**Title:** The conserved C2 phospholipid-binding Domain in Delta contributes to robust Notch signalling

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**Running title:** Role of Notch-ligand C2 domain

**Key words:**  
Notch ligands, C2 domain, Delta, Drosophila, phospholipid

**Short (1-2 sentences) summary of the findings and their significance**

We used a structure-guided approach to engineer precise changes in the conserved N-terminal C2 domain of endogenous Delta and Serrate by CRISPR/Cas9 engineering. Mutations perturbing one of the protruding loops that confer phospholipid binding properties resulted in defective Notch signalling in vivo, confirming the prediction that these features contribute to normal ligand activity.

**2-3 bullet points highlighting key results**

- Structures and biochemical properties of the N-terminal C2 domains in Drosophila Delta and Serrate are like those observed in human Notch ligands.
- CRISPR/Cas9 engineering of a protruding loop in Delta C2 domain resulted in defective Notch signalling in vivo.
- Regions conferring phospholipid binding in the C2 domain contribute to Notch ligand function.
Abstract

Accurate Notch signalling is critical for organism development and homeostasis. Fine-tuning of Notch-ligand interactions have substantial impact on signalling-outputs. Recent structural studies identified a conserved N-terminal C2 domain in human Notch ligands which conferred phospholipid binding in vitro. Here we show that *Drosophila* ligands Delta and Serrate adopt the same C2 domain structure with analogous variations in the loop regions, including the so-called β1-2 loop that has been associated with phospholipid binding. Mutations in the β 1-2 loop of Delta C2 domain retain Notch binding but have impaired ability to interact with phospholipids in vitro. To investigate its role in vivo we deleted five residues within the β 1-2 loop of endogenous Delta by CRISPR/Cas9 gene editing. Strikingly, this change compromised ligand function. The modified Delta enhanced phenotypes produced by Delta loss of function alleles and suppressed that of Notch alleles. As the modified protein was present on the cell surface in normal amounts, these results argue that C2 domain phospholipid-binding is necessary for robust signalling in vivo where the balance of trans and cis ligand-receptor interactions is finely tuned.
INTRODUCTION

The Notch signalling pathway is highly conserved and plays key roles in many aspects of development and homeostasis (Bray, 2016). Aberrant Notch signalling results in a number of inherited diseases and is associated with various cancers and other acquired disorders (Mašek & Andersson, 2017; Nowell & Radtke, 2017; Siebel & Lendahl, 2017; Monticone & Miele, 2021). As both the Notch receptors and the ligands are single-pass type I transmembrane proteins, signalling is initiated by direct protein–protein contact between adjacent cells, which may occur in some instances via long cell processes such as cytonemes (Boukhatmi et al, 2020; Cohen et al, 2010; Hunter et al, 2019; De Joussineau et al, 2003; Huang & Kornberg, 2015). Canonical Notch signalling involves a simple cascade, whereby ligand binding induces successive cleavages to release the Notch intracellular domain (NICD) which translocates to the nucleus and directly regulates gene expression with its binding partners (Bray, 2016; Kovall et al, 2017; Kovall & Blacklow, 2010; Kovall, 2008). One challenge is to understand how this simple core mechanism is modulated to ensure appropriate spatio-temporal regulation of the pathway. Mechanisms that fine-tune the ligand receptor interactions are likely to make important contributions.

All Notch ligands have a similar architecture, with an extracellular domain consisting of multiple (7, 8, or 16) epidermal growth factor (EGF) repeats, a so-called Delta/Serrate/Lag-2 (DSL) domain and a highly conserved N-terminal region (Kopan & Ilagan, 2009; D'Souza et al, 2008; Bray, 2006; Kovall & Blacklow, 2010). Receptor binding involves the N-terminal portion including the DSL and N-terminal domains (Luca et al, 2015, 2017; Cordle et al, 2008). Structural studies of the N-terminal region from human Delta and Jagged ligands revealed that it adopts a conformation characteristic of a phospholipid-binding C2 domain (Chillakuri et al, 2013; Kershaw et al, 2015). In agreement, these domains interact with phospholipid-containing liposomes in vitro, and exhibit ligand-specific preferences for liposomes of different composition (Suckling et al, 2017). Comparisons between mammalian Jagged and Delta type ligands revealed a diversity in the structures of the loops at the apex of the C2 domain which are implicated in membrane recognition in other C2 domain proteins (Suckling et al, 2017). A subset of missense mutations, which affect these loops in Jagged1 are associated with extrahepatic biliary atresia (EHBA)(Kohsaka et al, 2002).
Purified EHBA variants show reduced Notch activation in reporter cell assays, lead to a reduction in phospholipid binding, but do not alter Notch binding (Suckling et al, 2017). The C2 domain may therefore have a role in tuning the activity of the Notch ligands through its lipid-binding properties.

Mutations affecting the single Delta or Serrate (Jagged-like) ligands in Drosophila have well characterized consequences on development (e.g. (Heitzler & Simpson, 1991; Thomas et al, 1991; Fleming, 1998; Bishop et al, 1999; de Celis et al, 1996, 1997). Homozygous loss of ligand function leads to lethality but several defects, including wing venation abnormalities, are detected even in Delta heterozygotes, which have one normal gene copy (Dexter, 1914; Huppert et al, 1997; de Celis et al, 1997). As these defects occur when only one allele is mutated, it is evident that patterning is highly sensitive to ligand levels and activity. This therefore provides a powerful context in which to investigate the contributions from the apical C2 domain loops to ligand activity in vivo.

As the loop regions of the C2 domains are the most variable, we first set out to solve the structure of the C2 domains from the Drosophila Delta and Serrate ligands. This revealed similar prominent β1-2 and β5-6 loops to those in the C2 domain of the mammalian ligands that are thought to be responsible for the interaction with phospholipid head groups (Suckling et al, 2017). To test the functional contribution, we focussed on the β1-2 loop in Delta and used CRISPR/Cas9 genome editing to delete 5 amino acids so that we could analyse the impact on Notch activity during development. In vitro, such DIβ1-2 mutation(s), resulted in expression of stable protein with altered phospholipid-binding properties. Strikingly, in vivo the DIβ1-2 mutation compromised ligand function, exhibiting characteristics of reduced signalling activity. Our data therefore confirm the relevance of C2 domain loops for full ligand activity and, given their ability to confer lipid-binding, suggests that membrane-binding properties are important for robust signalling.
RESULTS

Structure and binding properties of the C2 domain of Drosophila ligands

To determine whether the Drosophila ligands adopt the same arrangement as their mammalian counterparts we solved the structures of the N-terminal region of Drosophila Delta and Serrate (Fig. 1) as well as the ligand binding region of Drosophila Notch (EGF11-13; Fig. EV1A-E). These were solved using molecular replacement of the individual domains from the human homologues to resolutions between 1.5 and 3.0 Å (Table 1, Fig. 1). When the new Drosophila ligand structures were overlaid on their mammalian equivalents, Jagged-1 and DLL4, it was evident that the core domain structure and arrangements of the fly ligands are highly conserved (Fig. 1A-C and Fig. EV1D; RMSD 2.5 Å for Delta and 3.1 Å for Serrate) as was the structure and domain arrangement of the Notch receptor ligand binding region (Fig. EV1D; RMSD 1.1 Å).

The conserved domain arrangement allows us to model the Notch-ligand complex by overlay of the Drosophila structures on the earlier structures of the mammalian complexes (Luca et al., 2015, 2017) with this leading to no significant clashes between the Notch and ligand coordinates. Notable exceptions to the overall conserved arrangements are the β1-2 and β5-6 loops which exhibit different lengths and folding in the ligands. These highly variable loops protrude apically from the C2 domain core and are positioned far from the Notch-binding interface.

Given the structural conservation with the mammalian ligands, it is likely that the Drosophila proteins exhibit similar properties. Purified N-terminal fragments (NE3 variants) were therefore used to test the liposome-binding capability of variants in which 5 amino acids were deleted from the β1-2 loop, hereafter referred to as Delta^{Δβ1-2}. The β1-2 loop was selected because of its importance for phospholipid binding in other C2-domain proteins (Hirano et al., 2019; Honigmann et al., 2013; Verdaguer et al., 1999) and because the genomic organization (present in a single exon) meant that the equivalent mutation could be engineered in vivo (as described below). Using a liposome composition of Phosphatidylcholine (PC): Phosphatidylserine (PS): Phosphatidylethanolamine-fluoroscein (PE) (80:15:5), we could detect binding of wild-type Delta (Delta^{WT}) fragment to the liposomes as seen for mammalian Notch ligands (Fig. 1D). This binding was compromised when the variable β1-2 loop was shortened, resulting in the deletion of residues GATGK; Delta^{Δβ1-2} fragment exhibited a significant
reduction in binding compared to that from Delta\textsuperscript{WT}. Likewise, the equivalent fragment containing Serrate\textsuperscript{Δβ1-2} loop deletion (removal of residues LRATK) also exhibited reduced binding to liposomes, although to a variable extent that was not reproducibly significant (Fig. 1D). This may be due to differences in the lipid-binding specificities because we have previously noted the heterogeneity of the C2 loops sequences in different ligand families and hypothesised that they may confer different lipid-binding specificities (Suckling \textit{et al}, 2017).

Purified Delta (NE3 fragment) also exhibited robust binding to a fragment of \textit{Drosophila} Notch (dNotch EGF11-13), which contains the core ligand-binding sites (Fig. 1E). This relies on the conventional contact sites because it is abolished by an alanine substitution in the DSL domain which replaces a key receptor-binding residue (F204). In comparison, the variant with the loop deletion, the Delta\textsuperscript{Δβ1-2} fragment, retained Notch binding as predicted from the fact that the loop is positioned far away from the Notch binding interface (Fig. 1A-C). We attribute the small difference in Notch binding compared to the wild-type to the slightly lower purity of the protein preparation (Fig EV1F) although we cannot rule out that the mutation causes a minor modification to the Notch interaction.

Together these data demonstrate that the C2 domain structure is conserved between species and that the properties detected in the mammalian ligands are also shared by the \textit{Drosophila} counterparts. The main source of variability is present in the N-terminal apical loops which nevertheless are important for liposome binding in \textit{Drosophila} Delta as in DLL-4 and Jagged-1 from mammals.

**Phenotypes produced by mutations in the ligand β1-2 loop**

The β1-2 and β5-6 loops generally make important contributions to phospholipid binding in C2 domains (Hirano \textit{et al}, 2019; Honigmann \textit{et al}, 2013; Verdaguer \textit{et al}, 1999). As the β1-2 loop in the \textit{Drosophila} ligands is encoded by a small sequence in a single exon (exon 2 of \textit{Delta} and exon 3 of \textit{Serrate}), it was the most amenable to mutagenesis by genomic engineering. Therefore, in order to study the importance of this loop for Notch signalling, the endogenous exons were replaced by modified exons where the coding sequence of the loops was deleted by CRISPR-mediated
homologous recombination. For each of the ligands, two gRNAs were designed to flank the target exon, and the recombination of the modified exon was promoted by a complementary sequence within which the β1-2 loop was replaced by a mutated version \( (D_{b1-2}^{b1-2}; \text{Fig. 2A and EV2A}) \). Successful recombination was identified by the presence of a dsRed marker that was subsequently removed and the mutations were confirmed by sequencing of the exon. As well as generating \( D_{b1-2}^{b1-2} \) mutations, we also recovered a deletion of the entire exon 2, \( D_{Exon2}^{b1-2} \) which removes a key part of the receptor-binding region and behaves as a null allele (Fig. EV2F).

Severe loss of Delta function, as with \( D_{Exon2}^{b1-2} \), results in lethality. In contrast \( D_{b1-2}^{b1-2} \) homozygotes were viable. Nevertheless \( D_{b1-2}^{b1-2} \) adult flies exhibited several visible phenotypes. Firstly, they had ectopic wing-vein material, with extra vein tissue detected around L2, L5 and the posterior cross-vein (Fig. 2B-C, arrowheads). Secondly, they had abnormal spacing of the microchaetae on the thorax (Fig. 2D-E). Both the venation and microchaetae defects are consistent with altered Notch pathway activity (Heitzler & Simpson, 1991; Vässin & Campos-Ortega, 1987), suggesting that localised mutations affecting the β1-2 loop impair the function of the Delta ligand.

The defects produced by \( D_{b1-2}^{b1-2} \) were relatively mild, and there were no disruptions to the wing margin (e.g. notching). In agreement, expression of genes \textit{cut} and \textit{deadpan} that require high levels of Notch signaling at the d/v boundary (Micchelli \textit{et al}, 1997; Babaoğlan \textit{et al}, 2013; San Juan \textit{et al}, 2012) was not disrupted in \( D_{b1-2}^{b1-2} \) mutants (Fig. EV3A) or in patches of \( D_{b1-2}^{b1-2} \) mutant cells (Fig. EV3B-E). Likewise, Ser\textit{f1-2} had normal wings (Fig. EV2B) but exhibited mild abnormalities associated with ectopic pigmentation of joints, that could also be indicative of compromised signalling. Together the data indicate that the specific deletion within the β1-2 loop has a detectable but mild effect on Notch ligand functions.

**Ligand β1-2 loop mutants exhibit reduced activity**

To further probe the consequences from the mutations in the C2 domain β1-2 loop, \( D_{b1-2}^{b1-2} \) was combined in trans with previously characterised deletions \( (Df(3R)Dl^{Fx3}) \) and loss of function (e.g. \( Dl^{Fev10} \)) \( Dl \) alleles. When heterozygous, the strong \( Dl \) alleles exhibit a robust and consistent wing vein phenotype, with “deltas” formed by extra vein
material along several of the veins (Fig. 3A-C – left panel). In combination with $Dl^{A/b1-2}$ this phenotype was strongly enhanced, so that more of the veins were affected and they became uneven and thickened (Fig. 3A-D – right panel). The enhancement of vein defects by $Dl^{A/b1-2}$ occurred in combinations with all $Dl$ alleles tested. Likewise, $Ser^{A/b1-2}$ had a similar effect. Full Notch activity in the wing veins also requires Serrate, as revealed by chromosomes carrying mutations in both $Dl$ and $Ser$, which have more severe phenotypes than $Dl$ mutations alone despite the fact that $Ser/+ \text{flies}$ have normal veins (Fig. EV2B). Combining $Ser^{A/b1-2}$ allele with this double mutant chromosome, enhanced the thickening of veins in a similar manner to $Dl^{A/b1-2}$ (Fig EV2C-D). The enhanced vein phenotypes indicate that deletions within the $\beta1-2$ loop of the C2 domain compromise ligand activity.

One unusual feature of the Notch pathway is that the ligand and receptor molecules can interact together in cis, when they are present on the same cell surface (De Celis & Bray, 1997; Micchelli et al, 1997). This cis-interaction is inhibitory and may be important to set a threshold that ensures a sharp response (Sprinzak et al, 2010). One manifestation of this balance is that the phenotypes produced by reduced Notch function are suppressed when combined with a $Delta$ loss of function allele (Fig.3F; De Celis & Bray, 2000). Notch heterozygous females have a characteristic wing-notching phenotype (Fig. 3E). When combined with $Dl^{A/b1-2}$ the wing notching phenotype was suppressed to a similar extent as with a classic $Delta$ allele (Fig.3E-F), suggesting that cis-interactions are also modified in this context. $Dl^{A/b1-2}$ also gave a modest and variable modification of the vein phenotype from Notch heterozygotes in a similar direction (Fig. EV2E). However, we note that the loop mutation is not sufficient to fully alleviate cis-inhibition, as we did not detect ectopic target gene expression when homozygous $Dl^{A/b1-2}$ mutant clones were juxtaposed with wild-type cells (Fig. EV3D-E; (Micchelli et al, 1997)).

Changes to signalling were also detected in another Notch-dependent process, the spacing between the sensory organs, microchaetae, on the notum. In the absence of Notch signalling an excess of sensory organ precursors are formed due to failure in lateral inhibition (Cohen et al, 2010; Heitzler & Simpson, 1991; De Joussineau et al, 2003; Sjöqvist & Andersson, 2017). Milder defects in Notch signalling lead to irregular
and reduced spacing between the sensory organ precursors with the consequence that there is an increase in the number of microchaete on the adult notum as seen in flies heterozygous for a deletion of Delta (e.g. $Df(3R)Dl^{Fx3}$/+; Fig. EV4A-B,E). As noted above $Dl^{A/0}$-2 homozygous flies had an increased density of microchaetae compared to wild-type (Fig. 2D-E and EV4C,E) and in combination with strong Delta alleles, $Dl^{A/0}$-2 led to a further increase in microchaetae numbers (Fig. EV4D-E; $Df(3R)Dl^{Fx3}$/$Dl^{A/0}$-2). Thus, as with the vein formation, the defects in microchaetae spacing indicate a reduced signaling potential for ligands with a shortened $\beta$1-2 loop, despite the fact that this change should not disrupt binding to the receptor per se (see Fig. 1E).

**$Dl^{A/0}$-2 has compromised Notch signalling in photoreceptor fate decisions**

Flies homozygous for $Dl^{A/0}$-2 also had mild roughening of the eyes. Notch activity is required at several stages in the development of the photoreceptors, including in the specification of R4 and R7 photoreceptors. The sequential differentiation of the eight neuronal photoreceptors (R cells) is initiated when a wave of differentiation (called Morphogenetic Furrow or MF) spreads from the posterior to the anterior region of the eye imaginal disc ((Şahin & Çelik, 2013); Fig. 4A). Notch activity in one cell of the five-cell cluster specifies R4 cell fate and can be detected by the expression of $E(spl)m\delta0.5$-lacZ, containing the Notch responsive $E(spl)m\delta$ enhancer ((Cooper & Bray, 1999); Fig. 4A,B). Reducing the levels of Delta, as seen in Delta heterozygotes $Df(3R)Dl^{Fx3}$/+, led to more variable expression of $E(spl)m\delta0.5$ (Fig. 4B). This was further enhanced in combination with $Dl^{A/0}$-2, so that many of the ommatidia exhibited very low levels of expression (Fig. 4B-C). No similar reduction occurred with $Dl^{A/0}$-2 heterozygotes (Fig. 4B) nor clones of $Dl^{A/0}$-2 homozygous mutant cells (Fig. EV5A-B") arguing that the decrease in activity in these conditions is not below the threshold needed for $E(spl)m\delta0.5$ activation. Nevertheless, the fact that the $Dl^{A/0}$-2 enhances the phenotype from the Delta deletion is consistent with it being compromised for productive Notch signalling.

**C2 Domain $\beta$1-2 loop mutation does not impair Delta trafficking**

Our results indicate that the $\beta$1-2 loop region of Delta C2 domain is required for full functionality. To investigate whether this involves a change in the localization or trafficking of Delta we generated mutant clones in the wing disc, a tissue where the
expression and localization of the ligand is well characterised. In late third instar stages, the expression of Delta is particularly enriched in two stripes flanking the DV boundary and in longitudinal stripes that prefigure the prospective wing veins (Fig. 5A,A'). In all regions of the disc, DIβ1-2 exhibited normal expression levels and it appeared to be localized at the apical membranes, at similar levels to wild-type Delta.

To confirm that the mutant protein was present on the surface we performed an antibody uptake assay (Le Borgne & Schweisguth, 2003). Wing imaginal discs were incubated ex vivo with an anti-DI antibody recognizing the extracellular domain at 4°C. Excess antibody was then washed away, and the tissues transferred to a permissive temperature (25°C) for 0 or 30 minutes so that the membrane localization, uptake and trafficking of bound antibody could be measured (Gomez-Lamarca et al, 2015). At zero minutes when antibody was bound to Delta on the cell-surface, similar levels were detected in control regions and in DIβ1-2 mutant clones (Fig. 5B,B'), indicating that the mutant protein was present on the cell surface. When endocytosis was allowed to proceed for 30 minutes, antibody bound Delta accumulated in puncta throughout the epithelial cells in both wild-type and DIβ1-2 tissue (Fig. 5C,C'). The uptake assays confirm therefore that the mutated protein reaches the cell surface normally and that its trafficking following endocytic uptake is not grossly affected, although we cannot rule out a subtle change.

**DISCUSSION**

C2 domain phospholipid binding properties are essential for membrane-targeting of many intracellular proteins. Notch ligands are unusual in having an extracellular N-terminal C2 domain (Chillakuri et al, 2013; Kershaw et al, 2015). This structure is present in all the human Notch ligands and retains the capacity to interact with liposomes (Suckling et al, 2017). Here, we showed that Drosophila Delta and Serrate also contain a globular C2 domain that confers the ability to bind to phospholipid-containing liposomes in vitro. The C2 domain structures are highly conserved, differing only in the length and orientation of several loops. A deletion mutation affecting one of these, a loop between the β1 and β2 strands of the C2 domain core, was sufficient to compromise liposome binding. This loop might therefore help to generate a “pocket”
capable of interacting with a specific type of lipid, e.g. phospholipid/glycosphingolipid and in this way influence productive Notch signalling.

A subset of human Jagged1 mutations that affect the loops at the apex of the C2 domain are associated with extrahepatic biliary atresia suggesting these regions are important for tuning the Notch signal in physiological contexts (Kohsaka et al, 2002; Suckling et al, 2017). Our results, from CRISPR engineering β1-2 loop mutations in Drosophila Delta and Serrate, support the conserved functional importance of the C2-domain loops. The mutated Delta exhibited reduced signaling activity in several different developmental contexts. The compromised signaling was most evident in genetic combinations with a strong loss of function allele or deletion of the locus, and was manifest by enhanced vein thickening, extra sensory bristles and reduced signalling during photoreceptor fate choice although there were no overt effects at the dorsal-ventral boundary. All of the processes affected involve highly dynamic signaling and are sensitive to subtle changes in signaling as evident from the defects in animals with reduced dosage of wild-type Delta (Df(3R)DfFx3/+). These results are consistent with the model that C2 domain loop regions are important for fine-tuning the Notch signal (Fig. 6A,B), as suggested by in vitro results where the binding of fluorescent liposomes to Jagged was modified/enhanced in the presence of a Notch1 11-13 fragment suggesting a coupling between C2 domain lipid binding and Notch binding, and by the changes in Notch activation seen with EHBA and related loop variants (Suckling et al, 2017).

There are several models for how C2 domain-mediated membrane interactions might impact on signalling. One possibility is that the spatial or temporal residence of Delta in the membrane may be affected by the C2 domain interactions. Evidence suggests that relative pools of the receptor and ligands, rather than absolute concentrations, are important for refining signalling outcomes due to the balance between cis-inhibition and trans-activation (Sprinzak et al, 2010). Models based on this relationship inferred that intrinsic noise would cause the width of the vein to become irregular, one characteristic of the phenotype produced from DfAβ/2. Loss of interaction with certain types of lipids might bias how the ligand interacts with the receptor. For example it could shift in favour of inhibition, producing a generalized reduction in signaling, despite there being similar amounts of proteins on the cell surface. However the ability of DfAβ/2 to suppress the phenotype from reduced Notch at the wing margin argues
that cis-inhibition is also compromised by the loop mutation in some contexts. This makes it more likely that C2 domain interactions fine-tune both the activating and inhibitory interactions, perhaps by modulating the length of time the ligand is diffusing in the membrane (Khait et al, 2016), and that the precise consequences may differ depending on the relative amounts of ligand and receptor present.

In summary, our structure-guided approach to make defined changes in the endogenous ligands has demonstrated the *in vivo* relevance of C2 domain loops for full activity in the physiological setting (Fig. 6A,B). This approach has uncovered subtle functional requirements that would unlikely be detected using *in vitro* or *in vivo* methods alone, highlighting the importance of using interdisciplinary methods to fully elucidate function.
MATERIALS AND METHODS

Protein Expression, Crystallisation and Structure Determination

Codon optimized open reading frames for constructs (synthesized by GeneArt®, Life Technologies Ltd., Paisley, UK), with recommended BiP signal peptide (for secretion), were subcloned into expression vector pEXS2-2 (Expres2ion® Biotechnologies, Horsholm, Denmark) using EcoRI and NotI restriction sites (see Table EV1 for primer sequences). Each construct was expressed as a monomer with a C-terminal 8xHis tag to facilitate purification. Purification was as described in Suckling et al. (Suckling et al., 2017). Drosophila (d) Delta and Serrate NE3 (C2 domain-DSL, EGF1, EGF2, EGF3) constructs were used for liposome- and Notch-binding assays. NE3 (residues 1-332 Delta; 1-388 Serrate), NE2 (residues 1-293 Delta, residues 1-349 Serrate), NE1(1-259 Delta, 1-314 Serrate) constructs were used to set up preliminary crystal trials. Delta NE1 and Serrate NE2 constructs produced best diffracting crystals. NotchEGF11-13 was produced using the same expression system, and the purified cleaved form used for crystallization.

The Notch receptor construct was concentrated to 24.2mg/ml in a buffer A (10mM Tris pH 7.5, 150mM NaCl, 10mM CaCl₂) and crystallised by the sitting drop method from 200nl + 200nl drops with mother liquor 0.1M MES pH 6.5, 1.8M Ammonium sulphate, 0.01M cobalt chloride. Crystals were cryoprotected by addition of 25% ethylene glycol and data collected at the European Synchrotron Radiation Facility, beamline ID29. Serrate was concentrated to 15.8 mg/ml in buffer A and crystallised by the sitting drop method from 200nl + 200nl drops with mother liquor 0.1M imidazole malate pH 7, 25% PEG4K and cryoprotected by addition of 25% ethylene glycol, 20 mM CaCl₂. Data were collected at Diamond Light Source, beamline I02. Delta was concentrated to 17.7 mg/ml and crystallised by the sitting drop method from 200nl + 200nl drops with mother liquor 0.1M Tris pH 8.5, 0.2M MgCl₂, 30% PEG4K and cryoprotected with 25% glycerol, 20mM CaCl₂. Data were collected at Diamond Light Source, beamline I04.

All structures were solved by molecular replacement using separated domains from the human homologues using program PHASER (McCoy et al, 2007) from program suite CCP4 (The CCP4 suite: programs for protein crystallography., 1994) rebuilt using BUCCANEER (Cowtan, 2006) and COOT (Emsley et al, 2010) and refined in
PHENIX (Liebschner et al., 2019). For all constructs data processing and model statistics are described in Table 1. Coordinates and data are deposited in the Protein Data Bank with accession codes 7ALJ, 7ALT, 7ALK for Notch, Serrate and Delta respectively.

**Liposome and Notch Binding assays**

Liposome binding assays were carried out as described in Suckling, Korona et al. (Suckling et al., 2017) using purified Delta/Serrate variants and liposomes comprising Phosphatidylcholine (PC): Phosphatidylserine (PS): Phosphatidylethanolamine-fluorescein (PE) in a 80:15:5 ratio. Liposomes were prepared as described in Chillakuri et al. (Chillakuri et al., 2013). Notch binding assays were carried out as described in (Suckling et al., 2017) using purified Delta variants and Notch EGF 11-13. The negative control Delta F204A variant reduces Notch/ligand binding at Site 2, by altering a key residue within the ligand DSL domain Notch-binding loop. We note there is some slight variability in the purity of the protein preparations (see Fig. EV1F).

**Drosophila melanogaster strains and genetics**

All *Drosophila melanogaster* stocks were grown on standard medium at 25°C. Alleles are as described in Flybase (Thurmond et al., 2019) and in particular the following were used to sensitize the genetic background: *Df*[^rev^10] (Heitzler & Simpson, 1991), *Df[^rev^10,^Ser^106] (Thomas et al., 1991), *Df(3R)Df[^Fx^3] (Vässin & Campos-Ortega, 1987), *N[^55e^11]* (#BL28813). *Df[^1^1^β^1]-2* clones were generated using FRT mediated recombination (Xu & Rubin, 1993) – recombination was promoted by heat-shock of 1h at 37°C 72h prior to dissection and analysis. The *E(spl)m[^δ^0.5]* reporter was used for analysis of R3/R4 determination in eye imaginal discs, (Cooper & Bray, 1999).

**Generation of β1-2 loop Notch Ligands mutants using CRISPR/Cas9**

Lines were generated by CRISPR-mediated homology repair (HR) strategy. As described in Fig 3 and FigEV2, two guideRNAs were designed to flank the target exon coding the β1-2 loop (see Table EV1 for primer sequences) and cloned into the guide RNA expression pCFD4 vector (Addgene #49411). The exon of interest and homology arms were cloned into donor template plasmid pHD-ScarlessDsRed (Addgene # 64703) using the Gibson Assembly Protocol (see Table EV1 for primer
sequences). Modifications to the exons were made using standard mutagenesis and PCR amplification prior to the co-injection of the guide RNAs and the donor template constructs into nos-Cas9 (#BL54591) embryos. Modifications included: 1) deletion of 15bp within β1-2 loop of DI (ΔDIβ1-2); 2) deletion of 15bp within β1-2 loop from Ser Exon 3 (ΔSerβ1-2); 3) deletion of region between the two gRNAs (ΔDlExon2). Δexon 2 could in principle produce a modified protein, the deletion would be in frame, but it would lack any signal peptide. As no residual protein was detected by antibody staining (Fig EV2F) the product is either not made or is unstable. Engineered flies were identified by expression of DsRed in the eyes and verified by genomic PCR sequencing. The transposable element containing the DsRed was removed subsequently by crossing to flies carrying PiggyBac Transposase (#BL32070).

**Immunostainings**

The following primary antibodies were used for Immunofluorescence staining: Goat anti-GFP (1:200, Abcam, ab6673), Mouse anti-Cut (1:20, Developmental studies hybridomw bank (DSHB)), Rat anti-DE-Cad2 (1:200, DSHB), Mouse anti-Delta (1:30, DSHB), Guinea pig anti-Delta (1:2000, a gift from Mark Muskavitch, (Huppert et al, 1997)), guinea pig Anti-Dpn (1:2000, a gift from Christos Delidakis), Mouse anti-NECD (1:50, DSHB), Rat anti-ELAV (1:200, DSHB), Mouse anti-β-Gal (1:1000, Promega, Z378A). Uptake assay was performed as described previously (Gomez-Lamarca et al, 2015).

**Adult tissues analysis**

For the analysis of the adult fly wings, female flies were collected in 70% Ethanol for 2 hours, rehydrated in PBS and one wing per fly was isolated and mounted in a 50% glycerol solution. To analyse the microchaete number, flies were collected in 70% Ethanol for 2 hours, rehydrated in PBS 1x and mounted on apple juice agar plates for imaging.

**Imaging and statistical analysis**

Immunostaining samples were imaged with Leica TCS SP8 microscopes (CAIC, University of Cambridge) at 40X magnification and 512/512 or 1024/1024 pixel resolutions. Images of the adult wings were taken using a Zeiss Axiophot microscope.
and images of the adult Notum were taken using the Leica MZ10F coupled with a camera Leica DFC3000G. ImageJ software was used to analyse images and polygon tool was used to measure the vein area on the region limited by the CV2, L4 and L5 veins on adult wings. The measurement of the wing notching was done by determining the tissue missing with the polygon tool after superimposing wings of the described genotypes with the reference wild type wing. The number of microchaete was assayed using a fixed area as reference on the Notum, as depicted by the white box on figure EV4A. For the analysis of the DI and Notch trafficking in $D_{I/D\text{-}2}$ mutant clones, a projection of 3-cell diameter was performed after re-slicing the images into the XZY axis in ImageJ software.

Statistics were calculated with GraphPad Prism. Comparisons between two groups were performed with a two-tailed unpaired t-test. Statistical differences among various groups were assessed with ordinary one-way ANOVA by comparison to the mean of the control column. For comparisons between two samples the student's T-Test was used. In the figures and figure legends ns indicates no significant difference; * P<0.1; ** P<0.001; ***P<0.0001; Multiple comparisons, we used ANOVA. Comparison between two samples, studentt;'s T-Test

Data Availability
Coordinates and data have been deposited in the RCSB Protein Data Bank (https://www.rcsb.org/) with accession codes 7ALJ, 7ALT, 7ALK for Notch, Serrate and Delta respectively.

Acknowledgments
We thank Mark Muskavitch, Christos Delidakis, the Bloomington Stock Center, the VDRC Stock Center and the Developmental Studies Hybridoma Bank for $Drosophila$ strains and antibodies. We thank Kat Millen for the technical help and for injections for the CRISPR engineering. We thank other members of SJB lab for valuable discussion. This work was funded by grants from BBSRC (BB/P006175/1 to SJB) and MRC (MR/L001187/1 to SML and PH; MR-R009317/1 to PH, MR/T014156/1 to SJB) and by a Wellcome Trust investigator Award (100298 to SML)
Author contributions
T.M and S.J.B designed the in vivo experiments. T.M performed the in vivo experiments. T.M and S.J.B analysed the data. P.H and S.M.L designed the in vitro experiments. B.K, Y.M and R.S purified ligand and receptor proteins. R.S and S.J performed structure determination. Y.M performed the Notch-binding experiments. B.K performed liposome-binding assays. B.K, Y.M, R.S, P.H and S.M.L analysed data. T.M, P.H, S.M.L and S.J.B wrote the manuscript.

Conflict of interest: The authors declare that they have no competing interests.
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Huang H & Kornberg TB (2015) Myoblast cytonemes mediate Wg signaling from the wing imaginal disc and Delta-Notch signaling to the air sac primordium. Elife 4: 1–22


McCoy AJ, Grosse-Kunstleve RW, Adams PD, Winn MD, Storoni LC & Read RJ


FIGURE LEGENDS

Figure 1. Structure and binding properties of *Drosophila* ligands

**(A,B)** Left panels. The structures of the N-terminal regions from *Drosophila* Delta (**A**) and Serrate (**B**) are shown in a cartoon representation (rainbow coloured from blue at N-terminus to red at C-terminus). These have been overlaid on their mammalian equivalents DLL-4 (**A**) and Jagged-1 (**B**) in the context of their complexes (PDB entries 4xlw and 5uk5 respectively) with Notch-1 (cartoon representation, coloured grey). The structure of isolated *Drosophila* Notch is also depicted in each panel (cartoon, rainbow coloured) superposed on the respective copy of mammalian Notch-1 (cartoon, grey) for each complex. The overlays demonstrate the high degree of conservation in domain structures and arrangements between the *Drosophila* and mammalian homologues. Right panels. A close-up view of the C2 domains of each ligand overlaid with their mammalian equivalent. These demonstrate conservation of overall fold but large differences in the apical loops, particularly in the β1-2 and β5-6 loops.

**(C)** Isolated structure of N-terminal Delta, with residues deleted in ∆β1-2 highlighted as red Van Der Waals spheres.

**(D,E)** Purified *Drosophila* Delta and Serrate NE3 proteins bind to liposomes (**D**) Binding is reduced in Delta$^{∆β1-2}$, but not to the Serrate equivalent using liposomes composed of PC:PS:PE-fluorescein (80:15:5). Notch binding to *Drosophila* Delta NE3 (**E**) WT and ∆β1-2 (Delta$^{∆β1-2}$) variants both bind to Notch, unlike variant with F204A substitution in DSL domain. Comparisons were performed with a two-tailed unpaired t-test. Values are shown as scattered data points with the dark lines representing the means. ns, no significant difference, **P < 0.01; ***P < 0.0001.

Figure 2. DI β1-2 loop mutant generated by genome editing.

**(A)** Two gRNAs flanking the DI Exon 2 were used to replace the exon with a modified version where 5 amino-acids in the β1-2 loop were removed. Red lettering highlights the genomic sequence of the β1-2 loop.

**(B-C)** Adult wings from DI$^{∆β1-2}$ flies. No defects are detected in wings from DI$^{∆β1-2}$/+ (**B**), Homozygous DI$^{∆β1-2}$/ DI$^{∆β1-2}$ have extra vein tissue near L5 and uneven L2 veins (arrowheads; C).
(D) Microchaetae are arranged in rows on the thorax of control (yw) flies; these become disordered and more dense in $Dl^{b1-2}/Dl^{b1-2}$. White rectangle indicates area scored for E

(E) Number of microchaetes per central area (white rectangle in D) in the indicated genotypes.

Data information: ***P<0.0001 (Unpaired t-test). Each dot represents an individual fly, and light or dark shading indicates individuals from independent genetic crosses.

On the violin plot, dashed line represents the median and the dotted lines show the quartiles. Scale bars correspond to 200 µm (B-C) and 500 µm (D).

**Figure 3.** $Dl^{b1-2}$ enhances vein thickening from loss of function $Dl$ alleles and suppresses Notch phenotype.

(A-C) Representative images of adult females wings in combinations of $Dl^{b1-2}$ with loss of function $Delta$ alleles. In combinations with $Dl^{rev10}$ (A), $Dl^{Exon2}$ (B), or $Df(3R)Dl^{Fx3}$ (C), vein thickening is strongly enhanced (right panels) compared to heterozygous mutants alone (left panels). Vertical square brackets indicate the regions used for vein thickness quantification.

(D) Quantification of wing-vein thickness in females of the indicated genotypes.

(E) Representative images of adult female wings demonstrate that $Dl^{b1-2}$ rescues the wing notching phenotype, caused by a Notch loss of function allele ($N^{55e11}$). Horizontal square bracket indicates the L5 vein “delta” at the intersection with the wing margin analyzed in EV2E.

(F) Quantification of wing notching in females of the indicated genotypes, $Dl^{b1-2}$ rescues notching in a similar manner to $Dl^{rev10}$.

Data information: ***, P<0.0001(Unpaired t-test). Light, dark shading indicates data points from independent genetic crosses. On the violin plots, dashed lines represent the median and the dotted lines show the quartiles. Scale bars A-E correspond to 200 µm.

**Figure 4.** $Dl^{b1-2}$ has compromised Notch response in photoreceptor fate decisions.
(A) Schematic representation of Notch reporter $E(spl)m\tilde{5}0.5$ expression during photoreceptor differentiation. Expression is initiated in R3 and R4 of the 5-cell pre-cluster and becomes restricted to R4 as Notch activity resolves. Light orange indicates photoreceptors with R3 in light green and R4 in dark green. MF marks the morphogenetic furrow, boxed region indicates the region shown in B.

(B) Equatorial region of eye imaginal discs where $E(spl)m\tilde{5}0.5$ expression (green) becomes restricted to a single photoreceptor in each cluster (magenta), as detected in control and $Dl^{1/21-2/1}$ discs (top panels). In $Df(3R)Df^{Fx3}$/+ and $Dl^{1/21-2/Df(3R)Df^{Fx3}}$ discs (bottom panels) $E(spl)m\tilde{5}0.5$ expression is reduced ($Df(3R)Df^{Fx3}$/+) or absent from several clusters ($Dl^{1/21-2/Df(3R)Df^{Fx3}}$) indicative of reduced Notch signalling. Scale bars correspond to 10 µm.

(C) Proportion of photoreceptors clusters that fail to express the $E(spl)m\tilde{5}0.5$ reporter in the indicated genotypes. ns, no significant difference, *** P<0.0001 (one-way ANOVA). On the violin plot, dashed line represents the median and the dotted lines show the quartiles.

Figure 5. $Dl^{1/21-2}$ exhibits normal sub-cellular localisation.

(A) Apical view of wing imaginal disc with homozygous $Dl^{1/21-2}$ clones (GFP negative) stained for Di (red) and Cadherin (blue). (A') Z-projection of apical layers spanning a $Dl^{1/21-2}$ clone (GFP negative) located at the DV boundary. No change in apical localisation of Di (red) and Cadherin (blue) is detected. Panels on the right show the Di (grayscale) apical localisation on the wing imaginal disc.

(B) Uptake assay at t=0. After exposure to extracellular anti-Di antibody, Di protein (red) is detected at similar levels apical to Cadherin (blue) in wild-type (GFP) and $Dl^{1/21-2}$ mutant tissue (GFP negative). (B') Cross-sectional view of B, Di protein (red) is present apically relative to Cadherin (Blue) which marks adherens junctions. Middle and lower panels show the crosssection in grayscale of the apical marker Cadherin (Cad) and Delta (Di), respectively.

(C) Uptake assay after 30 minutes, internalised anti-Di (red) enters the endocytic route and in the cross sectional view (C') can similarly be detected as puncta along the cell axis in wild-type (GFP) and homozygous $Dl^{1/21-2}$ (GFP negative) tissue. Middle and lower panels show the crosssection in grayscale of the apical marker Cadherin (Cad) and Delta (Di), respectively.
Figure 6. Schematic summarizing roles for ligand β1-2 loop.

(A, B) The C2, DSL domains and N-terminal EGF repeat in Delta (cyan) and the ligand binding region (EGF11-13) of Notch (magenta) are renditions from the structures obtained (Figure 1), other regions of the molecules are represented not to scale. The amino acids in the β1-2 loop of Delta are highlighted in yellow (A). (A) The interaction of Delta (cyan) in trans with the Notch receptor (magenta) is augmented by the C2 domain, possibly through contacts of the β1-2 loop (yellow), with the “receiving” cell membrane, to yield highest levels of signalling (black arrow; green indicates ligand induced cleavages). Phospholipid contacts from β1-2 loop in the same cell could also influence cis-interactions between Delta and Notch in the same cell. (B) A deletion of 5 amino acids within C2 domain β1-2 loop (no yellow) disrupts phospholipid interactions but does not prevent Delta from interacting with Notch. Activation of Notch signalling is weakened (grey arrow) and phenotypes from transheterozygous combinations suggest that cis-interactions between Delta and Notch are also modulated.
Table 1. Data collection and refinement statistics.

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Statistics for the highest-resolution shell are shown in parentheses.
Expanded View Figures: Titles and legends

Figure EV1. Individual structures and expression and purification of the DI variants and Notch in vitro.

(A) Comparison of β1-2 loop sequences from Drosophila and human Notch ligands. Disulphide bond connectivity within this region is represented by the yellow brackets; (B-E) The crystallographic asymmetric units are shown for the three structures (B) Delta - one copy of C2-DSL-EGF1 construct; (C) Serrate - two copies of C2-DSL-EGF1-EGF2 construct; (D) C2 domain of Drosophila Delta (rainbow coloring) overlaid on the same region of Drosophila Serrate (grey). Disulphide bonds are shown as yellow sticks and the bound calcium ion in the Serrate structure as a grey sphere. The β1-2 loop (blue for Delta, at top) adopts a different conformation in Delta than in Serrate despite conservation of the two stabilizing disulfides; (E) Notch - one copy of EGF11-13. All proteins are shown in a rainbow cartoon representation coloured from blue at the N-terminus of the construct to red at the C-terminus, except Serrate in D (grey). In B, C, E, Glycosylation is shown in a stick representation as are the side chains to which the glycans are attached and disulphide bonds. Figure drawn using PyMol (The PyMOL Molecular Graphics System, Version 2.0 Schrödinger, LLC.)

(F) Monomeric Drosophila Notch ligand (N-EGF3) and receptor (EGF11-13) fragments were expressed in S2 cells as C-terminal His-tagged fusion proteins, purified by IMAC and SEC and visualised by reducing (R) and non-reducing (NR) 10%SDS-PAGE. The DI variants expressed for in vitro experiments were DlWT as control, DlΔβ1-2 where the β1-2 loop was deleted, Dlβ1-2,5AA where the β1-2 loop was mutated to 5 Alanines. DlF204A was used as a negative control as this variant reduces Notch binding at Site 2, by altering a key residue within the ligand DSL domain. Additional weak bands in the DlΔβ1-2 sample, detected in both non-reducing and reducing conditions, indicate carry through of slight impurities.

Figure EV2. Ser β1-2 loop mutant enhances vein phenotypes induced by reduced DI and Ser activities.

(A) Strategy to remove β1-2 loop sequence from Exon 3 in Serrate. Red lettering highlights the sequence of the β1-2 loop of Ser.

(B) Adult wings from SerΔβ1-2 heterozygotes have no visible phenotype.
(C) Wings from $Dl^{1/11-2}$ and $Ser^{1/11-2}$ alleles in combination with $Df^{rev10};Ser^{Rx106}$; both mutants enhance vein thickening. Black square bracket indicates the region used for vein thickness quantification.

(D) Quantification of vein thickening in wings from females of indicated genotypes; ***$p<0.0001$ (Unpaired t-test). Light, dark shading indicates data points from two independent replicates.

(E) Quantification of L5 vein width at intersection with the wing margin of indicated genotypes; $Df^{rev10}$ strongly suppressed the small “delta” produced by $N^{55se11} /+$ ($P<0.0001$, one-way ANOVA), combinations with $Dl^{1/11-2}$ and $Ser^{1/11-2}$ resulted in mild and variable suppression of borderline significance (ns and $P<0.05$ respectively, one-way ANOVA). On the violin plots, dashed lines represent the median and the dotted lines show the quartiles.

(F) Gene diagram showing that the two gRNAs used to replace the Dl Exon 2 also generated a novel allele where the Exon2 was removed, $Dl^{ΔExon2}$. $Dl^{ΔExon2}$ mutant clones (green) in wing imaginal discs stained for Dl (red) and the apical marker Cadherin (blue); Dl protein is absent in the mutant cells, indicating that little or no stable protein is made most likely because the signal sequence has been deleted.

Data information: Scale bars represent 200µm(B,C), 50µm (F) and 10µm in the magnified image of F.

Figure EV3. Expression of Notch targets cut and dpn is unaffected by $Dl^{1/11-2}$ or $Ser^{1/11-2}$.

(A) Expression of Cut at the wing dorsal-ventral (DV) boundary is unperturbed in wing discs of the genotypes indicated.

(B-C) Cut (red) expression is similar in wild-type (MARCM>ry, green, B) or $Ser^{1/11-2}$ homozygous (MARCM>$Ser^{1/11-2}$, green, C) clones that intersect the DV boundary. Individual Cut expression for each experiment is shown in grayscale on the right subpanels.

(D,E) $Dl^{1/11-2}$ homozygous clones (green) that intersect the DV boundary retain expression of Dpn (red), higher magnification in (E). Individual Dpn expression is shown in grayscale on the right subpanels.

Data information: Scale bars: 50 µm (A-D) or 10 µm (E).
**Figure EV4.** $Dl^{A\beta-2}$ compromises signalling during microchaete selection.

(A-D) Microchaete distribution in the notum of adult flies of the following genotypes:

(A) Control, yw; (B) $Df(3R)Dl^{Fx3/+}$; (C) $Dl^{A\beta-2}/Dl^{A\beta-2}$; (D) $Df(3R)Dl^{Fx3}/Dl^{A\beta-2}$.

Homozygous $Dl^{A\beta-2}$ mutants and $Dl^{A\beta-2}$ combined with $Df(3R)Dl^{Fx3}$ increases the number of the microchaetes in the notum.

(E) Quantification of microchaete numbers which were scored in a central region (white rectangle in Fig. 2D) , ***$p<0.0001$ (Unpaired t-test). Each dot represents an individual fly, and light or dark shading indicates data points from two independent crosses. On the violin plot, dashed line represents the median and the dotted lines show the quartiles.

Data information: Scale bars (A-D) correspond to 500 µm.

**Figure EV5.** $E(spl)m\alpha0.5$ expression is retained in homozygous $Dl^{A\beta-2}$ clones.

(A) $Dl^{A\beta-2}$ homozygous clones (green) in eye imaginal discs with $E(spl)m\alpha0.5$ (red) reporter expression, ELAV (blue) marks all photoreceptors. (A’A’”) Higher magnification of the equatorial region (marked by white square in A), $E(spl)m\alpha0.5$ (red, A’; white, A’”) is detected in R4 of wild type and mutant clusters, including clusters where R3 is mutant.

(B) Large homozygous $Dl^{A\beta-2}$ clone (green), $E(spl)m\alpha0.5$ (red) expression is unaffected, ELAV (blue) marks all photoreceptors. (B’,B’”) Higher magnification of the equatorial region (marked by white square in B), $E(spl)m\alpha0.5$ (red, B’; white, B’”) is detected in R4 of wild type and mutant clusters, including clusters where R3 is mutant.

Data information: Scale bars correspond to 50 µm (A,B) or 10 µm (A’,B’).
Table EV1 - Primers list

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| Delta HA1 Rev | GTTAAGATGCGAGGTGCTCGACTTCTGTTTGGAGACCGC |
| Delta Exon2 Fw | CTATCTTCTAGGTAGGTTAAGGAAAGCCAAGCGAAGGTAAT |
| Delta Exon2 Rev | AACGTTATAAAGAAAGACAAACAAAAATATATTATGACTGATGAG |
| Delta HA2 Fw | TCAATCGAGGACAAATAATTTTTCTCTTACGTTAGTACAGAAAAATAGGCTAG |
| Delta HA2 Rev | TTCTTCCCGGGGAAGAAGCGCTTTATGCTACTGCTGTGCTAAATACTT |
| pH_DsRed_Dl bb 1 Fw | ATTTATGCCAGTAGCAATAACGCTCTTCCCCGGGAAAAGA |
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| pH_DsRed_Dl bb 2 Fw | GGCTTCAAAACAC AAAATCGAGCAGACCTGCAGTCTTAAAC |
| pH_DsRed_Dl bb 2 Rev | ACAGAATTAATATACTGCACTGCCCCGGGAA |
| Serrate HA1 Fw | CCAATTGCAGGTGCCTTTTGAAGGCCTAAACGAAAAAGA |
| Serrate HA1 Rev | TAAGATGCAGGGCTCTTCCCTGGCAGTTACATT |
| Serrate Exon3 Fw | TTCTAGGTTAAGGAAATTTGGGCTCAGGAGGAGGAG |
| Serrate Exon3 Rev | AGACTGCATTTTTACGTCTTTTACGGATAGGCTAG |
| Serrate HA2 Fw | TCGTTGTCAAAAGAAAAATATTAGGCTGAAAATGCGTCTTATTAGTGAC |
| Serrate HA2 Rev | TTCCGGCGGAGAAGCGGCGCCCATGATTCTGGTGC |
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<p>| <strong>Mutation of the β1-2 loops</strong> | |
| Deletion of DI β1-2 loop Fw | TGCCCGTGCAGGGAGTCTGGTC |
| Deletion of DI β1-2 loop Rev | GCTGGAGCTCTCAGGAGTCTGGTC |
| Deletion of Ser β1-2 loop Fw | ACGATAGGTAAGTCCATCTTTTAAGTTTATTAAAGGCCC |</p>
<table>
<thead>
<tr>
<th>Deletion of Ser β1-2 loop</th>
<th>TTCCGCTGGCATGCCG</th>
</tr>
</thead>
</table>

**Sequencing of the transformants**

<table>
<thead>
<tr>
<th>DI Ex2 seq</th>
<th>GTTCACAGTTCCGCG</th>
</tr>
</thead>
<tbody>
<tr>
<td>DI UP5 Fw</td>
<td>TGCCTGTGGCTGCTTC</td>
</tr>
<tr>
<td>DI 3Down Rev</td>
<td>AGGGAGAGGGATCGAGATAGAG</td>
</tr>
<tr>
<td>Ser Up5 Fw</td>
<td>TAGTAAGCAGCAAGAACCGGT</td>
</tr>
<tr>
<td>Ser 3Down Rev</td>
<td>ATCTCAGCAGGAGGATAG</td>
</tr>
<tr>
<td>Transp Rev</td>
<td>CAGACCGATAAAACACATGC</td>
</tr>
<tr>
<td>Transp Fw</td>
<td>ATGCACAGCGAGGATTC</td>
</tr>
</tbody>
</table>

**Protein expression**

<table>
<thead>
<tr>
<th>Delta β1-2 deletion Fw</th>
<th>AGCGGAGAGAGATGAGGCTGCTGGGAGTTGCAAGAC</th>
</tr>
</thead>
<tbody>
<tr>
<td>Delta β1-2 deletion Rev</td>
<td>GCAACTGCCAATCATCTCTCTCGGCTGAGG</td>
</tr>
<tr>
<td>Serrate β1-2 deletion Fw</td>
<td>GCAGCATGCCAGCGGAAACGGATAGGCTGCTGCGCATG</td>
</tr>
<tr>
<td>Serrate β1-2 deletion Rev</td>
<td>GCGAGCAGCCTATCGTTTCCGCTGGGATGCGG</td>
</tr>
<tr>
<td>Delta F204A Fw</td>
<td>GACGATAGCGCGGAGTTGCAAGAC</td>
</tr>
<tr>
<td>Delta F204A Rev</td>
<td>GTGGCCGGCGCTATCGTGCTGGGAG</td>
</tr>
<tr>
<td>pExS2-2 fusion Fw</td>
<td>GAAGGATGATCAGCTGAGCTGAGCT</td>
</tr>
<tr>
<td>pExS2-2 fusion Rev</td>
<td>CAAACTAAGATTTAGTTCAGATATCAG</td>
</tr>
</tbody>
</table>

A Delta:Notch
DII4:Notch-1 (pdb 4xlw)
B Serrate:Notch
Jagged-1:Notch-1 (pdb 5uk5)
C
D Liposome binding
E Notch binding
A

Strong NICD signal
Delta
Notch

Sending cell
Receiving cell

B

Weakened NICD signal
Delta
Notch

Sending cell
Receiving cell