Ubiquitylation-independent activation of Notch signalling by Delta

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Abstract Ubiquitylation (ubi) by the E3-ligases Mindbomb1 (Mib1) and Neuralized (Neur) is required for activation of the DSL ligands Delta (Dl) and Serrate (Ser) to activate Notch signalling. These ligases transfer ubiquitin to lysines of the ligands’ intracellular domains (ICDs), which sends them into an Epsin-dependent endocytic pathway. Here, we have tested the requirement of ubi of Dl for signalling. We found that Dl requires ubi for its full function, but can also signal in two ubi-independent modes, one dependent and one independent of Neur. We identified two neural lateral specification processes where Dl signals in an ubi-independent manner. Neur, which is needed for these processes, was shown to be able to activate Dl in an ubi-independent manner. Our analysis suggests that one important role of DSL protein ubi by Mib1 is their release from cis-inhibitory interactions with Notch, enabling them to trans-activate Notch on adjacent cells.

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Introduction

Signalling through the Notch signalling pathway is required in many developmental processes of probably all metazoans (Bray, 2006; Fiuza and Arias, 2007). Notch receptors are hetero-dimeric type 1 trans-membrane proteins, which are activated through ligands of the Delta/Serrate/Lag2 (DSL) protein family. The genome of Drosophila contains two DSL ligands, Delta (Dl) and Serrate (Ser), which are structurally related in their extracellular domain (ECD). Their binding elicits two consecutive proteolytic cleavages of Notch performed by Kuzbanian (Kuz)/ADAM10 and the γ-secretase complex, which results in the release of the intracellular domain (NICD) into the cytoplasm. The first, Kuz-mediated cleavage (S2-cleavage) occurs in the extracellular juxta-membrane region and removes the ecto-domain (NECD). The shedding of NECD triggers the second cleavage within the transmembrane domain by the γ-secretase (S3-cleavage), allowing the release of NICD. NICD enters the nucleus and associates with the CSL transcription factor Suppressor of Hairless (Su(H)) and co-factors to activate the transcription of target genes. In the absence of NICD, Su(H) acts as a repressor of transcription in association with Hairless (H) (Barolo et al., 2002; Brou et al., 1994).

A number of studies have demonstrated the importance of endocytosis for Notch signalling and regulation of its activity (Le Borgne et al., 2005a). During signalling, the ligands must be endocytosed in the signal-sending cell to activate Notch in the signal-receiving one. Two membrane-associated E3-ligases, Neuralized (Neur) and Mindbomb1 (Mib1), play a crucial role in the activation of the ligands and their endocytosis (Le Borgne et al., 2005a). E3-ligases mediate the ubiquitylation (ubi) of specific substrates at lysines (Ks). This label constitutes a common signal to initiate endocytosis...
Two enzymes known as Mindbomb1 and Neuralized activate Delta and Serrate. Both enzymes add a small unit called ubiquitin to specific locations on the signal proteins, but the effect that ubiquitin has on Notch signalling is not yet fully understood.

Berndt, Seib, Kim et al. have now examined fruit flies that had a variety of genetic mutations. These included some flies that could produce mutant versions of the Serrate and Delta proteins that lacked the locations to which ubiquitin normally attaches. The results of the experiments reveal that Delta requires ubiquitin, Mindbomb1 and Neuralized to work at full capacity. However, Delta could still perform some of its roles without ubiquitin. Neuralized and Delta can partner up to send some signals independently of ubiquitin, and Delta can even send some signals on its own. Serrate, on the other hand, does not work at all without ubiquitin.

The results presented by Berndt et al. help us to understand the role that ubiquitin plays in activating Notch signalling. Further work that builds on these findings could help to shed light on how uncontrolled Notch activation can contribute to a variety of diseases, including cancer, cardiovascular diseases and multiple sclerosis.

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the NECD. In the second model (recycling model), ligand signalling activity is thought to depend on events taking place after internalisation of the ligands into endosomes, such as enzymatic processing of the ligand into the active form or packaging into exosomes: accordingly, signalling activity would require recycling of the ligands to the cell surface to be able to activate Notch. Previous work has provided evidence for both models (reviewed in [Weinmaster and Fischer, 2011]).

Although Mib1 and Neur are clearly required for DSL signalling, their role is not entirely understood: (1) While endocytosis of Ser is virtually abolished in mib-1 mutant cells, that of DI is not obviously affected, although both Ser and DI function is impaired (Wang and Struhl, 2004). (2) Recent work suggests that Neur might have a function that is separable from its ligase function and required for endocytosis (Skwarek et al., 2007). (3) The phenotype of mib-1 mutants is milder than that of mutants of other genes involved in Notch signalling. These results raise the possibility that one or both ligands might possess an activity that is independent of ubi.

In addition to activating Notch in adjacent cells (trans-activation), the ligands engage in inhibitory interaction with Notch molecules present in the same cell (Klein et al., 1997; Micchelli et al., 1997). This cis-inhibition occurs if DSL ligands are expressed above a threshold level. The underlying mechanism is not well understood, but cis-inhibition can be suppressed by co-expression of Notch, indicating that the ratio between Notch and ligand concentration in a cell is an important parameter (Klein et al., 1997). Moreover, it has been shown that the ECD of the ligands is involved (Glittenberg et al., 2006). Whether the ICD is involved, or cis-inhibition is influenced by Mib1 and Neur is not established, although there is evidence that enhanced/diminished ligand endocytosis relieves/aggravates (respectively) cis-inhibition of Ser and DI (Glittenberg et al., 2006).

Here, we tested the activity of a DI variant (termed DIK2R) in which all 12 Ks of its ICD were replaced by structurally related arginines (Rs). We show that DIK2R possesses signalling activity. However, its activity is reduced compared to DI, indicating that ubi is required for the full function of DI. The reduction in activity is in part caused by increased cis-inhibitory activity of DIK2R indicating that the ICD and its ubi contribute to the degree/strength of cis-inhibition of a ligand. We found that an important function of Mib1 during normal signalling is the release of the ligands from cis-inhibitory interaction with Notch. Our results revealed that DI can signal in two ubi-independent modes, one dependent and one independent of Neur. The results also indicate that Neur and Mib1 have different mechanisms to activate DI. In contrast to Mib1, Neur can promote DI signalling in an ubi-independent manner. We identified neuron sibling specification during larval neurogenesis and also specification of the sensory organ precursor of the bristle sensillum as naturally occurring instances of DI/Notch signalling, where DI requires Neur, but ubi seems to be dispensable. Finally, we found an instance of DI induced Notch signalling that is independent of both Neur and ubi.

Results

mib-1 mutants exhibit residual Notch pathway activity, which depends on DI

During wing development of Drosophila interactions between dorsal and ventral cells mediated by the Notch pathway at the dorso-ventral (D/V) compartment boundary result in the establishment and maintenance of a stripe of expression of Wingless (Wg) that straddles the D/V boundary of the wing disc at the end of the third larval instar stage (Figure 1A, arrow) (Klein, 2001). During the initial phase, dorsal boundary cells signal to ventral boundary cells to activate Wg expression and to increase expression of DI (Micchelli et al., 1997; de Celis and Bray, 1997; de Celis et al., 1996; Troost and Klein, 2012). The ventral boundary cells signal back through DI to activate expression of Wg in dorsal cells (Figure 1B, step 1). After establishment, expression of Wg in boundary cells is maintained through a feedback loop (DS-loop, Figure 1B steps 2 and 3): it involves the induction of expression of DI and Ser in cells adjacent to the boundary cells by Wg signalling (Figure 1B step 2). Wg expression at the D/V boundary is then maintained by back signalling from these cells to boundary cells through the Notch ligands (Figure 1B step 3). Loss of the activity of Notch results in the loss of expression of Wg along the D/V boundary and a reduction of the diameter of its ring-like domains of expression in the proximal wing. The reduction depends on the degree of loss of Notch pathway activity. Strong loss of activity, as observed in mutants of the γ-secretase component Psn, reduces the inner ring-like domain to a spot or causes its complete absence (Figure 1C, arrowhead).
Figure 1. DI-induced activity of Notch in mib1 mutant wing imaginal discs. (A) Expression of Wg in a normal wing disc. (B) Establishment of the Wg expression domain along the DV boundary by Notch signalling. For further explanation, see text. (C–E) Expression of Wg in mutants of genes involved in Notch signalling. (C) Psn+, (D) Ser- and (E) mib1-mutant wing imaginal disc. The arrow in (A) points to the expression along the D/V boundary that is dependent on Notch signalling. It is lost in the mutants. The arrowhead in (A, C–E) points to the inner ring-like expression domain of Wg. Its diameter

Figure 1 continued on next page
is a measure of the loss of Notch pathway activity. (F) Expression of Gbe + Su(H) in a wildtype disc. (G–I) Expression of Wg (G, I), Gbe + Su(H) (H, I) in a mib1 mutant disc of the early third instar stage. The arrow highlights the expression along the D/V boundary. (J, K) Expression of vgBE (J, K) and Wg (K) in a mib1 mutant disc of the early third instar stage. The arrow points to the expression of vgBE and Wg along the D/V boundary. (L–O) Expression of Gbe + Su(H) in the notum of a wildtype (L), mib1 (M), and Notch depleted disc (N, O). The arrow highlights the S3 expression domain of Gbe + Su(H). S3 is present in mib1 mutants (M, arrow), but absent in Notch depleted nota (O, the arrow points to the expected location of S3).

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Figure supplement 1. Residual Dl induced Notch activity in mib1 mutant wing imaginal discs.
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(Klein and Martinez-Arias, 1998; Koelzer and Klein, 2006; Struhl and Greenwald, 1999a; Ye et al., 1999). Weaker loss of Notch activity, as observed in Ser mutants (Klein and Martinez-Arias, 1998), causes a reduction in diameter of the domains (Figure 1D, arrowhead).

The phenotype of mib1 null mutant wing discs resembles that of Ser mutants (Figure 1E, arrowhead; compare with D). We previously showed that residual activity of Dl is responsible for the weaker phenotype of Ser mutants (Klein and Martinez-Arias, 1999). The resemblance prompted us to search for residual Notch pathway activity in mib1 null mutants. We found that the Notch pathway is indeed weakly active. The Notch activity reporter Gbe + Su(H) was initially expressed along the D/V boundary in early third larval instar stages of mib1 mutant wing discs and lost only later in development (Figure 1F–I, arrow). Also the vestigial boundary enhancer (vgBE) (Williams et al., 1994), a direct endogenous target of the Notch pathway, was expressed in early wing discs (Figure 1J,K, arrow).

In the notum Gbe + Su(H) is expressed in four stripes (Figure 1L). We observed residual expression of stripe 3 in mib1 mutants (S3; Figure 1L,M, arrow). S3 expression is dependent on Notch signalling, since depletion of the activity of Notch by expression of N-RNAi in the notum with ciGal4 resulted in the complete abolishment of its expression (Figure 1N,O, arrow in O). Together, these results confirm that residual ligand-dependent Notch activity is present in mib1 mutants.

Hairless (H) encodes a member of the transcriptional repressor complex assembled around Su(H) in the absence of Notch activity to repress expression of the target genes of the pathway (Barolo et al., 2002). Loss of its function results in de-repression of a subset of target genes and enhancement of residual Notch signalling in Ser mutants, which is sufficient to re-establish the expression of Wg and Gbe + Su(H) along the D/V boundary of the wing disc (Troost and Klein, 2012; Klein et al., 2000). We observed that, as in the case of Ser mutants, concomitant loss of H function partly restores the expression of Wg and maintains that of Gbe + Su(H) along the D/V boundary in mib1 mutants (Figure 1—figure supplement 1A–C, arrow). The comparison with the expression of the dorsal selector Apterous (Ap) indicated that the expression of Gbe + Su(H) was restricted to dorsal boundary cells (Figure 1—figure supplement 1D–F). Since Ser is unable to activate the Notch pathway in dorsal cells because of the activity of Fng (Panin et al., 1997), this observation strongly suggests that Dl signalling is responsible for the re-appearance of the expression of the Notch target genes in mib1 H double mutants.

The results indicate that a residual activity of the Notch pathway, probably induced by Dl, is present in the absence of mib1 function. Since Mib1 is the only known E3-ligase for Notch ligands known to be involved during wing development, the results raise the possibility that Dl has an activity that is independent of ubi. Although it has been shown that Mib1 is absent in the discs mutant for the mib1 null alleles used here (Le Borgne et al., 2005), it might be possible that a long-lasting maternal component of mib1 exists that provides cells of the imaginal disc with residual Mib1 activity. It is also possible that another unidentified E3-ligase contributes to DI signalling. These issues are addressed in the following.

Activation of the Notch pathway by Delta in the absence of mib1

In the experiments described in the following, we expressed a set of UAS Dl constructs with ptcGal4 to test their activity in the wing imaginal disc. If not stated otherwise, the constructs are inserted in the attP landing site 51C (Flybase: M(3xP3-RFP.attP’)ZH-51C) to achieve similar expression levels and allow direct comparison (Bischof et al., 2007). All constructs are HA-tagged at their C-terminus.
ptcGal4 is expressed on the anterior side of the anterior-posterior boundary (A/P boundary) in a gradient that increases towards the posterior. Its posterior expression boundary coincides with the A/P boundary (Figure 2A–C, arrow in B). Continuous expression of DI-HA with ptcGal4 at 25°C induces ectopic expression of the Notch target gene Wg and the more sensitive Notch activity reporter Gbe + Su(H) in two stripes perpendicular to their normal expression along the D/V boundary, as has been reported for untagged versions (Doherty et al., 1996) (Figure 2D–F, white arrows in D, E). The anterior stripe (a in Figure 2E) is located in the anterior region of low expression, while the posterior stripe (p in Figure 2E) is adjacent to the ptc domain in posterior non-expressing boundary cells (Figure 2F). Wg is not induced in regions of high DI expression close to the A/P boundary (Figure 2D,E, yellow arrow, F), because of cis-inhibition of Notch by high DI concentrations (Doherty et al., 1996). The cis-inhibition also interrupts expression of endogenous Wg along the D/V boundary (Figure 2D,E, white and yellow arrows, F).

If DI is active in mib1 mutants, ectopic expression of DI-HA should induce ectopic activation of Notch in this mutant (Figure 2G–J). Indeed, expression of DI-HA in mib1 mutants caused weak and diffuse ectopic expression of Wg in the mib1 mutant discs (100% penetrance, n = 7; Figure 2I, arrow). In addition, the diameter of the ring-like domains of Wg is increased (Figure 2I, arrowhead, compare with H, arrowhead). Expression of the more sensitive Gbe + Su(H) was induced throughout the ptcGal4 expression domain, even in the notum where, in contrast to the wing, Notch activity does not induce expression of the ligands (Figure 2J, arrow and arrowhead). DI-HA was not able to induce ectopic expression of the activity reporters in Psn mutants, in which the pathway is interrupted on the signal-receiving side (Figure 2—figure supplement 1A–C). To rule out that residual activity of Mib1 supports DI signalling in the mutants, we generated a DI variant where the core of the Mib1 binding box in its ICD (ICD2, see Figure 3A, [Daskalaki et al., 2011]) is mutated to alanine (DI-HA2 ala). This variant induced activity of the Notch pathway in the mib1 mutant background in a manner comparable to DI-HA throughout the ptcGal4 domain (Figure 2KL, arrow and arrowhead, compare with I, J). This finding indicates that DI can activate the Notch pathway in a mib1 independent manner and strongly argues against a residual Mib1 activity in the mutants.

The expression of a Ser-HA variant, also inserted into the 51C landing site, failed to induce ectopic expression of Wg and Gbe + Su(H) in mib1 mutants, indicating that it is completely dependent on Mib1 (Figure 2M,N).

Altogether, the results confirm on the one hand a strong dependency of DI on the function of mib1. On the other hand, they indicate that DI, in contrast to Ser, can signal in the absence of the function of mib1 and raise the possibility that part of the activity of DI is independent of ubiq. Alternatively, it is possible that the residual activity of DI is induced by the activity of an unidentified DI specific E3-ligase.

**DI with a K-free ICD can signal in the absence of the function of mib1**

To discriminate between the possibilities, we generated a variant of DI where all 12 Ks of its ICD are replaced by the structurally similar arginine (R) (DIK2R-HA, Figure 3A) and inserted it into the genomic 51C landing site. It has been shown that a corresponding variant of DI1 is not ubiquitylated (Heuss et al., 2008). In agreement, we here found that, in contrast to DI-HA, DIK2R-HA is not ubiquitylated by Mib1 or Neur in S2 cells (Figure 3—figure supplement 1).

Expression of DIK2R-HA with ptcGal4 resulted in the interruption of Wg expression along the D/V boundary suggesting a strong negative effect on Notch signalling at the D/V boundary (Figure 3B, yellow arrow). The width of Wg interruption was much larger than that caused by DI-HA suggesting that DIK2R-HA has a significantly stronger cis-inhibitory activity compared to DI-HA (Figure 3B compare with Figure 2E,yellow arrow). In an attempt to quantify the increase in cis-inhibition, we measured the width of the gap induced in the expression domain of Wg by expression of DI-HA and DIK2R-HA (Figure 3—figure supplement 2A–C). We found that the gap induced by DI-HA was on average 3.3 (n = 10) nuclei and that induced by DIK2R-HA 16.7 nuclei (n = 9). This dramatic increase in the gap size of Wg expression confirms the strong increase in the cis-inhibitory abilities of DIK2R-HA.

However, DIK2R-HA was not completely inactive. It induced ectopic expression of Wg in posterior boundary cells in 44% of discs (n = 9; Figure 3E, arrows), suggesting that it still possesses signalling activity. Indeed, we found that DIK2R-HA consistently induced ectopic expression of the more sensitive Gbe + Su(H) reporter in two stripes throughout the ptcGal4 domain (Figure 3C, D, arrowhead).
Figure 2. DI induced activity in mib1 mutant wing imaginal discs. (A, B) Normal expression patterns of Gbe + Su(H) (A), Wg and ptc Gal4 (B). The arrow in (A) points to the S3 expression of Gbe + Su(H) and in (B) to the expression domain of ptcGal4. (C) The expression of ptcGal4 occurs in a gradient that increases from anterior until the A/P compartment boundary. It is not expressed in posterior cells and occurs perpendicular to the expression of Wg along the D/V boundary. A: anterior; P: posterior; D: dorsal; V: ventral. (D, E) Expression of DI-HA in a wildtype disc results in ectopic activation of
Wg (E, white arrows) and Gbe + Su(H)-lacZ (D, white arrows). The yellow arrow in (D, E) points to the intersection of ptcGal4 with the D/V boundary where the expression of Wg and Gbe + Su(H) is interrupted due to the cis-inhibitory effect of high Dl expression. Note that Gbe + Su(H) is ectopically induced throughout the whole ptcGal4 domain, which also runs through the notal region (arrowhead in D). In contrast the ectopic expression domain of Wg is restricted to the wing area and stops at the inner ring of Wg expression (arrowhead in E). (F) One stripe of the ectopic expression domains of Wg is located in regions of low expression of the anterior compartment. The second stripe is induced in posterior boundary cells that do not express Dl-HA in mib1 mutant wing discs.

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The following figure supplement is available for figure 2:

**Figure supplement 1.** DI-HA cannot induce activation of the Notch pathway in Psn mutant wing imaginal discs.

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DIK2R-HA was not able to induce expression of Gbe + Su(H) in Psn mutant discs, indicating that its expression depends on the activity of the Notch pathway (**Figure 3F**). Importantly, DIK2R-HA was able to induce strong expression of Gbe + Su(H) in mib1 null mutants, indicating that it can activate the Notch pathway in a mib1 independent manner (**Figure 3G–J**). Also weak ectopic expression of Wg was induced in a fraction of mib1 discs (18%, n = 17; **Figure 3I, arrow**). The activity of DIK2R-HA was independent of the presence of the Neur binding site in its ICD, as a variant where the Neur binding site (NEQNAV, see [Fontana and Posakony, 2009]; termed i1 in **Figure 3A**) is mutated to alanines (DIK2R11ala-HA) induces the expression of Gbe + Su(H) in mib1 mutants in a manner comparable to DIK2R-HA (**Figure 3K,L**, arrowheads, compare with G). This finding excludes the possibility that so far undetected weak ubiquitous expression of Neur activates DIK2R-HA in mib1 mutants.

In an earlier round of constructs UAS DIK2R-HA had been randomly inserted into the genome and a second- and third-chromosomal insertion had been selected for analysis. These constructs behaved in the same manner as the 51C insertion, but the phenotypes were more severe (**Figure 3—figure supplement 3**). Expression of each insertion by ptcGal4 resulted in a splitting of the wing primordium into two small halves (**Figure 3—figure supplement 3A–D**, yellow arrow in A, B, D). A similar split was observed if a dominant-negative variant of DI, DI^{stu}, or a Notch-RNAi construct is expressed (**Figure 3—figure supplement 3E–H**, yellow arrow in E, G). This similarity further confirms that the loss of the Ks strongly increases the cis-inhibitory abilities of DI. In contrast to Notch-RNAi and DI^{stu}, DIK2R-HA induced an ectopic stripe of Wg expression in the adjacent posterior boundary cells, indicating that it can activate Notch in trans in non-expressing neighbours (**Figure 3—figure supplement 3A–C**, arrowheads in A, B, arrow in C). The phenotype of DIK2R-HA expression could be mimicked by co-expression of N-RNAi and DI in wing imaginal discs (**Figure 3—figure supplement 3I,J**, compare with A, B) confirming the increase of cis-inhibitory abilities without a loss of trans-signalling for the K free DIK2R-HA.

We sought to suppress the strong cis-inhibitory effect of DIK2R-HA. In the case of DI-HA, this had been achieved through co-expression with Notch ([Klein et al., 1997; Doherty et al., 1996] (**Figure 3M**, yellow arrow, compare with **Figure 2E**). When co-expressed with Notch, DIK2R-HA (in 51C) produced the same phenotype. It induced a band of strong ectopic expression of Wg and Gbe + Su(H) and the endogenous expression of Wg along the D/V boundary was not interrupted (**Figure 3N–P**, yellow arrow in N). This finding shows that the ability of DIK2R-HA to activate the Notch pathway is partly obscured by its increased cis-inhibitory abilities. The co-expression of DI-HA and DIK2R-HA with Notch in mib1 mutants resulted in the induction of a broad stripe of expression of Gbe + Su(H) (**Figure 3Q–T**). However, induction of Wg expression was severely reduced or absent. Although we did not examine the mechanistic basis of this defect in depth, a possible explanation for the reduction of activity of the DI variants in mib1 mutants is the loss of the wing specific DS-loop. In line with this notion, we found that the ability of DI-HA and DIK2R-HA to induce expression of endogenous DI in mib1 mutant discs was severely reduced (**Figure 3—figure supplement**).
Figure 3. Expression of DIK2R-HA (inserted into the landing site 51C) with ptcGal4. (A) Schematic representation of DI-HA and DIK2R-HA and the localisation of the i1 and i2 binding boxes for Neur and Mib1 respectively in the ICD of DI. (B–D). Expression in wildtype discs interrupts the expression of Wg and of Gbe + Su(H) at the D/V boundary (B, C, yellow arrow). Expression Gbe + Su(H) is ectopically induced (C, D, arrowhead). (E) Ectopic Figure 3 continued on next page
expression of Wg (arrows) is observed in a fraction of the discs. (F) Expression of DIK2R-HA in Psn mutant discs does not induce Gbe + Su(H). (G–J) Expression of DIK2R-HA in mib1 mutant discs. (H–J) Magnification of the wing area of the disc shown in (G). The expression of DIK2R-HA causes the induction of two parallel stripes of Gbe + Su(H) (arrowheads in H, J) along the ptc domain. Wg expression is only weakly induced (I, arrow). (K, L) Expression of DIK2R14A-HA results in a similar induction of Gbe + Su(H) expression as DIK2R-HA (arrowheads). (M) Co-expression of DI-HA with Notch results in the induction of a broad stripe of expression of Wg in the wing area. Moreover, no gap in the expression domain of Wg along the D/V boundary is observed (yellow arrow, compare with Figure 2E). (N–P) A similar suppression of the increased cis-inhibitory effect of DIK2R-HA was observed upon co-expression with Notch (yellow arrow in N). (Q, R) Co-expression of DIK2R-HA with Notch in a mib1 mutant disc, reliably induces a broad stripe of Gbe + Su(H) expression (arrowhead). (S, T) Co-expression of Notch and DI-HA in mib1 mutants. A broad band of expression of Gbe + Su(H) is induced throughout the ptcGal4 domain (arrowhead). (U, V) DI Ser double mutant MARCM clones co-expressing DIK2R-HA and Notch. Wg is induced throughout the clone in the absence of endogenous ligands (arrows). (W, X) A mib1 mutant disc that bears DI Ser double mutant clones (loss of GFP) and expresses DIK2R-HA with ptcGal4. DIK2R-HA is able to induce expression of Gbe + Su(H) in the Ser DI double mutant cells (arrow).
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The following source data and figure supplements are available for figure 3:

**Figure supplement 1.** Ubiquitylation of DI-HA or DIK2R-HA in S2 cells by Neur and Mib1.
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**Figure supplement 2.** Quantification of the gap in the endogenous expression of Wg along the D/V boundary induced by cis-inhibition of the DI constructs.
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**Figure supplement 2—source data 1.** Quantification of cis-inhibition of DI variants.
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**Figure supplement 3.** The consequences of expression of a randomly inserted DIK2R-HA construct with ptcGal4.
DOI: https://doi.org/10.7554/eLife.27346.010

**Figure supplement 4.** Induction of DI expression by the DI variants.
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Therefore, most of the Notch activity in mib1 mutants is generated by expression of the exogenous DI constructs. It is probably too weak to consistently activate the expression of Wg.

Using the MARCM technique, we co-expressed DIK2R-HA with Notch in DI Ser double mutant cells and found that it induced expression of Wg in the mutant cells (Figure 3U,V; arrows). Moreover, we expressed DIK2R-HA in mib1 mutant wing discs and found that the activity marker Gbe + Su(H) was induced also in cells of DI Ser double mutant clones (Figure 3W,X, arrow). These results indicate that the activation of the Notch pathway by DIK2R-HA is independent of the endogenous ligands.

In summary, the results confirm that DIK2R-HA has a residual activity despite the loss of all Ks in its ICD. At least part of this activity is independent of mib1, and probably ubi-independent.

**DIK2R-HA displays enhanced stability**

In order to test whether DIK2R-HA is endocytosed, we monitored its sub-cellular distribution in comparison to the endosomal marker Rab7-YFP, which marks maturing endosomes (MEs) (Figure 4A–F; see also M and N). DI-HA was found at the apical membrane (Figure 4A, arrowhead) and in Rab7-positive MEs (Figure 4A–C, arrow), as previously described. We found that DIK2R-HA was distributed similarly to DI-HA at the apical membrane (Figure 4D, arrowhead) and in Rab7 positive MEs (Figure 4D–F, arrow) and localised with similar frequency at Rab7 positive MEs as DI-HA (Figure 4G). Thus, DIK2R-HA is endocytosed into approximately the same number of endosomes. This is in agreement with previous findings, which show that endocytosis of DI is not affected in a mib1 mutant background (Le Borgne et al., 2005).

Although the overall distribution in endosomes may not be affected by mib1 activity, we had earlier detected significant changes in endocytic rate of DI upon compromising the intracellular motifs mediating Mib1 interaction (Daskalaki et al., 2011). We therefore decided to assay possible stability differences between DI-HA and DIK2R-HA. For this purpose, we ubiquitously expressed DI-HA and DIK2R-HA for a limited time with tubGal4 tubGal80ts and evaluated their relative concentrations...
Figure 4. Endocytosis of DI-HA and DIK2R-HA. (A–C) Subcellular localisation of DI-HA and DIK2R-HA (D–F) in wing imaginal disc cells. The variants are detected at the apical plasma membrane (arrowhead in A, D) and in Rab7 positive MEs (arrows). (G) Quantification of the percentage of Rab7 positive endosomes that are also positive for the DI variants. (H) Summed fluorescence HA intensity of Rab7 positive vesicles in discs that either express UAS DI-HA or DIK2R-HA expressed with ciGal4. For further information of the method, see M and M. The intensity is significantly higher for DIK2R-HA

Figure 4 continued on next page
with Western blot analysis (eight independent blots). This analysis revealed that DlK2R-HA is expressed at an approximately 2.5 x higher level than Dl-HA (Figure 4I,J). Thus, the Ks in the ICD have a detectable negative effect on the stability of Dl (Daskalaki et al., 2011).

To pinpoint the phase in the endosomal pathway where the delay in degradation of DlK2R-HA is caused, we measured the fluorescence intensity of the HA staining of the Rab7 positive MEs shown in Figure 4A–F (see M and M for details). We found that the summed fluorescence intensity in the MEs was significantly higher in discs that expressed DlK2R-HA compared to discs that expressed Dl-HA (Figure 4H). This suggests that DlK2R-HA accumulates in MEs and is less efficiently transported to the lysosome than Dl-HA.

Next, we expressed Dl-HA and DlK2R-HA in a pulse-chase experiment using a combination of ciGal4 and tubGal80<sup>ts</sup>, and measured the time required for their degradation. After 16 hr of expression all cells of the anterior compartment reliably expressed either Dl-HA or DlK2R-HA (Figure 4—figure supplement 1A–E). After a chase of 24 hr, Dl-HA was completely degraded (Figure 4—figure supplement 1C,D). At this time point, we still observed high DlK2R-HA levels, also at the plasma membrane (Figure 4—figure supplement 1G,H, arrows in H). Low levels of DlK2R-HA were observed even after 36 hr of chase (Figure 4—figure supplement 1I,J). These results further confirm that the degradation of DlK2R-HA is delayed and indicate that at least two effects cause this delay: (1) Inefficient endocytosis and (2) Less efficient transport to the lysosome after endocytosis. Altogether, the results are consistent with the documented requirement of ubi at several steps in the endosomal pathway.

**A Dl variant that cannot bind to Mib1 and Neur possesses residual activity and displays increased cis-inhibition**

Dli1/2 is a randomly inserted variant that lacks the binding sites for Neur (ICD1) and Mib1 (ICD2) (Figure 3A). As a consequence, it is not ubiquitylated by these E3-ligases (Daskalaki et al., 2011). Based on the expression of Wg, the expression of Dli1/2 with ptcGal4 caused a phenotype that resembled that of DlK2R-HA, including the increased cis-inhibition judged from the large gap inflicted on the endogenous Wg stripe (Figure 5A, compare with Figure 3B). The quantification of the gap in Wg expression confirmed the increase in cis-inhibition (Figure 3—figure supplement 2).

This resemblance combined with our results raised the possibility that also Dli1/2 has residual activity, although a direct comparison of the two variants is not possible due to the unknown genomic location of Dli1/2. We monitored the induction of the more sensitive Gbe + Su(H) upon expression of Dli1/2. Indeed, we observed induction of ectopic expression of Gbe + Su(H) in wildtype, as well as in mi<sup>b1</sup> mutant wing discs (Figure 5B–F, arrowhead in B, arrow and arrowhead in E). Moreover, its co-expression with Notch induced a broad stripe of ectopic expression of Wg and Gbe + Su(H) (Figure 5G–I, arrow and arrowhead in G, H). Likewise, co-expression of Dli1/2 with Notch in mi<sup>b1</sup> mutant discs strongly induces expression of Gbe + Su(H) (Figure 5K,L), but not of Wg (Figure 5J,L). The activation of Wg expression achieved through co-expression of Notch and Dli1/2 in the wildtype is independent of the endogenous ligands, since it is also observed in Dl Ser double mutant MARCM clones (Figure 5M,N, arrow). The observed phenotypes are similar to those of expression of DlK2R-
**Figure 5.** Expression of Dli1/2 with ptcGal4. (A–C) Expression in wildtype discs causes an interruption of the expression of Wg along the D/V boundary at the point of intersection with the ptc domain (A, arrow) and induces ectopic expression of Gbe + Su(H) (B, arrowhead). (D–F) Expression of Dli1/2 in mib1 mutant discs fails to induce Wg expression (D), but induces strong expression of Gbe + Su(H) throughout the ptcGal4 domain (E, arrow and arrowhead). (G–I) Co-expression of Dli1/2 with Notch in the wildtype results in the induction of a broad stripe of ectopic expression of Wg (G, arrow)

Figure 5 continued on next page
and of Gbe + Su(H) (H, arrow and arrowhead) in a similar manner as co-expression of Notch with DlK2R-HA (Compare with Figure 3N-P, Figure 2J-L). (J-L) Co-expression of Dli1/2 with Notch in mib1 mutant results in the expansion of the ring-like expression domains of Wg (J, arrow) and ectopic expression of Gbe + Su(H) throughout the ptcGal4 domain (K, arrow and arrowhead). (M, N) Dl Ser double mutant MARCM clones (arrow, green in N) that co-express Dli1/2 with Notch. The mutant cells express Wg.

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Relief of cis-inhibition initiates ligand-dependent signalling in mib1 mutant wing imaginal discs

The results so far suggest that the loss of Notch activity in mib1 mutants may result from excessive cis-inhibition caused by endogenous levels of the DSL ligands. Cis-inhibition is determined by the ratio between the concentrations of DSL ligands versus Notch (Klein et al., 1997), implying a mechanism whereby DSL ligands and Notch expressed in the same cell engage in a non-productive stoichiometric complex (see Figure 6A). To determine the extent to which Notch signalling could be augmented in mib1 discs by relieving cis-inhibition, we made several manipulations that imbalance the stoichiometry between Notch and its ligands.

We induced Dl Ser double mutant clones in wildtype and mib1 mutant wing discs. It has been shown that the loss of function of Dl and Ser in cells of the wing pouch and the eye results in the activation of the Notch pathway by relieving cis-inhibition, when the mutant cells are located adjacent to ligand expressing wildtype cells (Michelli and Blair, 1999; Miller et al., 2009) (Figure 6A). Dl and Ser are widely expressed in the notum and in combination their patterns cover most of the notal area (Troost et al., 2015) (Figure 6—figure supplement 1A–D). Analysis of Dl Ser mutant clones revealed that cis-inhibition occurs also throughout a large part of the notum of normal wing discs: we observed activation of the Notch pathway in mutant boundary cells, indicated by the ectopic activation of Gbe + Su(H) (Figure 6B–D, arrowhead in C, D). Thus, cis-inhibition is not restricted to wing and eye development, but appears to be a frequently used mechanism to regulate Notch activity.

We next tested whether relief of cis-inhibition can also be observed in mib1 mutant discs. Indeed, strong activation of Gbe + Su(H) in mutant cells at the boundary of Dl Ser clones was observed in the notum, indicating that the loss of the ligands can result in strong activation of the Notch pathway also in the absence of mib1 function (Figure 6E–J, arrows). As expected, activation occurred only in areas where the ligands were expressed (Figure 6G–I). Note, that this Notch activity in Dl Ser mutant cells is induced by the activity of ligands in adjacent ligand-expressing cells, although these are devoid of the function of mib1 function. Since Ser is not able to activate the Notch pathway in the absence of mib1 function, even when over-expressed (see Figure 2M,N), the signalling must be mediated by Dl. Thus, Dl can signal in the absence of mib1 function without being over-expressed and ubiquitylated. The results also show that at least part of the loss of Notch activity in mib1 mutants is caused by cis-inhibition through the globally expressed ligands and supports the conclusion that Mib1 is required to overcome cis-inhibition during normal signalling.

The experiments so far were performed with the artificial reporter construct Gbe + Su(H). To confirm the results also with the promoter of an endogenous target gene, we monitored the expression of E(spl)mß-lacZ, which is expressed in a similar global manner as Gbe + Su(H) (Cooper et al., 2000) (Figure 6—figure supplement 1E–G). E(spl)mß behaved like Gbe + Su(H), as its expression was initiated in mutant boundary cells of Dl Ser double mutant clones in mib1 mutant discs (Figure 6—figure supplement 1I–L, arrowhead).
Figure 6. Analysis of cis-inhibition in mib1 null mutant wing discs in the early third instar stage. (A) Relief of cis-inhibition. In several areas of the notum of the wing disc cis-inhibition prevents significant Notch signalling (a). (b) Cells that are located at both sides of the boundary of a clone double mutant for Ser and DI. Mutant cells have lost their ligands and therefore the Notch receptor is no longer engaged in cis-inhibitory interactions and can be activated by the ligands of the adjacent DI and Ser positive cells on the other side of the boundary. Consequently the Notch pathway is activated in the...
mutant cells, which consequently activate the expression of the Notch target genes. (B–D) Cis-inhibition occurs in the notum. Dl Ser double mutant clones in the notum of a wildtype wing imaginal disc. (C, D) Magnification of the area with a clone highlighted by the arrow in (B). The clone is labelled by the absence of GFP. Expression of Gbe + Su(H) is induced in the mutant cells at the clone boundary because of relief of cis-inhibition (arrowhead). (E–J) Expression of Gbe + Su(H) is also induced in Dl Ser double mutant cells at the boundary of Dl Ser double mutant clones in mib1 mutant cells in the notal area (arrows). Thus, ligand dependent Notch activation caused by relief of cis-inhibition also occurs in the absence of mib1 function. (K, L) A large Dl Ser double mutant clone that covers most of the notal area of a mib1 mutant disc. The S3 expression of Gbe + Su(H) normally present in mib1 mutant discs is lost. The arrows point to the expected region of S3 expression (compare with Figure 1M, arrow). (M–O) Depletion of Notch function in the anterior compartment of a wing disc by expression of Notch-RNAi with ciGal4. It causes the induction of Gbe + Su(H) expression in the posterior cells adjacent to the ciGal4 expression domain (O, arrows). The arrowhead in (O) points to the unrelated expression of Gbe + Su(H) in the trachea, which is attached to the imaginal disc, but is no part of it. (P, Q) Loss of Neur function does not affect the S3 expression of Gbe + Su(H) in mib1 mutant discs. neur mutant clones are labelled by the absence of GFP. The arrow points to S3 expression in a clone.

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The following figure supplement is available for figure 6.

Figure supplement 1. Activation of E(spl)mß expression in mib1 discs through relief of cis-inhibition.

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Cis-inhibition is mutual: the non-productive cis DSL-Notch complex inhibits the activity of both receptor and ligand (Becam et al., 2010). We therefore next tested whether removing Notch expression in a group of cells would release the ligands from cis-inhibition and would result in trans-activation of the Notch pathway in adjacent cells. We created this situation by expression of a Notch-RNAi construct that efficiently depletes Notch expression if expressed with ciGal4 in the anterior compartment (Figure 6M–O). In this situation, anterior cells that lack Notch were exposed to posterior Notch expressing cells at the A/P boundary. Consequently, the endogenous ligands of the anterior cells released from cis-inhibition should signal to posterior boundary cells. Indeed, this was observed in mib1 mutant wing imaginal discs: The expression of Gbe + Su(H) was induced in adjacent posterior boundary cells (Figure 6O, arrows). This result confirms that endogenous Dl can signal in the absence of mib1 function. It also indicates that DI, not engaged in cis-interaction, can to some extent out-compete cis-interacting ligands in adjacent cells for interaction with Notch.

An instance of DI/Notch signalling that is independent of Mib1 and Neur

In mib1 mutant wing discs residual Gbe + Su(H) expression (S3) is observed (Figure 1L, arrow). S3 expression is lost in mib1 discs if the notum lacks expression of the ligands in this area, indicating that S3 is induced by DI-dependent Notch signalling (Figure 6K, L, arrow). However, expression of S3 in mib1 mutant discs is independent of Neur, as it is still detected in neur mutant clones (Figure 6P, Q). We observed a S3-like residual expression domain of E(spl)mß-lacZ in mib1 mutants (Figure 6—figure supplement 1H, J, arrow). The depletion of Notch with Notch-RNAi also abolished the expression of S3-like, indicating that it is dependent on Notch (Figure 6—figure supplement 1M–O). Hence, we have identified a region of Notch signalling in the late larval notum that is induced by DI independently of Mib1 and Neur and probably of ubi altogether.

Neur can activate DI independently of ubi

DSL signalling during wing development depends on Mib1. The alternative E3-ligase for DI, Neur, is only detected in sensory organ precursors, a small late arising minority of cells in the wing pouch. Hence, the experiments presented so far investigated the connection of Mib1 with the Ks in the ICD of DI. In order to test the connection with Neur, we co-expressed it together with the DI variants in a mib1 null mutant background. Expression of Neur alone had no effect on the expression of Wg in normal discs (Figure 7A). Co-expression of Neur with DI-HA in wildtype wing discs induces strong ectopic expression of Wg similar to that already seen with DI-HA alone (Figure 7B, arrows; compare with Figure 2E). However, we consistently observe a loss of the anterior stripe of Wg expression in the dorsal half of the pouch (Figure 7B, arrowhead). As previously reported, expression of Neur in mib1 mutants re-established the expression of Wg along the D/V boundary in the area of the ptcGal4 domain (Le Borgne et al., 2005; Wang and Struhl, 2005) (Figure 7C, arrow). The co-expression of Neur and DI-HA in mib1 mutant wing discs resulted in induction of an ectopic
Figure 7. Activation of DI-HA and DIK2R-HA by Neur. With exception of (H, I) ptcGal4 was used for expression of the constructs. (A) Expression of Neur has no effect on expression of Wg along the D/V boundary. (B) Co-expression of Neur and DI-HA causes a strong ectopic expression of Wg (arrows). The arrowhead points to the dorsal half where the anterior stripe of ectopic Wg expression is suppressed. (C) Expression of Neur in mib1 mutants restores expression of Wg along the D/V boundary (arrow). (D) Co-expression of DI-HA with Neur in mib1 mutants induces ectopic Wg expression in a
expression domain of Wg that was comparable to that observed in wildtype discs (Figure 7D, arrows; compare with B).

To our surprise, co-expression with Neur dramatically enhanced also the weak activity of DIK2R-HA in normal and also mib1 mutant wing discs. Strong ectopic expression of Wg was observed in both genetic backgrounds (Figure 7E,F, arrows). We confirmed the enhancement of DIK2R-HA by Neur with an independently generated second chromosome UAS neur insertion line (Lai and Rubin, 2001) (Figure 7G, arrows). Note, that the strength of Wg expression achieved by co-expression of Neur with DIK2R-HA or DI-HA in mib1 mutants was comparable (compare Figure 7D with F). This suggests that the Ks in the ICD of DI are of minor importance for its activation by Neur. Moreover, cis-inhibition of DIK2R-HA is reduced to a level comparable to DI-HA (Figure 7E, compare with B). Using MARCM, we found that the activation of the Notch pathway by the co-expression of Neur and DIK2R-HA occurred in the absence of endogenous DI and Ser (Figure 7H,I, arrow). This excludes the possibility that the endogenous ligands are involved in the induction of the ectopic Notch activity. We co-expressed Mib1 together with DIK2R-HA and found that in contrast to Neur, co-expression of Mib1 failed to modulate the activity of DIK2R-HA (Figure 7J, yellow arrow, compare with E). Thus, the enhancing effect on DIK2R-HA is a unique property of Neur.

NeurΔNHR1 is a variant that is unable to bind to the ICD of DI, because it lacks the necessary NHR1 binding domain (Commissio and Boulianne, 2007). In contrast to Neur, NeurΔNHR1 failed to rescue the expression of Wg along the D/V boundary in mib1 mutant discs and did not enhance the signalling activity of DIK2R-HA and DI-HA in wildtype and mutant discs (Figure 7K-N). Moreover, co-expression of Neur did not change the phenotype of Dl1/2, which lacks the Neur binding site in its ICD (Figure 7O, compare with Figure 5A). Hence, Neur must directly bind to the ICD of DI-HA and DIK2R-HA to promote their signalling activity and reduce cis-inhibition of DIK2R-HA. The results reveal fundamental differences between Neur and Mib1 in the activation of DI.

We next asked whether Neur requires its E3-ligase activity for activation of DI-HA and DIK2R-HA. To do so, we used a variant that lacks the RF, which catalyses the ubi reaction (NeurΔRF) (Pavlopoulos et al., 2001). The subcellular localisation of this variant resembles that of Neur, indicating that the RF is not necessary for localisation of Neur (Yeh et al., 2001). We have confirmed the correct localisation for the constructs used in our experiments (Figure 7—figure supplement 1). Similar to Neur, expression of NeurΔRF alone did not affect expression of Wg along the D/V boundary and did not significantly affect the activity of DI-HA in wildtype wing discs (Figure 7P,Q). However, unlike Neur, it does not abolish the dorsal anterior stripe of ectopic Wg expression observed if DI-HA is expressed alone (Figure 7O, arrowhead, compare with B). It was also unable to rescue Wg expression in mib1 mutants (Figure 7R). Nevertheless, NeurΔRF strongly enhanced the activity of DI-
HA in mib1 mutant discs (Figure 7S). In comparison to Neur, a significant difference in the strength of the ectopic expression of Wg was observed between wildtype and mib1 mutant discs (compare Figure 7Q, S and B–D), indicating a requirement of the RF for full activity. However, the result indicates that Neur can activate Dl in a manner independently of catalysing ubi of its ICD or other components of the endocytic machinery. NeurΔRF was also able to enhance the activity of DlK2R in wildtype discs, indicated by the induction of a stripe of ectopic expression of Wg in posterior boundary cells (Figure 7T, arrows). A weak stripe was induced if both proteins were expressed even in mib1 mutant discs (Figure 7U–W, arrows). Altogether, the results show that Neur, in contrast to Mib1, can activate Dl in an ubi-independent manner. However, comparison revealed that the induction of Wg expression upon co-expression of Dl and NeurΔRF was stronger than that of DlK2R and NeurΔRF (compare Figure 7S with V). In case of Neur the induction of Notch activity was comparable. This suggests that the Ks in the ICD of Dl are important for full activity, especially when Neur lacks its ability to ubiquitylate. The reason for this paradoxical requirement is not clear at the moment. A possible explanation could be a contribution of the Ks to the conformation of the ICD in a manner that cannot be completely replaced by the introduced Rs.

**Ubi-independent Dl signalling is sufficient for specification of neural fates**

Since Neur can stimulate Dl activity even in the absence of Ks in its ICD, we tested the activity of the various Dl variants in contexts, where Notch signalling is more dependent on endogenous Neur. We had shown earlier that two such developmental processes are (a) the asymmetric cell division of ganglion mother cells (GMCs) and (b) the selection of the sensory organ precursor (SOP) of macrochaetae in the larval notum.

During GMC asymmetric division the daughter cells rely on Dl/Notch signalling to turn on Hey expression in one of the two siblings (Monastirioti et al., 2010) (Figure 8A–B). neur loss of function severely compromises this signalling, whereas Mib1 only plays an accessory role (Monastirioti et al., 2010): 80% of neur-clones lose Hey expression (Figure 8C, C'), which rises to 100% in a mib1 background. However, mib1 in the presence of Neur shows no loss of Hey. Using the MARCM system, we substituted endogenous Dl by DlK2R-HA or Dli1/2 in the developing late larval CNS. The clones were scored in the central brain and ventral nerve cord. As a control, we confirmed that removal of both endogenous ligands results in 96% of the lineages (clones) being Hey negative (Figure 8D, D') and this is fully rescued (0% Hey negative) by UAS DI-HA, but not Dli1/2 (Figure 8E–F). When we expressed DlK2R-HA in Dl Ser MARCM clones, we obtained a full rescue of Hey expression; only 4% of lineages were Hey negative (Figure 8G, G'). Thus, unlike the wing pouch, where loss of Ks severely compromises Dl activity, in this context DlK2R-HA was able to sustain Notch signalling. Since Neur is needed but Ks are dispensable, we wondered whether the catalytic activity of Neur is necessary. We therefore generated neur mutant MARCM clones expressing NeurΔRF. These showed complete rescue of Hey (100% of clones are positive; Figure 8H, H'). Taken together with the DlK2R-HA result, this supports a model where ubi is not required for Dl signalling in the context of GMC asymmetric cell division.

Another Neur-dependent Notch signalling context is the selection of the sensory organ precursor (SOP) of the bristle sensillum in the notum. This occurs within proneural clusters, which are defined by the expression of proneural genes by Dl induced Notch signalling (Modolell and Campuzano, 1998) (Figure 9A). We recently introduced a MARCM based test system to test variants of Dl in this selection process (Pitsouli and Delidakis, 2005). MARCM clones double mutant for Dl and Ser cause the development of clusters of SOPs instead of single ones, termed neurogenic phenotype (Figure 9A). This phenotype can be revealed by anti Hindsight (Hnt) staining, which labels mature SOPs (Figure 9B, C, arrows). The neurogenic phenotype is completely suppressed by expression of DI-HA (Pitsouli and Delidakis, 2005) (Figure 9D, E, arrows). Similarly, DlK2R-HA strongly suppressed the Dl Ser mutant neurogenic phenotype (Figure 9F, G, arrows). This result suggests that the ubi-independent mechanism is sufficient for selection of most SOPs. To test whether Dik2R1α-Ser-HA that lacks the Neur binding site. Dik2R1α-Ser-HA failed to suppress the neurogenic phenotype in Dl Ser double mutant MARCM clones, indicating that direct binding of Neur is required for the selection of the SOP by Dik2R-HA (Figure 9H, I, arrows).
Figure 8. (A) Graphic of the larval CNS and a neuroblast lineage. Arrows depict events of Notch signalling. (B) A brain hemisphere (area boxed in A) with clonal lineages in green. One marked lineage is highlighted by the arrow. The large cell nearest to the arrow is the neuroblast. Green: GFP, Red: Hey, Blue: Delta. (C–H) Enlarged areas of brain lobes containing marked clones of various genotypes, highlighted by arrows. In all panels GFP (green) marks...
Lqf binds ubiquitylated cargo by two UIMs. In order to further evaluate the role of ubi of the DSL ligands during SOP selection, we used a “reading “defective variant of Lqf in which UIM1 was mutated and UIM2 deleted (Xie et al., 2012). This GFP-tagged variant, termed LqfUIM1<sup>3E/3A</sup>Δ UIM2-GFP is controlled by the endogenous lqf promoter and cannot recognise ubiquitylated cargo (Xie et al., 2012). As previously shown, LqfUIM1<sup>3E/3A</sup>Δ UIM2-GFP partially rescues lqf null mutant flies, which normally die during embryogenesis, to the pharate adult stage (Xie et al., 2012). The pharate adults displayed severe patterning defects that were similar to those described for mib1 mutants. The phenotype will be described in detail elsewhere. Importantly, the rescued flies displayed a nearly normal bristle pattern with only the occasional duplicated large bristle and a higher density of small bristles (Figure 9J–L, arrowhead in K, L). A similar bristle phenotype was described for mib1 mutants (Le Borgne et al., 2005). The analysis of the wing imaginal discs of the LqfUIM1<sup>3E/3A</sup>Δ UIM2-GFP rescued lqf flies revealed a nearly normal pattern of SOPs, just like in mib1 mutants (Figure 9M,O, compare with A). Altogether, these data confirm that ubi-independent, but Neur dependent DSL signalling prevails during SOP selection. Note, that complete loss of lqf function caused a neurogenic phenotype in mib1 mutant discs (Figure 9P, arrow), indicating that Lqf (but not its UIMs) is required for the ubi-independent selection of the SOP.

**Discussion**

Previous work established that the activity of the members of the DSL family depends on the function of the E3-ligases Neur and Mib1 that sends them into Lqf/Epsin dependent endocytosis. This activating endocytosis is different from bulk endocytosis, which is - in the case of DI - not dependent on Mib1 or Lqf (Le Borgne et al., 2005; Wang and Struhl, 2005). In order to account for these dependencies on E3-ligases, it has been suggested that ubi of the DSL protein ICDs by Neur and Mib1 initiates a special endocytosis event (Weinmaster and Fischer, 2011). This event either creates a pulling force that is essential for Notch S2 cleavage and subsequent ecto-domain shedding, or sends the ligands through a recycling pathway where they mature into the active form. However, it is not clear why in a given process only one of the two E3-ligases is required or why Neur strongly affects the endocytosis of DI, but not of Ser, whereas Mib1 affects the endocytosis of the ligands in the opposite way (Le Borgne et al., 2005; Wang and Struhl, 2005; Le Borgne and Schweisguth, 2003).

Here, we further assessed the role of Ks and ubi during activation of DI, using different tissues of *Drosophila* as test systems. One main finding of our work is that the activity of DI has three components: One is dependent on Mib1/Neur/E3-ligase activity and probably ubi, the second is Neur-dependent, but ubi-independent and finally the third is independent of any of these factors. Our analysis suggests that the ubi-independent activities probably account for the residual Notch activity observed in mib1 null mutants. We identified two normal developmental processes, GMC division and SOP selection, which rely on the ubi-independent/Neur-dependent activity of DI. The only example of a ubi-independent/Neur-independent case of DI signalling is the S3 notum region expression of Gbe + Su(H) and the endogenous target gene E(spl)mß. Unfortunately, it is not known in which developmental process this expression domain is involved. The latter ubi-/Neur-independent pathway appears to be restricted to DI as we did not observe ectopic activation of Gbe + Su(H) upon Ser expression in mib1 mutants.

Recent evolutionary studies suggest that the ubi-independent signalling mode of DI might be the evolutionary ancient one. *Trichoplax adherens* is the only known member of the basal metazoan phylum Placozoa. It possesses all crucial elements of the Notch pathway members except Mib1 and...
Figure 9. Specification of SOPs in an ubi-independent manner. SOPs are revealed by anti Hnt staining. (A) Schematic drawing of the distribution of the SOPs of the large bristles in a wildtype disc (red cells). In the wildtype the nascent SOP sends an inhibitory signal via the Notch pathway to prevent SOP development in its neighbours. The loss of the activity of the Notch pathway, e.g. in clones mutant for genes involved in Notch signalling, results in the formation of clusters of SOPs (neurogenic phenotype). (B–I) Di Ser double mutant MARCM clones. The SOPs are revealed by Hnt expression. Clones Figure 9 continued on next page
are labelled by GFP (green). (B, C) A DI Ser double mutant MARCM clone. The clone includes a proneural cluster (arrow). Many cells have adopted the SOP fate in the absence of DI and Ser function (neurogenic phenotype). (D, E) The neurogenic phenotype is suppressed and a single SOP is observed in a DI Ser double mutant MARCM clone that expresses DI-HA (arrows), indicating that selection of the SOP is restored. (F, G) A similar rescue of the selection process is observed when DIK2R-HA is expressed in DI Ser double mutant MARCM clones (arrows). (H, I) In contrast no rescue is observed when DIK2R<sup>1<sub>H</sub></sup> is expressed in the clones (arrows). (J) Notum of a wildtype fly showing the regular bristle pattern. (K, L) Notum of a lqf mutant fly rescued by Lqf UIM1<sup>3E/3A</sup>−ΔUIM2-GFP. (L) A magnification of the notal area of the fly shown in (K). An occasional duplication of a large bristle is highlighted by the arrowhead. The density of the smaller bristles is increased compared to the wildtype. However, most bristles are well separated from each other. (M, O) Comparative analysis of SOP formation in wing discs of lqf mutant flies rescued by Lqf UIM1<sup>3E/3A</sup>−ΔUIM2-GFP and mib1 mutant discs. The arrow highlights a SOP at the posterior dorso-central position. A nearly wildtype pattern of SOPs is observed in both genotypes. The arrowhead in (M) points to two adjacent developing SOPs that probably give rise to the bristle duplication observed in the imago (compare with K, L, arrowhead). However, no neurogenic phenotype is observed and the pattern of SOPs of both genotypes is similar. (P) lqf mutant clones in the notum of a mib1 mutant wing disc revealed by absence of GFP (green). The arrow highlights the region where the clone includes a proneural cluster. The loss of the function of lqf causes a neurogenic phenotype.

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Neur (Gazave, 2009). The E3-ligases appear to be later additions. Moreover, the only ligand present in the Trichoplax genome belongs to the DI family. In the light of our results, it is possible that Notch signalling in Trichoplax occurs solely by the probably more ancient ubi-independent mechanism. The Ser-like ligands were probably only introduced later in evolution, after the recruitment of the E3-ligases.

Besides the discovery of the ubi-independent modes, we here confirm that the Ks in the ICD of DI and Ser are important for their signalling abilities. In addition, our results indicate that the loss of the Ks in DI also results in a defect in ligand degradation and delayed endocytosis, suggesting the Ks in the ICD are required for the correct degradation. This is in good agreement with our previous finding that the endocytosis of DI1/2 that lacks the Mib1 and Neur binding sites is delayed (Daskalaki et al., 2011). We did not find significant differences in the number of Rab7 positive endosomes that are also positive for DI-HA or DIK2R-HA, however the total amount of DIK2R-HA on these endosomes was increased compared to DI-HA. Together with the persistence of DIK2R-HA in the plasma membrane, we favour a model where ICD ubi enhances trafficking of DI-HA at many stations along the endocytic route and ultimately increases its degradation.

A novel finding is that Neur and Mib1 activate DI by different mechanisms. In contrast to Mib1, Neur can activate DIK2R-HA to a level that is very similar to that of DI-HA upon co-expression, also in the absence of mib1 function. This activation requires direct binding of Neur to the ICD of DI. This indicates that Neur can strongly activate DI in a manner independently of ubi of its ICD. The findings are in agreement with the finding that Neur is involved in processes, which appear to depend on ubi-independent DI signalling, e. g. the selection of neural fates in the brain and the PNS. Previous experiments suggested that Mib1 cannot rescue the embryonic defects of neur mutants (Le Borgne et al., 2005). It is likely that Mib1’s lack of this second, ubi-independent function is the reason for this failure. One possibility for the activation of DI by Neur might be that it acts as an adapter that connects DI to the endocytic machinery in the identified developmental processes. In this way DI could be incorporated into endocytic vesicles without ubi. This adaptor function of Neur is consistent with the observation that it frequently co-traffics with DI and that expression of DI can dramatically change the subcellular distribution of Neur (Daskalaki et al., 2011; Skwarek et al., 2007). Recent work indicates that this adapter function might be more widespread among E3-ligases than anticipated since it has been shown that also the E3-ligase Suppressor of deltex, which plays an important role in endosomal trafficking of Notch, induces the endocytosis of Notch in an ubi-independent manner (Yamada et al., 2011).

The selection of the SOP occurs within groups of cells, termed proneural clusters, which are defined by the expression of proneural genes (Modolell and Campuzano, 1998). It has previously been shown that this selection occurs within a subgroup of the proneural cluster (Troost et al., 2015). It is termed the Neur group, since the cell that first expresses Neur will develop into the SOP and inhibits its neighbours by increased DI signalling. The requirement for Neur to increase DI signalling is puzzling since the cells of the subgroup also express Mib1. The discovery of the ubi-independent activation of DI by Neur provides an explanation why Neur might be required for correct SOP
selection, since it could boost DI signalling in a different manner than Mib1. Interestingly, our results indicate that also Lqf contributes to SOP selection by an ubi-independent function, as the process is to a large extent rescued by LqfUIM1\textsuperscript{3E/3A}-\textsuperscript{Δ}UIM2-GFP in lqf mutants. Hence, Lqf might be part of the Neur adapter complex.

Another so far not well-appreciated finding is that the ICD contributes to cis-inhibition. It has previously been shown that the ECD of the ligands is involved in cis-inhibition (Fleming et al., 2013; Glittenberg et al., 2006; Li and Baker, 2004). Here, we found that the cis-inhibitory ability of DI strongly increases if the Ks in its ICD are lost. We also showed that loss of mib1 results in a global increase of cis-inhibition by endogenously expressed DSL ligands. Thus, it appears that both parts of DI are required for cis-inhibition: The ECD appears to execute the necessary physical interactions with Notch, while the ICD adjusts the strength of cis-inhibition via recruiting E3-ligases and promoting ubi, which suppresses cis-inhibition.

How might DI/Notch cis-inhibition be suppressed by ubi of the ligand ICD? The ubiquitin moiety alone might already promote the separation of the DI/Notch cis-pair on the cell surface. Alternatively, ubi enables binding of Epsin/Lqf via the UIMs, which separates the pair. The similarity between the phenotype of mib1 mutants and the LqfUIM1\textsuperscript{3E/3A}-\textsuperscript{Δ}UIM2-GFP rescued lqf mutants favours the second possibility. Another possibility is that ubi/Lqf sends DI into the recycling pathway as previously suggested (Wang and Struhl, 2004). Separation of the DI/Notch ECDs is a likely event during this recycling thanks to the acidic pH in the endosome lumen. It is possible that the majority of newly synthesised DI (and probably Ser) on its way to the cell surface is initially engaged in cis-interaction with Notch. Unless their separation is promoted by ubi, only a minor amount of DI at the surface would be free to signal. This diminished signalling would justifi the reduced (but not eliminated) Notch activity, observed in mib1 mutants (e.g. S3 expression of Gbe + Su(H)).

We found that co-expression of Neur also reduces the cis-inhibitory effect of DIK2R-HA and at the same time strongly enhances its signalling activity. To do so, Neur must directly bind to the ICD of DIK2R-HA. These results raise the possibility that the binding of Neur might directly separate the DI/Notch cis-pair or, as a possible endocytic adapter, stimulates endocytic recycling upon which the cis-pair is separated. Future work is needed to clarify whether the cis-pair separation is the only mechanism through which ubi activates DSL signalling, or whether it also contributes to the pulling force which seems to be a prerequisite for Notch-activation in several contexts.

Materials and methods

Drosophila stocks and genetics

\textit{mi}b\textsuperscript{1}, \textit{mi}b\textsuperscript{1}, \textit{mi}b\textsuperscript{1/2}, \textit{mi}b\textsuperscript{1/2}EY09780 (Le Borgne et al., 2005a), lqf\textsuperscript{21} (Xie et al., 2012) and lqf\textsuperscript{227} (FRT2A (Wang and Struhl, 2004), Psn\textsuperscript{C2} FRT2A (Struhl and Greenwald, 1999b), D\textsuperscript{Fev10} Ser\textsuperscript{RX2} FRT82B (Michelli et al., 1997), H\textsuperscript{E31} (Bang and Posakony, 1992), Ser- (RRID:BDSC\_59284) and DI-MIMIC-GFP (RRID:BDSC\_59819) (Nagarkar-Jaiswal et al., 2015). \textit{Neur}\textsuperscript{1} (Commissio and Boulianne, 2007). All used alleles are null alleles.

UAS constructs: UAS \textit{neur} (Weinmaster and Fischer, 2011; Lai et al., 2001), UAS \textit{neur-myc} (third chromosome) and UAS \textit{neur\textsuperscript{Δ}RF-GFP} (Pavlopoulos et al., 2001), UAS \textit{neur\textsuperscript{Δ}NHRI-V5} (Commissio and Boulianne, 2007), UAS \textit{N-LV} (Loewer et al., 2004), UAS \textit{mi}b\textsuperscript{1} (Lai et al., 2005), UAS \textit{N-RNAi} (RRID:BDSC\_7078), UAS \textit{Di}i1 and UAS \textit{Dii1/2} (Daska\textsuperscript{λ}\textit{laki et al., 2011}), UAS D\textsuperscript{Fev1} (a gift of D. Horowicz and D. Henrique).

Further stocks: ptc\textit{Gal4} (RRID:BDSC\_2017), tub\textit{Rab7-YFP} (Marois et al., 2006), Gbe+Su(H)-lacZ (Furriols and Bray, 2001), tub\textit{Gal4} tub\textit{Gal80\textsuperscript{18}} (RRID:BDSC\_5138, RRID:BDSC\_7018), ci\textit{Gal4} (Croker et al., 2006) and vg\textit{BE-GFP} (Zecca and Struhl, 2007).

Generation of constructs

DI-HA was generated by introduction of a synthesized fragment from the NdeI restriction site of the ICD onwards. The DIK2R-HA was generated by replacing the ICD of DI-HA by a synthesised ICD in which all lysines are replaced by arginines. Constructs were cloned in pUAST attB and inserted into the landing site at 51C. Gene synthesis was performed by GenScript. DIK2R-HA was used to generate DIK2R\textsuperscript{1}ala-HA (NEQNAV to A, see [Fontana and Posakony, 2009]) and DI\textsuperscript{2}ala-HA (IKNTWWDK to A) by site directed mutagenesis. All constructs were sequenced before injection and subsequently
inserted into the attP landing site at 51C. The LqUIM13E/3A-ΔUIM2-GFP attB construct was obtained from J. Fischer (described in [Xie et al., 2012]) and inserted into the attP22A landing site. Transgenesis was partly performed by BestGene Inc. The necessary primers were purchased from Sigma-Aldrich.

**Clonal analysis**

MARCM and conventional clones were induced during the first larval instar stage (24–48 hr after egg laying).

**Western blot analysis**

Wing imaginal discs were dissected from third instar larvae and boiled in Laemmli buffer at 95°C for 10 min. Each lysate contained at least 10 discs. Proteins were separated by SDS-PAGE gel electrophoresis and blotted according to standard protocols. Antibodies used: HA (anti-HA, High Affinity, Roche (3F10) RRID:AB_390919), alpha-Tubulin (Sigma-Aldrich T5168, RRID:AB_477579), HRP-conjugated secondary antibodies were purchased from Jackson Immuno Research.

**Ubiquitylation of Dl-HA and DlK2R-HA**

**Transfections**

S2-DGRC cells (RRID: CVCL_Z992) were transfected with the calcium phosphate precipitation method. The cell line was obtained from Drosophila Genomics Resource Center, supported by NIH grant ZP400D010949. Its identity was confirmed by visual inspection of the cell morphology and its growth kinetics in M3 medium (SIGMA-ALDRICH 58398) supplemented with 10% Fetal Bovine Serum (GIBCO 10270). A mycoplasma test is usually not done for S2 cells.

For each sample 10⁷ cells were used (10 cm dish). The medium was supplemented with biotin (50 μM) for 24 hr and the lysosomal inhibitor E64 for 5–6 hr. Finally, the cells were collected 96 hr post-transfection. For each transfection, the following amounts of plasmids were used: 5 μg of pIZDeltaHA.His(6) or pIZDeltaK2R.HA.His(6)(this work), 2.5 μg pAC5-Gal4 (Addgene #24344), 2.5 μg of Actin5C-bioUbi(6)-BirA-puro or Actin5C-BirA-puro and 5 μg of pUAST-EGFP:Neuralized or pUAST-Mindbomb1:HisMyc or in the case of the negative controls pUAST-attB as a stuffer plasmid.

**Lysis and neutravidin pull-down**

The cells were washed 3 times in 1X PBS to remove excess of free biotin. The cells were resuspended in lysis buffer (0.5 ml/10 cm dish; 8 M urea, 1% SDS, 50 mM N-ethylmaleimide, 1 × protease inhibitor cocktail (Roche) in PBS, and sonicated (4 cycles 30 s ON, 30 s OFF). For the pull-down of the total biotinylated and ubiquitylated proteins, cell extracts were incubated with 50 ul of clear bed volume of high-capacity NeutrAvidin-agarose beads (Thermo Scientific, Waltham, Massachusetts, USA) overnight at room temperature. Washes were done according to Franco et al.: 2 × WB1, 3 × WB2, 1 × WB3, 3 × WB4, 1 × WB5, 1 × WB6 [WB1: 8 M urea, 0.25% SDS in PBS; WB2: 6 M guanidine hydrochloride in PBS; WB3: 6.4 M urea, 1 M NaCl, 0.2% SDS in PBS (prewarmed to 37°C); WB4: 4 M urea, 1 M NaCl, 10% isopropanol, 10% ethanol, 0.2% SDS in PBS; WB5: 8 M urea, 1% SDS in PBS; WB6: 2% SDS in PBS]. Samples were eluted in 100 ml of 4 × Laemmli sample buffer with 100 mM DTT by two cycles of heating (5 min; 99°C), with vortexing in between.

**References:** (Pirone et al., 2017; Franco et al., 2011)

**Immunostaining and microscopy**

Antibody staining was performed according to standard protocols ([Klein, 2006]. Antibodies used: anti-Wg antibody (DSHB Iowa RRID:AB_528512), anti-β-Gal (Cappel/MP Biomedicals RRID:AB_2313831), anti-Rab7 ([Tanaka and Nakamura, 2008](RRID:AB_2569808)). Fluorochrome-conjugated secondary antibodies were purchased from Invitrogen/Molecular Probes. Images were obtained with a Zeiss AxioImager Z1 Microscope equipped with a Zeiss Apotome or a Leica SP6 confocal microscope.

**Imaris image analysis for DI-HA and DIK2R-HA in Rab7 positive vesicles**

For quantitative analysis of DI-HA/DIK2R-HA in Rab7 positive vesicles, the image analysis software ‘Imaris’ (RRID:SCR_007370, Bitplane, Zurich, Switzerland) was used. Z-Stacks of imaginal discs were
acquired with a Zeiss AxioImager Z1 Microscope equipped with a Zeiss Apotome, applying the same microscopy hardware settings (e.g. exposure time) to ensure reproducibility between datasets. Individual Rab7 vesicle volumes were identified utilizing the ‘spot feature’ of Imaris with an initial size of 0.4 mm. Regrowing of volume, based on the local contrast diameter threshold was allowed later on. A quality threshold was placed on resulting spots to minimise background volumes. For further analysis only vesicles were included, which contain at least voxels with a DI-HA/DIK2R-HA signal intensity threshold of 1. Finally, the absolute cargo of DI-HA/DIK2R-HA signal in Rab7 vesicles was defined as the median of the absolute signal sum per vesicle. The average of these values over several biological replicates is displayed in the graph. Statistics were applied as described above to calculate significance levels. The procedure of vesicle volume definition, thresholding and downstream analysis was the same for all datasets. In total 19373 (DI-HA) and 15720 (DIK2R-HA) vesicles of 3 discs for each genotype were included in the analysis.

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