC - 3' 393 were annealed (pU6-sgRNA-wdp-1); and 5' - CTTCGAGTGGCCATTGATCACCTGG - 3' and 5' -394 AAACCCAGGTGATCAATGGCCACTC - 3' (pU6-sgRNA-wdp-2) were annealed and ligated in the 395 BbsI-digested pU6-BbsI-chiRNA plasmid (a gift from Melissa Harrison, Kate O'Connor-Giles, and Jill 396 Wildonger; Addgene #45946) (Gratz et al., 2013). A mixture of 50 ng/µl of pU6-sgRNA-wdp-1 and pU6-397 sgRNA-wdp-2 was injected into the embryos of the vas-Cas9 flies, which express Cas9 under the control 398 of the germline vasa regulatory elements (Gratz et al., 2014), by BestGene Inc. The wdp^{KO.ACDS} allele was 399 screened by PCR and verified by Sanger sequencing. 400 To generate the *wdp^{KLHA}* allele, we constructed a donor plasmid, which contained a Gly-Gly-Ser linker, 401 402 smGFP-HA, and approximately 1-kb homology arms to wdp flanking the linker and smGFP-HA, for 403 homology-directed repair. smGFP-HA and the *wdp* homology sequences on either side of the targeted 404 DSB were PCR-amplified from pJFRC201-10XUAS-FRT>STOP>FRT-myr:smGFP-HA (a gift from 405 Gerald Rubin; Addgene plasmid #63166) (Nern et al., 2015) and genomic DNA extracted from the vas-406 Cas9 flies, respectively. These fragments were cloned into the pHD-DsRed-attP backbone (a gift from 407 Melissa Harrison, Kate O'Connor-Giles and Jill Wildonger; Addgene #51019) (Gratz et al., 2014) using 408 NEBuilder HiFi DNA Assembly Master Mix (NEB, E2621S). Similarly, we generated a donor plasmid 409 with OLLAS tags amplified from pJFRC210-10XUAS-FRT>STOP>FRT-myr:smGFP-OLLAS (a gift 410 from Gerald Rubin; Addgene plasmid #63170) (Nern et al., 2015). A mixture of 50 ng/µl of pU6-sgRNA-411 wdp-2 and 125 ng/µl of each donor plasmid was injected into the vas-Cas9 embryos by BestGene Inc. The wdp^{KL,HA} and wdp^{KL,OLLAS} alleles were screened by PCR and verified by Sanger sequencing. 412 413 414 Flies were raised on a standard cornmeal fly medium at 25°C unless otherwise indicated. 415 416 417 **Mosaic analysis** 418

419 The *wdp^{KO. ACDS}* homozygous clones were generated by FLP/FRT-mediated mitotic recombination (T. Xu & Rubin, 1993). The FLP expression was induced by *nub-GAL4 UAS-FLP*.

- 421
- 422

423 Immunohistochemstry

424

425 Third-instar larval imaginal discs were stained as described previously (Takemura & Adachi-Yamada,

426 2011) with some modifications. Wing discs were dissected from third-instar wandering larvae in

427 phosphate-buffered saline (PBS, pH 7.4) and subsequently fixed in 3.7% formaldehyde in PBS for 15 min

428 at room temperature. After three 10-min washes with PBST (PBS containing 0.1% (vol/vol) Triton X-100

[Sigma, T8532]), the samples were incubated in primary antibodies overnight at 4°C. After three 10-min

washes with PBST, the samples were incubated with Alexa Fluor-conjugated secondary antibodies
 (1:500, Thermo Fisher Scientific) overnight at 4°C or 2 hours at room temperature. After three 10-min

431 (1.500, Thermo Fisher Scientific) overlight at 4 C of 2 hours at room temperature. After three ro-min 432 washes with PBST, the samples were stained with 1 μ g/ml DAPI (Thermo Fisher Scientific, 62248) and

432 washes with 1 DS1, the samples were standed with 1 µg/m DA11 (Thermo Fisher Scientific, 0
 433 subsequently mounted in VECTASHIELD Antifade Mounting Medium (Vector Laboratories.

Burlingame, CA, H-1000). F-actin was stained with Alexa Fluor 568 phalloidin (Thermo Fisher

435 Scientific, A12380). Adult midguts were dissected and immunostained as previously described

436 (Takemura & Nakato, 2017). Images were acquired on a LSM710 confocal microscope (Carl Zeiss,

437 Oberkochen, Germany). For quantification of Ptc staining, images were acquired with the same condition,

438 and fluorescence intensity was measured in a set area with Fiji (Schindelin et al., 2012).

439

440

441 Antibodies

442

443 The primary antibodies used were as follows: mouse anti-Ptc Apa 1 (1:20, Developmental Studies

444 Hybridoma Bank [DSHB], Iowa City, IA, deposited by Isabel Guerrero) (Capdevila et al., 1994), rat anti-

445 Ci 2A1 (1:20, DSHB, deposited by Robert Holmgren) (Motzny & Holmgren, 1995), chicken anti-β-

446 Galactosidase (1:2000, Abcam), mouse anti-En 4D9 (1:20, DSHB, deposited by Corey Goodman)

447 (Riggleman, Schedl, & Wieschaus, 1990), rabbit anti-pH3 (1:1000, Millipore, 06-570), rat anti-HA 3F10

448 (1:200, Roche, 11867423001), rabbit anti-HA C29F4 (1:1000, Cell Signaling, 3724), mouse anti-Smo

449 20C6 (1:50, DSHB, deposited by Philip Beachy) (Lum, Zhang, et al., 2003b), rabbit anti-pSmad3

450 (1:1000, Epitomics, 1880-1) (Smith, Machamer, Kim, Hays, & Marques, 2012), rabbit anti-Salm (1:30, a

gift from Scott Selleck), mouse anti-Dll 1:500 (1:500, a gift from Dianne Duncan) (D. M. Duncan,
Burgess, & Duncan, 1998), guinea pig anti-Sens (1:1000, a gift from Hugo Bellen) (Nolo, Abbott, &

453 Bellen, 2000), rabbit anti-cleaved Caspase-3 (1:200, Cell Signaling, 9661), rat anti-OLLAS L2 (1:500,

454 Novus Biologicals, NBP1-06713), mouse anti-Arm N2 7A1 (1:50, DSHB, deposited by Eric Wieschaus)

455 (Riggleman et al., 1990), mouse anti-Pros MR1A (1:50, DSHB, deposited by C.Q. Doe) (Campbell et al.,

456 1994), and mouse anti-Fas3 7G10 (1:50, DSHB, deposited by Corey Goodman) (Patel, Snow, &

457 Goodman, 1987). Alexa488, Alexa548, Alexa564 and Alexa633-conjugated secondary antibodies

458 (Thermo Fisher Scientific) were used at a dilution of 1:500.

459

460

461 Adult wing preparation

462

463 The left wings from female flies were dissected and mounted on slides using Canada balsam (Benz

464 Microscope, BB0020) as previously described (Takemura & Adachi-Yamada, 2011). Images were taken

using a ZEISS Stemi SV 11 microscope equipped with a Jenoptic ProgRes C3 digital camera.

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468

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- 476
- 477

478 Competing interests479

The authors declare no competing or financial interests.

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482 References

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- Akiyama, T., Kamimura, K., Firkus, C., Takeo, S., Shimmi, O., & Nakato, H. (2008). Dally regulates
 Dpp morphogen gradient formation by stabilizing Dpp on the cell surface. *Developmental Biology*,
 313(1), 408–419. http://doi.org/10.1016/j.vdbio.2007.10.035
- Baena-Lopez, L. A., Nojima, H., & Vincent, J.-P. (2012). Integration of morphogen signalling within the
 growth regulatory network. *Current Opinion in Cell Biology*, 24(2), 166–172.

489 http://doi.org/10.1016/j.ceb.2011.12.010

- Bandtlow, C. E., & Zimmermann, D. R. (2000). Proteoglycans in the developing brain: new conceptual
 insights for old proteins. *Physiological Reviews*, 80(4), 1267–1290.
- 492 http://doi.org/10.1152/physrev.2000.80.4.1267
- Bech-Hansen, N. T., Naylor, M. J., Maybaum, T. A., Sparkes, R. L., Koop, B., Birch, D. G., et al. (2000).
 Mutations in NYX, encoding the leucine-rich proteoglycan nyctalopin, cause X-linked complete
 congenital stationary night blindness. *Nature Genetics*, 26(3), 319–323. http://doi.org/10.1038/81619
- 495 congenital stationary night bindness. *Nature Genetics*, 26(5), 519–523. http://doi.org/10.1058/81619
 496 Bischof, J., Maeda, R. K., Hediger, M., Karch, F., & Basler, K. (2007). An optimized transgenesis system
 497 for Drosophila using germ-line-specific phiC31 integrases. *Proceedings of the National Academy of* 498 *Sciences of the United States of America*, 104(9), 3312–3317.
- 499 http://doi.org/10.1073/pnas.0611511104
- Briscoe, J., & Thérond, P. P. (2013). The mechanisms of Hedgehog signalling and its roles in
 development and disease. *Nature Reviews. Molecular Cell Biology*, 14(7), 416–429.
 http://doi.org/10.1038/nrm3598
- Calleja, M., Moreno, E., Pelaz, S., & Morata, G. (1996). Visualization of gene expression in living adult
 Drosophila. *Science (New York, NY)*, 274(5285), 252–255.
- Campbell, G., Göring, H., Lin, T., Spana, E., Andersson, S., Doe, C. Q., & Tomlinson, A. (1994). RK2, a
 glial-specific homeodomain protein required for embryonic nerve cord condensation and viability in
 Drosophila. *Development (Cambridge, England)*, *120*(10), 2957–2966.
- Capdevila, J., & Guerrero, I. (1994). Targeted expression of the signaling molecule decapentaplegic
 induces pattern duplications and growth alterations in Drosophila wings. *The EMBO Journal*, 13(19),
 4459–4468.
- Capdevila, J., Pariente, F., Sampedro, J., Alonso, J. L., & Guerrero, I. (1994). Subcellular localization of
 the segment polarity protein patched suggests an interaction with the wingless reception complex in
 Drosophila embryos. *Development (Cambridge, England)*, *120*(4), 987–998.
- 514 Coles, C. H., Shen, Y., Tenney, A. P., Siebold, C., Sutton, G. C., Lu, W., et al. (2011). Proteoglycan515 specific molecular switch for RPTPσ clustering and neuronal extension. *Science (New York, NY)*,
 516 332(6028), 484–488. http://doi.org/10.1126/science.1200840
- 517 Cortes, M., Baria, A. T., & Schwartz, N. B. (2009). Sulfation of chondroitin sulfate proteoglycans is
 518 necessary for proper Indian hedgehog signaling in the developing growth plate. *Development* 519 (*Cambridge, England*), 136(10), 1697–1706. http://doi.org/10.1242/dev.030742
- Denef, N., Neubüser, D., Perez, L., & Cohen, S. M. (2000). Hedgehog induces opposite changes in
 turnover and subcellular localization of patched and smoothened. *Cell*, 102(4), 521–531.
- 522 Desbordes, S. C., & Sanson, B. (2003). The glypican Dally-like is required for Hedgehog signalling in the
 523 embryonic epidermis of Drosophila. *Development (Cambridge, England)*, 130(25), 6245–6255.
 524 http://doi.org/10.1242/dev.00874
- Duncan, D. M., Burgess, E. A., & Duncan, I. (1998). Control of distal antennal identity and tarsal
 development in Drosophila by spineless-aristapedia, a homolog of the mammalian dioxin receptor.
 Genes & Development, 12(9), 1290–1303.
- Esko, J. D., & Zhang, L. (1996). Influence of core protein sequence on glycosaminoglycan assembly.
 Current Opinion in Structural Biology, 6(5), 663–670.
- Gradilla, A.-C., & Guerrero, I. (2013). Hedgehog on the move: a precise spatial control of Hedgehog
 dispersion shapes the gradient. *Current Opinion in Genetics & Development*, 23(4), 363–373.
 http://doi.org/10.1016/j.gde.2013.04.011

- 533 Gratz, S. J., Cummings, A. M., Nguyen, J. N., Hamm, D. C., Donohue, L. K., Harrison, M. M., et al.
- (2013). Genome engineering of Drosophila with the CRISPR RNA-guided Cas9 nuclease. *Genetics*,
 194(4), 1029–1035. http://doi.org/10.1534/genetics.113.152710
- Gratz, S. J., Ukken, F. P., Rubinstein, C. D., Thiede, G., Donohue, L. K., Cummings, A. M., & O'ConnorGiles, K. M. (2014). Highly specific and efficient CRISPR/Cas9-catalyzed homology-directed repair
 in Drosophila. *Genetics*, 196(4), 961–971. http://doi.org/10.1534/genetics.113.160713
- Groth, A. C., Fish, M., Nusse, R., & Calos, M. P. (2004). Construction of transgenic Drosophila by using
 the site-specific integrase from phage phiC31. *Genetics*, 166(4), 1775–1782.
- Hartl, T. A., & Scott, M. P. (2014). Wing tips: The wing disc as a platform for studying Hedgehog
 signaling. *Methods*, 68(1), 199–206. http://doi.org/10.1016/j.ymeth.2014.02.002
- Hayashi, S., Ito, K., Sado, Y., Taniguchi, M., Akimoto, A., Takeuchi, H., et al. (2002). GETDB, a
 database compiling expression patterns and molecular locations of a collection of Gal4 enhancer
 traps. *Genesis (New York, NY : 2000)*, 34(1-2), 58–61. http://doi.org/10.1002/gene.10137
- Hayashi, Y., Sexton, T. R., Dejima, K., Perry, D. W., Takemura, M., Kobayashi, S., et al. (2012).
 Glypicans regulate JAK/STAT signaling and distribution of the Unpaired morphogen. *Development* (*Cambridge, England*), 139(22), 4162–4171. http://doi.org/10.1242/dev.078055
- 549 Holt, C. E., & Dickson, B. J. (2005). Sugar Codes for Axons? *Neuron*, 46(2), 169–172.
- 550 http://doi.org/10.1016/j.neuron.2005.03.021
- Hu, Y., Flockhart, I., Vinayagam, A., Bergwitz, C., Berger, B., Perrimon, N., & Mohr, S. E. (2011). An
 integrative approach to ortholog prediction for disease-focused and other functional studies. *BMC Bioinformatics*, 12(1), 357. http://doi.org/10.1186/1471-2105-12-357
- Huff, J. L., Kingsley, K. L., Miller, J. M., & Hoshizaki, D. K. (2002). Drosophila windpipe codes for a
 leucine-rich repeat protein expressed in the developing trachea. *Mechanisms of Development*, 111(1-2), 173–176.
- Jia, J., Tong, C., Wang, B., Luo, L., & Jiang, J. (2004). Hedgehog signalling activity of Smoothened
 requires phosphorylation by protein kinase A and casein kinase I. *Nature*, 432(7020), 1045–1050.
 http://doi.org/10.1038/nature03179
- Kantor, D. B., Chivatakarn, O., Peer, K. L., Oster, S. F., Inatani, M., Hansen, M. J., et al. (2004).
 Semaphorin 5A is a bifunctional axon guidance cue regulated by heparan and chondroitin sulfate proteoglycans. *Neuron*, 44(6), 961–975. http://doi.org/10.1016/j.neuron.2004.12.002
- Kim, M.-S., Saunders, A. M., Hamaoka, B. Y., Beachy, P. A., & Leahy, D. J. (2011). Structure of the
 protein core of the glypican Dally-like and localization of a region important for hedgehog signaling.
 Proceedings of the National Academy of Sciences of the United States of America, 108(32), 13112–
 13117. http://doi.org/10.1073/pnas.1109877108
- Kudo, A., Shigenobu, S., Kadota, K., Nozawa, M., Shibata, T. F., Ishikawa, Y., & Matsuo, T. (2017).
 Comparative analysis of the brain transcriptome in a hyper-aggressive fruit fly, Drosophila
 prolongata. *Insect Biochemistry and Molecular Biology*, *82*, 11–20.
 http://doi.org/10.1016/j.jbmb.2017.01.006
- 571 Lander, A. D., & Selleck, S. B. (2000). The elusive functions of proteoglycans: in vivo veritas. *The* 572 *Journal of Cell Biology*, 148(2), 227–232.
- Lee, J. J., Kessler, von, D. P., Parks, S., & Beachy, P. A. (1992). Secretion and localized transcription
 suggest a role in positional signaling for products of the segmentation gene hedgehog. *Cell*, *71*(1),
 33–50.
- Lee, J.-S., & Chien, C.-B. (2004). When sugars guide axons: insights from heparan sulphate proteoglycan
 mutants. *Nature Reviews Genetics*, 5(12), 923–935. http://doi.org/10.1038/nrg1490
- Lin, D. M., & Goodman, C. S. (1994). Ectopic and increased expression of Fasciclin II alters motoneuron
 growth cone guidance. *Neuron*, *13*(3), 507–523.
- Lindahl, U., & Li, J.-P. (2009). Interactions between heparan sulfate and proteins-design and functional
 implications. *International Review of Cell and Molecular Biology*, 276, 105–159.
- 582 http://doi.org/10.1016/S1937-6448(09)76003-4

- Lu, J., Wang, D., & Shen, J. (2017). Hedgehog signalling is required for cell survival in Drosophila wing
 pouch cells. *Scientific Reports*, 7(1), 11317. http://doi.org/10.1038/s41598-017-10550-4
- Lum, L., Yao, S., Mozer, B., Rovescalli, A., Kessler, Von, D., Nirenberg, M., & Beachy, P. A. (2003a).
 Identification of Hedgehog pathway components by RNAi in Drosophila cultured cells. *Science (New York, NY)*, 299(5615), 2039–2045. http://doi.org/10.1126/science.1081403
- Lum, L., Zhang, C., Oh, S., Mann, R. K., Kessler, von, D. P., Taipale, J., et al. (2003b). Hedgehog signal
 transduction via Smoothened association with a cytoplasmic complex scaffolded by the atypical
 kinesin, Costal-2. *Molecular Cell*, *12*(5), 1261–1274.
- Matsumoto, Y., Irie, F., Inatani, M., Tessier-Lavigne, M., & Yamaguchi, Y. (2007). Netrin-1/DCC
 signaling in commissural axon guidance requires cell-autonomous expression of heparan sulfate. *The Journal of Neuroscience : the Official Journal of the Society for Neuroscience*, 27(16), 4342–4350.
 http://doi.org/10.1523/JNEUROSCI.0700-07.2007
- Momota, R., Naito, I., Ninomiya, Y., & Ohtsuka, A. (2011). Drosophila type XV/XVIII collagen, Mp, is
 involved in Wingless distribution. *Matrix Biology : Journal of the International Society for Matrix Biology*, 30(4), 258–266. http://doi.org/10.1016/j.matbio.2011.03.008
- Motzny, C. K., & Holmgren, R. (1995). The Drosophila cubitus interruptus protein and its role in the
 wingless and hedgehog signal transduction pathways. *Mechanisms of Development*, 52(1), 137–150.
- Mullor, J. L., Calleja, M., Capdevila, J., & Guerrero, I. (1997). Hedgehog activity, independent of
 decapentaplegic, participates in wing disc patterning. *Development (Cambridge, England)*, 124(6),
 1227–1237.
- Nakato, H., & Li, J.-P. (2016). Functions of Heparan Sulfate Proteoglycans in Development: Insights
 From Drosophila Models. *International Review of Cell and Molecular Biology*, 325, 275–293.
 http://doi.org/10.1016/bs.ircmb.2016.02.008
- Nern, A., Pfeiffer, B. D., & Rubin, G. M. (2015). Optimized tools for multicolor stochastic labeling reveal
 diverse stereotyped cell arrangements in the fly visual system. *Proceedings of the National Academy of Sciences of the United States of America*, 112(22), E2967–76.
- 609 http://doi.org/10.1073/pnas.1506763112
- Noborn, F., Gomez Toledo, A., Green, A., Nasir, W., Sihlbom, C., Nilsson, J., & Larson, G. (2016). Site specific identification of heparan and chondroitin sulfate glycosaminoglycans in hybrid
 proteoglycans. *Scientific Reports*, 6(1), 34537. http://doi.org/10.1038/srep34537
- Noborn, F., Gomez Toledo, A., Nasir, W., Nilsson, J., Dierker, T., Kjellén, L., & Larson, G. (2018).
 Expanding the chondroitin glycoproteome of Caenorhabditis elegans. *Journal of Biological Chemistry*, 293(1), 379–389. http://doi.org/10.1074/jbc.M117.807800
- Noborn, F., Gomez Toledo, A., Sihlbom, C., Lengqvist, J., Fries, E., Kjellén, L., et al. (2015).
 Identification of chondroitin sulfate linkage region glycopeptides reveals prohormones as a novel
 class of proteoglycans. *Molecular & Cellular Proteomics : MCP*, 14(1), 41–49.
 http://doi.org/10.1074/mcp.M114.043703
- Nolo, R., Abbott, L. A., & Bellen, H. J. (2000). Senseless, a Zn finger transcription factor, is necessary
 and sufficient for sensory organ development in Drosophila. *Cell*, 102(3), 349–362.
- O'Keefe, D. D., Thor, S., & Thomas, J. B. (1998). Function and specificity of LIM domains in Drosophila
 nervous system and wing development. *Development (Cambridge, England)*, *125*(19), 3915–3923.
- Olson, S. K., Bishop, J. R., Yates, J. R., Oegema, K., & Esko, J. D. (2006). Identification of novel
 chondroitin proteoglycans in Caenorhabditis elegans: embryonic cell division depends on CPG-1 and
 CPG-2. *The Journal of Cell Biology*, *173*(6), 985–994. http://doi.org/10.1083/jcb.200603003
- Patel, N. H., Martín-Blanco, E., Coleman, K. G., Poole, S. J., Ellis, M. C., Kornberg, T. B., & Goodman,
 C. S. (1989). Expression of engrailed proteins in arthropods, annelids, and chordates. *Cell*, 58(5),
 955–968.
- 630 Patel, N. H., Snow, P. M., & Goodman, C. S. (1987). Characterization and cloning of fasciclin III: a
- 631 glycoprotein expressed on a subset of neurons and axon pathways in Drosophila. *Cell*, 48(6), 975–
 632 988.

- Perrimon, N., & Bernfield, M. (2000). Specificities of heparan sulphate proteoglycans in developmental
 processes. *Nature*, 404(6779), 725–728. http://doi.org/10.1038/35008000
- Pfeiffer, B. D., Ngo, T.-T. B., Hibbard, K. L., Murphy, C., Jenett, A., Truman, J. W., & Rubin, G. M.
 (2010). Refinement of tools for targeted gene expression in Drosophila. *Genetics*, 186(2), 735–755. http://doi.org/10.1534/genetics.110.119917
- Poulain, F. E., & Yost, H. J. (2015). Heparan sulfate proteoglycans: a sugar code for vertebrate
 development? *Development (Cambridge, England)*, 142(20), 3456–3467.
 http://doi.org/10.1242/dev.098178
- 640 http://doi.org/10.1242/dev.098178
 641 Pusch, C. M., Zeitz, C., Brandau, O., Pesch, K., Achatz, J
- Pusch, C. M., Zeitz, C., Brandau, O., Pesch, K., Achatz, H., Feil, S., et al. (2000). The complete form of
 X-linked congenital stationary night blindness is caused by mutations in a gene encoding a leucine rich repeat protein. *Nature Genetics*, 26(3), 324–327. http://doi.org/10.1038/81627
- Ragkousi, K., & Gibson, M. C. (2014). Cell division and the maintenance of epithelial order. *The Journal of Cell Biology*, 207(2), 181–188. http://doi.org/10.1083/jcb.201408044
- Ren, W., Zhang, Y., Li, M., Wu, L., Wang, G., Baeg, G.-H., et al. (2015). Windpipe controls Drosophila
 intestinal homeostasis by regulating JAK/STAT pathway via promoting receptor endocytosis and
 lysosomal degradation. *PLoS Genetics*, 11(4), e1005180.
- 649 http://doi.org/10.1371/journal.pgen.1005180
- Ren, X., Yang, Z., Xu, J., Sun, J., Mao, D., Hu, Y., et al. (2014). Enhanced specificity and efficiency of
 the CRISPR/Cas9 system with optimized sgRNA parameters in Drosophila. *CellReports*, 9(3), 1151–
 1162. http://doi.org/10.1016/j.celrep.2014.09.044
- Riggleman, B., Schedl, P., & Wieschaus, E. (1990). Spatial expression of the Drosophila segment polarity
 gene armadillo is posttranscriptionally regulated by wingless. *Cell*, 63(3), 549–560.
- Saied-Santiago, K., & Bülow, H. E. (2018). Diverse roles for glycosaminoglycans in neural patterning.
 Developmental Dynamics : an Official Publication of the American Association of Anatomists,
 247(1), 54–74. http://doi.org/10.1002/dvdy.24555
- Schaefer, L., & Iozzo, R. V. (2008). Biological functions of the small leucine-rich proteoglycans: from
 genetics to signal transduction. *The Journal of Biological Chemistry*, 283(31), 21305–21309.
 http://doi.org/10.1074/jbc.R800020200
- Schindelin, J., Arganda-Carreras, I., Frise, E., Kaynig, V., Longair, M., Pietzsch, T., et al. (2012). Fiji: an
 open-source platform for biological-image analysis. *Nature Methods*, 9(7), 676–682.
 http://doi.org/10.1038/nmeth.2019
- Silver, J., & Miller, J. H. (2004). Regeneration beyond the glial scar. *Nature Reviews Neuroscience*, 5(2),
 146–156. http://doi.org/10.1038/nrn1326
- Smith, R. B., Machamer, J. B., Kim, N. C., Hays, T. S., & Marques, G. (2012). Relay of retrograde
 synaptogenic signals through axonal transport of BMP receptors. *Journal of Cell Science*, *125*(Pt 16),
 3752–3764. http://doi.org/10.1242/jcs.094292
- Strigini, M., & Cohen, S. M. (1997). A Hedgehog activity gradient contributes to AP axial patterning of
 the Drosophila wing. *Development (Cambridge, England)*, 124(22), 4697–4705.
- Tabata, T., & Takei, Y. (2004). Morphogens, their identification and regulation. *Development* (*Cambridge, England*), 131(4), 703–712. http://doi.org/10.1242/dev.01043
- Takei, Y. (2004). Three Drosophila EXT genes shape morphogen gradients through synthesis of heparan
 sulfate proteoglycans, *131*(1), 73–82. http://doi.org/10.1242/dev.00913
- Takemura, M., & Adachi-Yamada, T. (2011). Cell death and selective adhesion reorganize the
 dorsoventral boundary for zigzag patterning of Drosophila wing margin hairs. *Developmental Biology*, 357(2), 336–346. http://doi.org/10.1016/j.vdbio.2011.07.007
- Takemura, M., & Nakato, H. (2015). Genetic approaches in the study of heparan sulfate functions in
 Drosophila. *Methods in Molecular Biology (Clifton, N.J.), 1229*(Chapter 38), 497–505.
 http://doi.org/10.1007/978-1-4939-1714-3 38
- Takemura, M., & Nakato, H. (2017). Drosophila Sulf1 is required for the termination of intestinal stem
 cell division during regeneration. *Journal of Cell Science*, *130*(2), 332–343.
- 683 http://doi.org/10.1242/jcs.195305

- Tanimoto, H., Itoh, S., Dijke, ten, P., & Tabata, T. (2000). Hedgehog creates a gradient of DPP activity in
 Drosophila wing imaginal discs. *Molecular Cell*, 5(1), 59–71.
- Tavsanli, B. C., Ostrin, E. J., Burgess, H. K., Middlebrooks, B. W., Pham, T. A., & Mardon, G. (2004).
 Structure-function analysis of the Drosophila retinal determination protein Dachshund.
 Developmental Biology, 272(1), 231–247. http://doi.org/10.1016/j.vdbio.2004.05.005
- Townley, R. A., & Bülow, H. E. (2018). Deciphering functional glycosaminoglycan motifs in development. *Current Opinion in Structural Biology*, *50*, 144–154.
- 691 http://doi.org/10.1016/j.sbi.2018.03.011
- Toyoda, H., Kinoshita-Toyoda, A., & Selleck, S. B. (2000). Structural analysis of glycosaminoglycans in
 Drosophila and Caenorhabditis elegans and demonstration that tout-velu, a Drosophila gene related
 to EXT tumor suppressors, affects heparan sulfate in vivo. *The Journal of Biological Chemistry*,
 275(4), 2269–2275.
- Van Vactor, D., Wall, D. P., & Johnson, K. G. (2006). Heparan sulfate proteoglycans and the emergence
 of neuronal connectivity. *Current Opinion in Neurobiology*, *16*(1), 40–51.
 http://doi.org/10.1016/j.conb.2006.01.011
- Viswanathan, S., Williams, M. E., Bloss, E. B., Stasevich, T. J., Speer, C. M., Nern, A., et al. (2015).
 High-performance probes for light and electron microscopy. *Nature Methods*, *12*(6), 568–576.
 http://doi.org/10.1038/nmeth.3365
- Wang, J., Al-Ouran, R., Hu, Y., Kim, S.-Y., Wan, Y.-W., Wangler, M. F., et al. (2017). MARRVEL:
 Integration of Human and Model Organism Genetic Resources to Facilitate Functional Annotation of
 the Human Genome. *American Journal of Human Genetics*.
 http://doi.org/10.1016/j.ajhg.2017.04.010
- Wang, L., Dankert, H., Perona, P., & Anderson, D. J. (2008). A common genetic target for environmental
 and heritable influences on aggressiveness in Drosophila. *Proceedings of the National Academy of Sciences of the United States of America*, 105(15), 5657–5663.
 http://doi.org/10.1073/pnas.0801327105
- Whalen, D. M., Malinauskas, T., Gilbert, R. J. C., & Siebold, C. (2013). Structural insights into
 proteoglycan-shaped Hedgehog signaling. *Proceedings of the National Academy of Sciences of the* United States of America, 110(41), 16420–16425. http://doi.org/10.1073/pnas.1310097110
- Williams, E. H., Pappano, W. N., Saunders, A. M., Kim, M.-S., Leahy, D. J., & Beachy, P. A. (2010).
 Dally-like core protein and its mammalian homologues mediate stimulatory and inhibitory effects on
 Hedgehog signal response. *Proceedings of the National Academy of Sciences of the United States of America*, 107(13), 5869–5874. http://doi.org/10.1073/pnas.1001777107
- Xu, D., & Esko, J. D. (2014). Demystifying heparan sulfate-protein interactions. *Annual Review of Biochemistry*, 83(1), 129–157. http://doi.org/10.1146/annurev-biochem-060713-035314
- Xu, T., & Rubin, G. M. (1993). Analysis of genetic mosaics in developing and adult Drosophila tissues.
 Development (Cambridge, England), 117(4), 1223–1237.
- Yan, D., Wu, Y., Yang, Y., Belenkaya, T. Y., Tang, X., & Lin, X. (2010). The cell-surface proteins
 Dally-like and Ihog differentially regulate Hedgehog signaling strength and range during
 development. *Development (Cambridge, England)*, 137(12), 2033–2044.
- 724 http://doi.org/10.1242/dev.045740
- Zecca, M., Basler, K., & Struhl, G. (1995). Sequential organizing activities of engrailed, hedgehog and
 decapentaplegic in the Drosophila wing. *Development (Cambridge, England)*, 121(8), 2265–2278.
- Zecca, M., Basler, K., & Struhl, G. (1996). Direct and long-range action of a wingless morphogen
 gradient. *Cell*, 87(5), 833–844. http://doi.org/10.5167/uzh-996
- Zhang, F., McLellan, J. S., Ayala, A. M., Leahy, D. J., & Linhardt, R. J. (2007). Kinetic and structural
 studies on interactions between heparin or heparan sulfate and proteins of the hedgehog signaling
 pathway. *Biochemistry*, 46(13), 3933–3941. http://doi.org/10.1021/bi6025424
- 732 Zhu, A. J., Zheng, L., Suyama, K., & Scott, M. P. (2003). Altered localization of Drosophila Smoothened
- 733 protein activates Hedgehog signal transduction. *Genes & Development*, *17*(10), 1240–1252.
- 734 http://doi.org/10.1101/gad.1080803





738 Figure 1. Identification of Wdp as a novel CSPG in *Drosophila*. (A) A scheme for identifying CSPGs

in *Drosophila*. The workflow includes the enrichment of proteoglycans from fly extract, enzymatic

- hydrolysis and subsequent analysis and interpretation of mass spectra. (**B** and **C**) MS2 fragment mass
- spectra of Wdp protein (UniProt: Q9W266) showing two unique CS-glycopeptides. (B) Peptide
- 742 (SDQVEGSGDLSETNMELK) identified with one hexasaccharide structure and one methionine
- oxidation (m/z 983.38; 3+) (C) Peptide (EEHIVKDEDEDDEGSGSGGGLLIIPDPSK) identified with two
- hexasaccharide structures where one of the hexasaccharides were modified with one phosphate
- modification (m/z 1276.76; 4+). The asterisk denotes the second isotopic peak.



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748 Figure 2. Overexpression of *wdp* reduces the Hh-signaling-active domain. (A–F) Control wing discs

(A, C, and E) and wing discs overexpressing wdp with ap-GAL4 (ap>GFP+wdp) (B, D, and F) were

immunostained for the expression of Ptc, Ci (A and B), *dpp-lacZ* (C and D), En, and *hh-lacZ* (E and F).

The expression domains of Ptc, Ci, and *dpp-lacZ* were reduced by *wdp* overexpression in the dorsal

compartment compared to those in the ventral compartment. En expression induced by high-level Hh

signaling in the anterior compartment is diminished by *wdp* overexpression. Note that the *hh-lacZ*

expression is not affected by *wdp* overexpression. Nuclei were stained with DAPI (C and D). Anterior to

755 the left; dorsal to the top. Scale bars: 50 μ m.



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Figure 3. Wdp negatively regulates Hh signaling in a GAG-dependent manner. (A) A schematic

drawing of wild-type Wdp and a mutant form of Wdp (Wdp^{ΔGAG}). (**B–D**) A control adult wing (B) and adult wings expressing *UAS-wdp* (C) or *UAS-wdp^{\Delta GAG}* (D) with *Bx^{MS1096}-GAL4*. (**E** and **F**) Wing discs expressing *UAS-wdp^{\Delta GAG}* with *ap-GAL4* were immunostained for Ptc, Ci (E) and *dpp-lacZ* (F). (**G–I**) Wing discs expressing *UAS-3xMyc:wdp* (G), *UAS-3xMyc:wdp^{\Delta LRRs}* (H), and *UAS-3xMyc:wdp^{\Delta ICD}* (I) were</sup>

immunostained for Ptc. Nuclei were stained with DAPI. Scale bars: 50 µm.



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Figure 4. Wdp expression in the wing disc. (A) A schematic of CRISPR–Cas9-mediated gene editing of 768 769 wdp for generating wdp^{KLHA}. Ten copies of an HA epitope tag (smGFP-HA) were inserted in frame near 770 the stop codon of the *wdp* coding sequence (CDS). Only the last exon is shown. CDS, the black box; 771 smGFP-HA, the magenta box; LHA, left homology arm; RHA, right homology arm; Cas9 target site, the open triangle. (**B–D**) Wing discs homozygous for $wdp^{KI.HA}$ were stained with Alexa Fluor 568-conjugated 772 phalloidin (F-actin), anti-phospho histone H3 antibody (pH3, mitotic nuclei), and anti-HA antibody. 773 774 Apical (B) and basal (C) sections of the same disc are shown. Intense staining of Wdp:HA was observed on the basal side of wing disc epithelium (C-C'''). An optical cross section shows the accumulation of 775 776 Wdp:HA in the basal projection of apically translocating mitotic cells (D-D'''). (E) Wdp:HA is not detectable in the dorsal compartment of a wing disc expressing wdp^{RNAi} (TRiP.HMC06302) with ap-777

778 *GAL4*. Nuclei were stained with DAPI (D and E). Scale bars: 50 μm.





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782 Figure 5. Loss of wdp leads to increased Hh signaling. (A) A wing disc expressing UAS-wdp^{RNAi} 783 (TRiP.HMC06302) with *ap-GAL4* was immunostained for Ptc and Ci. (B and C) Signal intensity plots of 784 the Ptc expression in the dorsal compartment (red) and ventral compartment (blue) in wing discs expressing UAS-FLP (B) or UAS-wdp^{RNAi} (C). Solid lines indicate the average intensity of Ptc staining 785 and shaded areas show the standard error of the mean. (**D**) A wing disc expressing UAS-wdp^{RNAi} with ap-786 787 GAL4 was immunostained for dpp-lacZ. Nuclei were stained with DAPI. (E) A schematic of the generation of a wdp loss-of-function allele ($wdp^{KO.ACDS}$) lacking most of the wdp CDS using the CRISPR-788 Cas9 system. (F) A PCR-based genotyping result for the wild-type (WT) and $wdp^{KO.\Delta CDS}$ allele (KO) 789 using a primer set shown as cyan arrows in E. (G) Genomic sequence of the wdp endogenous locus 790 791 targeted by CRISPR-Cas9 to delete most of the wdp CDS. A small insertion is shown in green. (H) Somatic mosaic clones of *wdp^{KO}* were induced in the wing pouch using *nub-GAL4 UAS-FLP*. 792 Homozygous wdp^{KO} mutant cells are marked by loss of GFP. Increased Ptc expression was observed in 793 794 wdp mutant clones (yellow arrows) (I, J) A control adult wing (I) and a wing expressing UAS-wdp^{RNAi} 795 (TRiP.HM05118) (J) by nub-GAL4. Scale bars: 50 µm. 796



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Figure 6. Wdp negatively regulates Smo cell surface accumulation. (A, B) A control wing disc and a wing disc expressing UAS-wdp^{RNAi} (TRiP.HMC06302) with *ap*-GAL4 were immunostained for the

expression of Smo. Nuclei were stained with DAPI. Scale bars: 50 µm. (C) A model for how Wdp modulates Hh signaling. Wdp inhibits Smo cell surface accumulation thereby reducing Hh signaling activity.





Figure S1. Effect of *wdp* **overexpression on Dpp and Wg signaling.** (A–F) Control wing discs (A and D) and wing discs expressing *UAS-wdp* (B and E) or *UAS-wdp*^{AGAG} (C and F) with *ap-GAL4* were immunostained for pMad (A–C) and Salm (D–F) as read-outs of Dpp signaling. (G–L) Control wing discs (G and J) and wing discs expressing *UAS-wdp* (H and K) or *UAS-wdp*^{AGAG} (I and L) with *hh-GAL4* were immunostained for Dll (G–I) and Sens (J–L) as read-outs of Wg signaling (Nolo et al., 2000; Zecca, Basler, & Struhl, 1996). Nuclei were stained with DAPI. Scale bars: 50 µm.



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816 Figure S2. Overexpression of *wdp* in the wing disc induces apoptosis. (A–C) A control wing disc (A)

and wing discs expressing UAS-wdp (B) or UAS-wdp^{ΔGAG} (C) with ap-GAL4 were immunostained for 817

cleaved Caspase-3 as a marker of apoptotic cells. (D) An optical cross section of a wing disc expressing 818

UAS-wdp with ap-GAL4. Signals for cleaved Caspase-3 (cvan) and F-actin (magenta) are shown. (E) A 819

wing disc expressing UAS-smo^{RNAi} (TRiP.HMC03577) with *ap-GAL4* was immunostained for Ptc. (F and **G**) Wing discs expressing UAS-smo^{RNAi} (F) or UAS-ptc (G) with *ap-GAL4* were immunostained for 820

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822 cleaved Caspase-3. (H) A wing disc coexpressing UAS-wdp and UAS-FLAG:smo^{Act} was immunostained

823 for cleaved Caspase-3. Nuclei were stained with DAPI. Scale bars: 50 µm.

| | | SP | LRRs | GAG attachment sites | | ТМ |
|------------|---------------------------------------|----|------|----------------------|--|----|
| | 3xMyc:Wdp | | | | | |
| | 3xMyc:Wdp ^{∆GAG} | | | | | |
| | 3xMyc:Wdp ^{∆LRR} | | | | | |
| 825 826 | <mark>3xMyc:Wdp^{∆ICD}</mark> | | | | | |

- 827 Figure S3. A schematic of 3xMyc-tagged wild-type and mutant Wdp constructs. The
- magenta box denotes 3 copies of a Myc tag. The cyan box indicates the signal peptide (SP). The
- 829 yellow box shows the LRR motifs. The red line indicates the CS attachment sites. The green box
- 830 shows the transmembrane domain (TM).

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834 Figure S4. Expression patterns of Wdp in the eye disc, adult posterior midgut, ovary and testis. (A, **B**) Apical (A) and basal (B) sections of a wing disc homozygous for $wdp^{KLOLLAS}$ was stained with anti-835 OLLAS antibody. (C-H) An eye disc, adult midgut, ovary, and testis with one or two copies of wdp^{KLHA} 836 were immunostained with anti-HA antibody (magenta). (C, D) Apical (C) and basal (D) sections of an eye 837 disc. Neurons are marked by *elav-GAL4 UAS-mCD8;GFP* (*elav>mCD8;GFP*, green). (E) An adult 838 839 midgut showing ISCs/enteroblasts (escargot-GAL4 UAS-GFP [esg>GFP]), enteroendocrine cells 840 (nuclear Prospero [Pros]), cell membrane (Armadillo [Arm]), and Wdp:HA. (F) An optical cross section 841 of an adult midgut showing ISCs/enteroblasts (esg>GFP) and F-actin (stained by phalloidin-Alexa568). 842 (G) Wdp:HA expression is detected in the egg chambers (stage 6 and later) in the ovary. Wdp:HA is also detectable in the tracheal system. (H) Wdp:HA expression is detected in the sheath cells and the tracheal 843 844 system in the testis. Nuclei were stained with DAPI. Scale bars: 50 µm (A, B, C, D, and G), 25 µm (E, F, 845 and H).

| Figure | Panel | Genotype |
|--------|---------|--|
| 2 | Α | ap-GAL4 UAS-GFP/+ |
| 2 | В | ap-GAL4 UAS-GFP/+; UAS-wdp/+ |
| 2 | С | ap-GAL4 UAS-GFP dpp-lacZ/+ |
| 2 | D | ap-GAL4 UAS-GFP dpp-lacZ/+; UAS-wdp/+ |
| 2 | Е | ap-GAL4 UAS-GFP hh-lacZ/+ |
| 2 | F | ap-GAL4 UAS-GFP hh-lacZ/+; UAS-wdp/+ |
| 3 | В | Bx^{MS1096} - $GAL4/+$ |
| 3 | С | Bx ^{MS1096} -GAL4/+; UAS-wdp/+ |
| 3 | D | Bx^{MS1096} -GAL4/+; UAS-wdp ^{ΔGAG} |
| 3 | Е | ap-GAL4 UAS-GFP/+; UAS-wdp ^{4GAG} /+ |
| 3 | F | ap-GAL4 UAS-GFP dpp-lacZ/+; UAS-wdp ^{ΔGAG} /+ |
| 3 | G | ap-GAL4 UAS-GFP/+; UAS-3xMyc:wdp/+ |
| 3 | Н | ap-GAL4 UAS-GFP/+; UAS-3xMyc:wdp ^{ALRRs} /+ |
| 3 | Ι | ap-GAL4 UAS-GFP/+; UAS-3xMyc:wdp ^{AICD} /+ |
| 4 | B, C, D | wdp ^{KI.HA} /wdp ^{KI.HA} |
| 4 | E | ap-GAL4 UAS-GFP wdp ^{KI.HA} /UAS-wdp ^{RNAi.HMC06302} |
| 5 | A, C | ap-GAL4 UAS-GFP/UAS-wdp ^{RNAi.HMC06302} |
| 5 | В | ap-GAL4 UAS-GFP/UAS-wdp ^{RNAi.HMC06302} |
| 5 | D | ap-GAL4 UAS-GFP dpp-lacZ/UAS-wdp ^{RNAi.HMC06302} |
| 5 | Н | nub-GAL4, FRT42D wdp ^{KO. ACDS} /FRT42D 2xubi-GFP; UAS-FLP/+ |
| 5 | Ι | nub-GAL4/+ |
| 5 | J | nub-GAL4/+; UAS-wdp ^{RNAi.HMS05118} /+ |
| 6 | Α | ap-GAL4 UAS-GFP/+ |
| 6 | В | ap-GAL4 UAS-GFP/UAS-wdp ^{RNAi.HMC06302} |
| S1 | A, D | ap-GAL4 UAS-GFP/+ |
| S1 | В, Е | ap-GAL4 UAS-GFP/+; UAS-wdp/+ |
| S1 | C, F | ap-GAL4 UAS-GFP/+; UAS-wdp ^{AGAG} /+ |
| S1 | G, J | hh-GAL4 UAS-tdTomato/+ |
| S1 | Н, К | hh-GAL4 UAS-tdTomato/UAS-wdp |
| S1 | I, L | hh-GAL4 UAS-tdTomato/UAS-wdp ^{AGAG} |
| S2 | A, D | ap-GAL4 UAS-GFP/+ |
| S2 | В | ap-GAL4 UAS-GFP/+; UAS-wdp/+ |
| S2 | С | ap-GAL4 UAS-GFP/+; UAS- wdp ^{AGAG} /+ |
| S2 | E, F | ap-GAL4 UAS-GFP/UAS-smo ^{RNAi,HMC03577} |
| S2 | G | ap-GAL4 UAS-GFP/UAS-ptc |
| S2 | Н | ap-GAL4 UAS-GFP/UAS-FLAG:smo ^{Act} ; UAS-wdp/+ |
| S4 | A, B | wdp ^{KI.OLLAS} /wdp ^{KI.OLLAS} |
| S4 | C, D | elav-GAL4 UAS-mCD8:GFP/+; wdp ^{KLHA} /+ |
| S4 | E, F | esg-GAL4 UAS-GFP/+; wdp ^{KI.HA} /+ |
| S4 | G, H | wdp ^{KLHA} /wdp ^{KLHA} |

846 Table S1. Genotypes of *Drosophila* strains used in each figure.

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