

Notch Signalling in context

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Sarah Bray is Professor of Developmental Biology at the University of Cambridge. Her career began with studies of translational regulation in sea urchin embryos (with Tim Hunt), for her PhD. From there she moved to Harvard for her postdoctoral research, working with two groups (Jay Hirsh and Fotis Kafatos) that were harnessing transgenesis to study gene regulation. There she discovered the first member of the Grainy head family of transcription factors. Her research into the Notch pathway began when she returned to University of Cambridge to set up her own group. Their early work investigating Notch pathway functions in *Drosophila* began to highlight the diverse roles that the pathway plays in development. Since then her group have continued to investigate the basis for this diversity, with an emphasis on the mechanisms at the transcriptional heart of the pathway.

Key Points:

- The highly conserved Notch cell-cell signalling pathway operates in many different contexts where the consequences can differ widely, despite the fact that the core pathway is very simple.
- Many different types of regulation contribute to the differing outcomes, ranging from tissue level co-ordination to nuclear governance.
- The pattern of expression of the ligands, receptors and critical modifying enzymes is one level of regulation that is common to many signalling pathways. However, the one-to-one interaction between ligand and receptor places extra emphasis on this, especially as they can cis-inhibit one another when present in the same cells.
- 'Topological' tissue organization and the extent of cell-cell contacts is likely to be of unusual significance in influencing the levels of Notch activation, because the ligands are trans-membrane proteins.
- Nuclear context, in the form of cell-type specific transcription factors and chromatin organization is a primary level of control in generating qualitatively different outcomes from Notch activation. In addition, the wiring of the regulatory network within the signal receiving cells contributes to the diversity of responses and to the nature of its cross-talk with other signalling pathways.
- Together the mechanisms make the Notch pathway versatile and able to undertake many different roles. But they are also susceptible to perturbations, and may be a contributory factor in Notch-related diseases.

Abstract

Since it became evident that the highly conserved Notch signalling pathway functions in many different developmental and homeostatic processes, questions have arisen about how this pathway can achieve such diverse outcomes. With a direct route from the membrane to the nucleus, the Notch pathway has fewer opportunities for regulation than many other signalling pathways, yet it generates exquisitely patterned structures. More confusingly, its activity promotes growth in some circumstances but cell death in others. This review will consider the regulatory mechanisms that shape the activity of the Notch pathway and its outcome, enabling it to generate biological consequences that are appropriate for each context.

Introduction:

Despite the fact that the core Notch pathway operates in vastly different developmental and disease contexts, from stem cell regulation and heart morphogenesis to cancers and cardiomyopathies, it is relatively simple in its operation. Ligand-mediated activation induces a series of proteolytic cleavages in members of the Notch family of receptors, which release the Notch intracellular domain, NICD;. (**Figure 1a** and **Box 1**). Once released, the NICD enters the nucleus and, together with the DNA-binding protein CSL (CBF1/Suppressor of Hairless/Lag-1) and the co-activator Mastermind (Mam), stimulates transcription of target genes (**Figure 1a**)¹⁻⁴. Thus, no amplification of the signal can occur, unlike in many other pathways, and no intermediates are present between the membrane and the nucleus —NICD takes responsibility for implementing pathway activation.

Given this relative simplicity — receptor–ligand interactions release the bioactive NICD — how can the canonical Notch pathway coordinate so many diverse biological outcomes? One Notch receptor and two ligands — Delta and Serrate (Ser) — exist in *Drosophila melanogaster* while vertebrates have four Notch paralogues (Notch1–4) and a similar diversification of ligands (referred to as Delta-like ligands (DLLs) and Jagged in mammals). This repertoire clearly cannot account for the diversity of Notch pathway signalling outcomes, indicating that other mechanisms of regulation exist. In this Review, we will illustrate features that enable the Notch pathway to function differently according to its setting and that help to explain the myriad of roles that the pathway has in development and disease. It is only possible to draw on a small subset of examples owing to space limitations, but these examples highlight fundamental principles that should be widely applicable across the different contexts where the Notch pathway operates.

Ligand-receptor landscapes [L1 Heading]

Early models suggested that Notch-mediated developmental patterning occurred when stochastic differences in ligand–receptor interactions directed signalling within fields of cells with near uniform levels of ligand expression. It now seems that this scenario is the exception rather than the rule: the expression profiles of ligands, receptors and several modifying enzymes have important roles in defining the outcome. Furthermore, other signalling pathways can regulate these expression patterns to augment, inhibit or modulate Notch pathway activity, providing an important mechanism of crosstalk.

Expression of canonical ligands and receptors [L2 heading]

The classic paradigm of lateral inhibition, which is a common phenomenon during precursor patterning, assumed that the driving force for signalling arose through stochastic differences in ligand levels. The initial bias would be reinforced by a negative-feedback loop so that ligand expression became repressed in signal-receiving cells, generating a spaced distribution of signalling cells and receiving cells that explained the distribution of precursors⁵. It now seems that signalling rarely

relies on uniform ligands; rather, their spatial and temporal regulation directs Notch signalling profiles in many contexts and is an important mode for regulation by other signalling pathways. Although much regulation of ligand expression occurs at the transcriptional level, micro-RNAs (miRs) also contribute. For example, miR8/miR200 targets the 3'UTRs of Ser and Jagged1 (Jag1) to fine-tune the levels of protein produced in *D. melanogaster* and in human tissues, respectively^{6,7}.

One example of how ligand expression dictates the spatial pattern of signalling occurs during the formation of the growth organizer in *D. melanogaster* wing imaginal discs. Here, Ser is produced exclusively in the dorsal territory owing to its regulation by the spatially restricted Apterous transcription factor^{8,9}, and specifically generates a stripe of Notch activity in adjacent ventral cells to create a boundary that organizes growth of the tissue⁸. Dynamic changes in expression of both ligand and receptor also help to drive oscillations in Notch signaling¹⁰ such as occur during somitogenesis in the presomitic mesoderm. Here, Notch1 expression is dependent on its' pathway activity which might help to reinforce signalling, whereas ligand expression is Wnt regulated, thus ensuring that the pattern of Notch activity is instructed by another main component involved in the somite clock^{11,12}.

In addition, interplay between different ligands is frequently required to set the correct balance of precursor cells in a number of situations. For example, in the ear, a broad 'inductive' signal by Ser/ Jag1 confers the neurogenic potential of the placode^{13,14} before subsequent dispersed expression of DLLs in emerging precursors inhibits the surrounding cells to ensure a single precursor is generated at each position¹⁵⁻¹⁸. Likewise, to establish the branching pattern of blood vessels the production of DLL4 in tip cells prevents neighbouring cells from adopting the same fate^{19,20}, whereas Jag1 acts as a potent proangiogenic regulator that antagonizes DLL4–Notch signalling to favour new sprouting.

Although the fundamental consequences of ligand binding are the same -- cleavage of Notch receptors to produce NICD -- they nevertheless often elicit different outcomes²¹. One possibility is that different ligands bring about different strengths (or durations) of intracellular signal. For example, Jag1 and DLL1 have lower measured affinities for Notch1 than does DLL4, possibly owing to differences in the orientation of the amino -terminal and Delta/Serrate/Lag-2 (DSL) domains involved in the binding interface²². Indeed, in the haemangiogenic endothelium, specification of haematopoietic stem cells involves a low-strength Jag1-dependent signal whereas specification of endothelial arterial cells requires a high-strength Dll4 signal²³. Strikingly, cells that received the low-strength Jag1-induced Notch activity appeared unresponsive to a high-strength DLL4 signal. Although it is unclear how this lack of response arises, if the expression of Jag1 precedes that of DLL4, it could switch on an inhibitory programme or cause cell re-arrangements to disrupt contacts with DLL4. Similarly, in the inner ear Jag1 might elicit a lower level of Notch1 activation than DLL1 does, as Jag1 can induce the expression of Hey1, which requires a low threshold, but not Hes5, which requires a higher level of signalling²⁴.

In animals with multiple Notch paralogues, the deployment of different receptors will also influence signalling outcomes. In humans, mutations in different paralogues have different disease consequences (for example, only defects in Notch3 cause CADASIL (cerebral autosomal-dominant arteriopathy with subcortical infarcts and leukoencephalopathy), emphasizing different receptor roles^{21,25,26}). Although these differences might be partly explained by variations in their expression patterns, examples exist where individual paralogues — for example, Notch1 and Notch2 — make different contributions even when expressed in identical patterns. Indeed, Notch1 and Notch2 were found to have opposite effects on the growth of one specific

tumor cell-type²⁷. However, when the intracellular domains of Notch1 and Notch2 were swapped within each normal gene context their intrinsic activities were similar^{28, 29} but, in some contexts, the amount or stability of the NICD moieties differed²⁹. This may reflect differences in their potential for post-translational modifications. These observations emphasize the possibility that, ultimately, the amount or duration of the NICD 'signal' produced will depend on the specific ligand–receptor pairs that are engaged, as well as the availability of the metalloproteases essential for the activating cleavages. Whether such NICD differences would result in quantitative or in qualitative differences in the downstream signalling output is an important question that remains to be resolved.

Contributions from *cis* inhibition [L2 heading]

Notch signalling is also highly sensitive to the relative levels of ligands and receptors owing to *cis* inhibition, an inhibitory interaction that occurs when ligand and receptor are present in the same cell (**Figure 2**)³⁰. First identified through genetic experiments in *D. melanogaster*^{31, 32}, the precise mechanism for *cis* inhibition remains elusive, although some studies suggest that Notch molecules are targeted for degradation once they have undergone ligand interaction^{33 34}.

Modelling experiments suggest that an ultrasensitive switch between two mutually exclusive cell states: signal-sending and signal-receiving could be generated by Notch having a sharp ligand threshold for *cis* inhibition combined with a graded response to *trans*-acting ligand³⁵. One example of a *cis*-inhibitory interaction affecting the signalling outcome occurs during photoreceptor specification in *D. melanogaster*: here, loss of DI-mediated *cis* inhibition reversed the direction of lateral signalling, thereby generating the wrong complement of photoreceptors³⁶. Likewise, *cis* inhibition helps to stabilize tip and stalk fates during angiogenesis, and hence prevents hybrid tip–stalk cells forming³⁷. In addition, different Notch receptors might also *cis* inhibit one another, adding another potential mechanism to fine-tune signal reception³⁸. The balance between *cis*-interactions and *trans*-interactions is thus likely to be important in signalling outcomes (**Figure 2a**).

Intriguingly, the mammalian ligand DLL3 might operate only in a *cis*-inhibitory mode. In cell-based assays, DLL3 was unable to activate signalling in *trans* but, when co-expressed with Notch1, it could prevent Notch1 responding to other ligands on neighbouring cells³⁹. *In vivo*, the loss of DLL3 led to increased Notch activity during T-cell development⁴⁰ and to defects in Notch1 signalling in the presomitic mesoderm⁴¹. It is not fully clear why DLL3 might have a uniquely inhibitory role, although its highly divergent DSL domain might be a contributory factor; however, by doing so, it adds another strand of pathway regulation in vertebrates.

Deployment of Fringes and other modifying enzymes [L2 heading]

Further influencing Notch signalling is the presence of enzymes that modify the extracellular domains (ECDs) of ligands and receptors and modulate their ability to signal (**Figure 2b**). Although modifications to ligands might be important, most focus has been on receptors, the activity of which is profoundly affected by glycosylation of EGF repeats in the ECD. O-fucosyltransferases, which add fucose to serine and threonine residues, and O-glucosyltransferases, which add glucose to serines, are essential for optimal Notch signalling^{42, 43}. Many O-fucose monosaccharides on Notch can subsequently be extended (with N-acetylglucosamine) by the Fringe proteins, with differing effects on Notch, depending on the sites modified and the ligands present⁴².

Fringe-mediated modifications could influence specific ligand–receptor interactions. Structural studies indicate that elongation of an O-fucose on EGF repeat 12 of Notch

receptors could provide additional energetic contributions to Notch–ligand interfaces²². This modification enhanced binding of DLL1 and Jag1 to a greater extent than binding to DLL4, possibly because the inherent affinity of DLL4 is relatively high even in the absence of glycosylation²². Whether or not the enhanced ligand binding translates into increased pathway activity is uncertain. In some cases, where the affinity of Jag1 for the receptor was increased by the presence of Fringe proteins, the effects on transcriptional output were reduced⁴⁴. Concomitant changes to *cis* inhibition, to competition with other ligands or to other sites in the receptor are possible explanations for this apparent anomaly.

Fringe proteins are thus likely to modulate the ability of cells to send and receive signals in a manner that is highly dependent on the cocktail of ligands (and Fringe proteins) present (**Figure 2b**). For example, in *D. melanogaster* wing discs, modifications by Fringe render Notch insensitive to Ser⁴⁵. Here, co-expression of Ser and Fringe in dorsal cells guarantees that Ser can only signal to the adjacent ventral territory⁸. At the same time, Dll is enriched in ventral cells and its binding is enhanced by Fringe modifications^{45,46}. The combined effects generate a stripe of cells with Notch activity that straddles the boundary. Likewise, when high levels of both Jag1 and Fringe are present in stalk cells, Fringe is thought to make Jag1 into an effective competitor for the more signalling proficient DLL4, which prevents DLL4 from signalling between adjacent stalk cells, and thereby inhibits excessive sprouting⁴⁷. Similarly, during ventricular development, temporal modulation of Mfng enables sequential Notch activation to drive different morphological processes; MFng and Dll4 downregulation in the endocardium allows these cells to respond to myocardial Jag1/ Jag2 and generate a functional ventricular wall⁴⁸. Finally, in the spinal cord, where Fringes enhance DLL-activated Notch signalling and block that of Jag1, the consequence of their patterned expression (controlled by homeodomain proteins) is domain-wide Notch activation by either Dll or Jag1, and a suppression of signalling across progenitor domain boundaries⁴⁹.

Experiments modelling the outcome of Fringe modifications have illustrated how these could impact on the relative signalling capabilities of cells. For example, cells expressing Lunatic Fringe (Lfng) or Manic Fringe (Mfng) in combination with Jag1 and Notch1 acquired the ability to send and receive signals simultaneously, but only using different ligands⁵⁰. This observation could be explained if Fringe modifications weaken *cis* interactions between Jag1 and Notch1, consequently making high surface levels of both available, and also prevent Notch1 from being *trans* activated by Jag1 (**Figure 2b**). Notch1 could thus only receive signals from DLL on adjacent cells but Jag1 would itself be free to signal to neighbouring cells. Such a model fits well with the observations at the dorsal–ventral boundary of the *D. melanogaster* wing and with the tip/stalk decision in angiogenesis, and highlights the profound effect of the patterned deployment of these molecules on signalling outcomes.

Topological context [L1 heading]

Notch receptors and their ligands are transmembrane proteins. Furthermore, endocytosis is known to be required for ligand activity^{51,52}. Mechanisms that transport the proteins to and from the correct places in the cell are therefore likely to have an important impact on signalling. Although it has largely been assumed that ligand–receptor interactions occur at sites where cells are tightly opposed, several observations are challenging this assumption, and highlight that the dimensions and stability of the contact area might have important roles.

Endocytic trafficking and ubiquitin ligases [L2 heading]

Factors that modulate endocytosis and trafficking of receptors and ligands have a number of important consequences on pathway activity^{53, 54}. First, endocytosis of the ligand after it has engaged with its Notch receptor is thought to generate the force on the receptor that exposes the protease cleavage site within the negative regulatory (NRR) 'cage',^{55, 56, 57} **(Figure 1b)**. In the absence of certain E3 ubiquitin ligases of the Mindbomb or Neuralized families, ligand endocytosis is prevented with concomitant loss of signalling^{58, 59, 60-63, 64}. As ubiquitylation of the ligand intracellular domains by these enzymes is critical for ligand activity, the presence of Mib and/or Neur determines which cells can send a signal. During asymmetric division, for example, directional Notch signalling is achieved, in part, by the polarized segregation of these E3 ligases into one of two daughter cells, depending on the underlying cell polarity^{65, 66, 67}.

Second, receptor trafficking affects pathway activity not only by determining receptor levels on the cell surface but also because ligand-independent activation can occur during this process^{53, 54, 68}. Normally, while some of the endocytosed Notch is likely recycled to the membrane, a large fraction is targeted for degradation. When this fraction of the receptor fails to be properly routed into the inner luminal vesicles of maturing lysosomes — for example, in response to mutations that affect the ESCRT complex⁶⁹⁻⁷² this can result in ligand-independent Notch activation, although the precise mechanisms that underly this phenomenon are unknown. Most likely, the conditions encountered in the lysosome promote ligand-independent activation by destabilizing the NRR. Proteins that regulate endolysosomal transport have a concomitant effect on Notch trafficking that can, in some cases, result in receptor activation in normal physiological conditions. These include the E3 ligases Deltex and Itch/Suppressor of Deltex, which, by ubiquitylating Notch, alter both the receptor levels on the membrane and the amount of ligand-independent activation⁷³⁻⁷⁵.

Numb is another factor that regulates cell-fate decisions via its powerful effects on receptor trafficking; its depletion in several lineages results in ectopic Notch activity. During sensory organ development, Numb is differentially segregated into one of two daughter cells, in which it inhibits Notch activity to bring about specific 'Notch off' fates⁷⁶⁻⁸⁰. Numb achieves this inhibition either by promoting Notch internalization or by altering the route taken after Notch is endocytosed^{81, 82}. Notably, this phenomenon only occurs in certain contexts, despite Numb being present more widely. In *D. melanogaster*, this specificity depends on the adaptor Sanpodo, which couples Numb localization with Notch trafficking⁸²⁻⁸⁴. Adaptors analogous to Sanpodo are likely to perform a similar task in other species.

Third, spatial regulation of trafficking also has the potential to affect the geometry of signalling. For example, endocytic depletion of Notch from the cleavage furrow after the division of sensory organ precursor cells in *D. melanogaster* is important to enable unidirectional signalling in the progeny⁸⁵. The asymmetrical distribution of endosomes containing receptors and/or ligands is also associated with a bias in signalling to promote specific cell fates in neural lineages^{86, 87}. In addition, by affecting the activity and/or localization of the γ -secretase complex, which is required for the release of the NICD, proteins that organize cellular polarity (for example, Crumbs⁸⁸ and hibiris⁸⁹) can consequently affect Notch signalling. Finally, beyond regulating the activity of the ligands and receptor, endocytic trafficking might also be important for localizing functional pools of ligand and receptor to the appropriate cellular subdomains, although many questions remain about where these subdomains are located.

Tissue architecture and morphology [L2 heading]

Because Notch signalling occurs between cells that are in contact with one another, the organization of the tissue is likely to influence the levels or patterns of signalling, such that the strength and periodicity of signalling could depend on the extent or durability of the adhesive contacts between cells. For example, models suggest that, under conditions in which the extent of cell–cell contact is greater than the diffusion range of the ligand within the membrane, the signal generated will be proportional to the contact area⁹⁰. Signalling might also be mediated by dynamic cellular protrusions such as filopodia, which, importantly, could extend the distance over which signalling occurs⁹¹⁻⁹⁴. The nature of the cell–cell contacts could therefore have important consequences for the functional outcomes of signalling (**Figure 3a**).

Adherens junctions (AJs) mediate cell–cell adhesion and are important for effective Notch signalling in some contexts. Thus, during vertebrate neurogenesis, the dimensions of the contacts made by the ‘end-feet’ and the integrity of their AJs are important for nascent neurons to engage in effective DLL–Notch signalling with their progenitors to prevent them differentiating. Disrupting the AJs down regulated Notch signalling and caused precocious neurogenesis⁹⁵, whereas manipulations that expanded the size of the apical domain enhanced signalling and reduced neurogenesis⁹⁶.

Strong adhesion between posterior lateral mesoderm and somite cells, mediated by the junctional adhesion molecules Jam1a and Jam2a, is also important for generating a sufficiently high Notch signal to specify the haematopoietic lineage in zebrafish⁹⁷. The contact surface area between migrating lateral mesoderm cells and the somite was decreased when *jam1a* or *jam2a* was depleted, and correlated with decreased activation of Notch signalling. As this phenotype could be rescued by widespread expression of ligands, the transduction pathway remained functional, leading to the model that dimensions or stability of contacts was important. The deployment of specific adhesion molecules can thus create a unique topological opportunity for signalling to occur.

Signalling might not always require stable cell–cell contacts, transient interactions can be sufficient to deliver Notch activity and switch cells to a specific cell fate. Neural crest cells expressing DLL could elicit signalling in myotome Notch1-containing cells that they contacted, in a ‘kiss and run’ mode⁹⁸. Likewise, during angiogenic sprouting, because cells constantly shuffle their positions, a presumptive stalk cell might encounter high levels of Notch for only a brief period before losing contact⁹⁹ (**Figure 3b**). Transient structures may also extend the range over which a cell can signal. For example, filopodia extend from DI-producing cells on the *D. melanogaster* notum and their disruption (by specific genetic mutations) perturbed the spacing of sensory organ precursor cells (**Figure 3b**)⁹¹. Similarly, dynamic filopodia project between neurogenic progenitors and radial glial cells during signalling in the mammalian neocortical progenitor cell niche, although their functional relevance has not yet been tested¹⁰⁰. And during the formation and maintenance of pigmented stripes in zebrafish, signalling is mediated via long cellular protrusions that extend between xanophores and melanophores. Interestingly the geometry switches: first the protrusions carry vesicles of DLL from xanophores to promote melanophore stripe consolidation⁹³ (**Figure 3b**). Later, in adults, the melanophores extend protrusions towards interstripe DLL expressing xanophores to receive signals necessary for their own survival⁹⁴.

Filopodia-like protrusions are more transient than AJs, indicating that a prolonged stable contact point might not be essential for ligand–receptor signalling, although the extent of their contribution to Notch signalling remains to be determined. Modelling experiments suggest that the strength of the signal mediated through

filopodia will depend principally on the diffusion of ligands or receptors⁹⁰, so that factors that regulate their trafficking would be influential. Furthermore, it is possible that the types of downstream response might differ according to the way the ligand is presented, — for example, filopodia might provide a burst of signal whereas more stable cellular junctions could generate a more sustained signal.

Nuclear context [L1 heading]

The output of Notch signalling primarily relies on NICD entering the nucleus; mechanisms that set the nuclear context are therefore critically important as they determine the gene expression programme and consequent physiological outcome. For example, in the outer proliferation centre of the *D. melanogaster* optic lobes, Notch signalling induces neurons to die or to survive, depending on the transcription factors present¹⁰¹. Likewise, the contrasting oncogenic and tumor suppressor roles of Notch activity that occur in different tissue contexts are likely the consequences of differential cell-type specific transcriptional programmes¹⁰². Nuclear mechanisms that confer such context-specific Notch responses are potentially of broad relevance to other signalling pathways, although the precise components might differ.

Key features underpinning the nuclear response [L2 heading]

A key aspect of specificity will logically result from the presence of cell-type-specific or stage-type-specific transcription factors that alter the selection of responsive genes. Indeed, if CSL itself could function as a 'pioneer', binding to sites in dense nucleosome-covered 'closed' enhancers, it would be difficult to account for the cell-type specificity of responsive genes — how, then, could cell death genes be selected only in specific neuronal progeny to correctly programme neural networks, for example?¹⁰¹. In agreement with this notion, CSL has a higher affinity for motifs at the edge of nucleosomes, suggesting it binds preferentially to open chromatin¹⁰³. Furthermore, the motifs bound by CSL in *Drosophila* cells are located preferentially within regions of primed or active chromatin, making it likely that response specificity is aided by many motifs being hidden in chromatin environments that are not accessible to CSL¹⁰⁴ (**Figure 4**). However, even though the NICD–CSL complex prefers ready-primed chromatin regions, it is not just an idle passenger — the consequences of its recruitment include large-scale changes in histone acetylation, removal of repressive complexes and enhanced accessibility of the DNA¹⁰⁴⁻¹⁰⁸ (**Figure 4**).

The original 'switch' model proposed that, prior to Notch activation, CSL was present at enhancers in a complex with co-repressors that kept the enhancer silenced by recruiting histone deacetylases or other modifying enzymes¹⁰⁹. NICD was thought to displace the co-repressors from CSL to render the target enhancers active. However, this model in its simplest form has been challenged. First, the co-repressors KyoT2 and MINT bind to CSL with similar affinities as NICD does, making it hard to explain how NICD could displace them^{110, 111}. Resolving this question will however also require detailed knowledge of the relative stoichiometries of all the proteins concerned, including NICD. Second, CSL binding dramatically increases at target enhancers following Notch activation, arguing that this event is quite dynamic^{105, 112-114}. Thus, it is likely that an exchange of entire complexes (rather than co-activators substituting for co-repressors on DNA-resident CSL) occurs, similar to the model for hormone receptors, with the NICD–CSL complex being 'captured' to increase the amount or duration of its binding (**Figure 4**).

Although further studies are needed to determine the extent to which enhancers are 'marked' by CSL prior to Notch activation, CSL-binding co-repressors are nevertheless likely to contribute to the regulatory landscapes. First, loss or down-

modulation of CSL leads to derepression of tumour-promoting genes in several contexts^{115, 116}. Second, as several co-repressors bind directly to CSL in a manner that would preclude concomitant NICD binding, they will titrate the availability of CSL and hence set a threshold that NICD would need to exceed in order to activate transcription^{110, 111, 117-119}. Finally, the different types of co-repressor might give rise to different types of repression complex, especially if they differ in their ability to recruit chromatin-modifying complexes (**Box 2**). Thus, by binding to their target sites transiently, certain types of CSL-repressor complex might help to make the enhancers more refractory to the effects of NICD. Uncovering their contributions will be important for deciphering the regulatory landscapes that NICD encounters.

Cooperation with transcription factors

To ensure that the appropriate tissue-/cell-type response occurs, mechanisms that direct NICD to the appropriate enhancers must exist. Although the execution of some roles of Notch signalling relies on a set of common targets, the HES family of bHLH-containing proteins, other roles depend on diverse transcriptional responses¹²⁰. These responses are likely to be achieved through close cooperation between CSL complexes and other transcription factors (**Figure 5**), either because specific configurations of binding motifs allow direct interactions between transcription factors and NICD-CSL or because nearby motifs recruit transcription factors that help recruit NICD-CSL indirectly by modifying chromatin. At the same time, other transcription factors might block CSL recruitment to specific enhancers.

One well-characterized example of an enhancer 'signature' is the so-called SPS+A site in Notch-regulated genes during neural precursor specification. This signature combines a pair of specifically orientated CSL motifs (known as SPS) with a binding site for the pro-neural basic helix-loop-helix protein Daughterless (referred to as A) and allows NICD-CSL to interact directly with Daughterless, thereby conferring Notch-bHLH synergy¹²¹. Similarly, in *Caenorhabditis elegans*, the *ref-1* enhancer contains four predicted binding sites for GATA transcription factors that are required for Notch-dependent endodermal expression¹²² and that probably facilitate NICD-CSL recruitment via a direct interaction between CSL and the GATA factor. A third example involves the mouse *Foxp3* gene, in which an overlapping CSL-nuclear factor κ B (NF- κ B) binding site within the promoter facilitates cooperative regulation by Notch3 and canonical NF- κ B signalling¹²³.

By contrast, other examples indicate that cooperation can occur without a distinctive enhancer signature. An association between Runx proteins and NICD-CSL complexes exists in several cellular contexts, with CSL-bound regions enriched for Runx sites^{105, 108, 124} but without any very precise arrangement of their motifs. Runx proteins are required for CSL to be recruited to these enhancers and it seems likely that the mechanism involves changes in the local chromatin organization to 'expose' the CSL-binding motifs¹⁰⁴ (**Figure 4**). Other examples in which specific transcription factors are important exist (see **Box 2**), but it is as yet unclear whether direct or indirect mechanisms are involved and it remains to be determined whether subsets of transcription factors have a special relationship with NICD-CSL because they introduce a specific partner or chromatin conformation or, alternatively, whether any transcription factor binding in proximity to a CSL-binding motif might be sufficient to render an enhancer responsive.

Notably, binding of specific transcription factors can also prevent enhancers from responding to Notch. A well-characterized example is the zinc-finger transcription factor Ikaros, which restricts the Notch responsiveness of many T-cell targets, including *Hes1* and *Myc*, by binding to their enhancers¹²⁵⁻¹²⁷ (**Figure 4**). In the

absence of Ikaros, Notch target genes that are normally shut off in thymocytes were persistently expressed and other normally inactive or weakly upregulated genes became strongly induced by Notch¹²⁵. Conversely, re-expressing Ikaros could repress Notch1 target genes, including Myc, in T-cell acute lymphoblastic leukaemia (T-ALL) cells¹²⁷. Among other transcription factors that inhibit the activity of NICD–CSL complexes, several do so by binding to the complex rather than by blocking enhancer binding^{128, 129}. For example, the transcriptional repressor BCL6 inhibits the expression of ESR1¹³⁰ either by preventing the recruitment of Mam to NICD or by recruiting the histone deacetylase SIRT1 to promote deacetylation of neighbouring histones¹³¹. Finally, BEN-SOLO proteins can bind to nearby sites on DNA and directly contact CSL to antagonize Notch activity during neurogenesis¹³². Direct and indirect negative regulators are therefore likely to have widespread roles in setting the transcriptional landscape.

Chromatin context and indirect mechanisms [L2 heading]

If target enhancers need to be present in the appropriate chromatin complex to be bound by CSL, epigenetic mechanisms that alter the accessibility of enhancers should have an important influence on gene expression in responding cells¹³³, especially during developmental (and other) transitions when the Notch-responsive programme needs to change between one state and the next (**Figure 5**). Several examples illustrate that such transitions might be coordinated by stage-specific transcription factors in conjunction with chromatin-modifying complexes. For example, Hamlet/Evi1, when recruited to targets in nascent *D. melanogaster* olfactory neurons, appears to enable a modified response in a subsequent round of Notch signalling by altering histone methylation and density to erase the Notch state inherited from the parental cell¹³⁴. BCL6 similarly mediates stable epigenetic repression of Hes5 by recruiting SIRT1 during the neurogenic transition in mouse cortical progenitors¹³¹. Finally, Pax6/Eyeless blocks the ability of NICD to promote tumours in older generation progenitors in some *D. melanogaster* neural stem cell lineages, where it prevents transcriptional activation of direct target genes possibly through BRG1-associated factor (BAF)–SWI/SNF-related chromatin remodelling complex¹³⁵.

Polycomb complex-mediated silencing of target enhancers is also likely to shape the Notch response. Although their action is reversible, two multiprotein Polycomb repressive complexes (PRCs) confer heritable repressive states, and their presence at many Notch-regulated genes in embryonic stem cells indicates their potential importance¹³³. The inability of Notch to drive cardiac regeneration in adult rat cardiomyocytes is also attributed to PRC-mediated repression¹³⁶. Likewise, the activity of PRCs curtailed NICD-mediated activation of target genes in *D. melanogaster* cells and in human T-ALL cells^{107, 137}.

Many other chromatin regulatory complexes have been found to influence Notch activity *in vivo* (see **Box 2**). Although the functional data indicate that these complexes contribute to the landscape of target genes, it remains challenging to distinguish whether they do so via specific or non-specific mechanisms. Nevertheless, as Notch signalling is sensitive to changes in the activity of several chromatin-regulatory complexes, chromatin organization might be particularly significant for NICD responsiveness.

Factors that modify the stability or activity of NICD [L2 heading]

Another potentially potent way to regulate the Notch pathway is by modulating the activity or stability of NICD (**Figure 4**). Relatively little is known about these aspects of its regulation post cleavage, but NICD could be subject to post-translational modification or might interact with other proteins that modulate its nuclear levels

and/or activity. Yes-associated protein (YAP, a key effector of the Hippo tumour suppressor pathway) and Smad3 (an intracellular transducer of transforming growth factor- β signals) both augmented the activity of NICD independently of DNA binding^{138, 139}.

In some contexts, the stability of NICD is affected by its interactions with F-box and WD repeat domain-containing 7 (FBW7), the substrate-recognition component in a ubiquitin ligase complex. Cancer-associated mutations in FBW7 were identified in patients with γ -secretase insensitive T-ALL and correlated with increased levels of NICD activity¹⁴⁰. FBW7 has been shown to bind directly to NICD, promoting its polyubiquitylation and proteasomal degradation^{140, 141}, and, as this interaction is regulated by CDK8 (a nuclear serine/threonine kinase that functions as a transcriptional regulator), it was thought to terminate NICD activity at its transcriptional targets¹⁴². However, FBW7 affects many other substrates¹⁴³ and regulates the association of CDK8 with the Mediator complex¹⁴⁴, making it tricky to distinguish the significance of direct NICD regulation in many contexts. Furthermore, although FBW7 mutation affects several developmental processes linked to altered Notch1 activity in vertebrates^{145, 146}, mutations in the *D. melanogaster* homologue have not uncovered an equivalent role. Thus, the extent of direct NICD regulation by FBW7 remains to be clarified, and the existence of other E3 ligases that perform similar roles to terminate NICD activity merits further exploration.

SIRT1 and coactivator-associated arginine methyltransferase 1 (CARM1) are speculated to act as rheostats by modulating the activity of NICD. SIRT1 directly associates with NICD and attenuates Notch activity in zebrafish endothelial cells¹⁴⁷, and CARM1 methylates NICD and regulates the duration of transcriptional responses from some target enhancers¹⁴⁸. Other modifications may attenuate the transcriptional activity of NICD or affect its nuclear localization. These include hydroxylation, mediated by factor inhibiting hypoxia-inducible factor (FIH) in response to changes in oxygen levels during myogenesis¹⁴⁹; phosphorylation, by glycogen synthase kinase-3 (GSK-3)¹⁵⁰ or AKT¹⁵¹; and ubiquitylation, conferred by the HECT ubiquitin ligase WWP2^{152, 153}. Post-translational modifications targeted to positions within the six ankyrin repeats can prevent NICD from forming its tripartite activation complex with CSL and Mam¹⁵⁴⁻¹⁵⁶. If the relevant enzymes were recruited to specific targets, these modifications could nevertheless lead to differential effects on gene expression. Modifying enzymes potentially provide mechanisms of cross-talk with other pathways, as suggested by the purification of multiple kinases with NICD¹⁵⁷, and are, in some cases, specific to one paralogue. Clearly, more knowledge about when and where NICD is modified and what effects each modification has on its activity will be important for understanding its operations in a given milieu.

Network context [L1 heading]

Interpreting the context-specific effects of Notch will ultimately require that we understand the wiring of the regulatory networks in which it operates. Although this presents an enormous challenge, indications already exist as to how differences between cell types affect Notch pathway outcomes, as illustrated by the examples below.

Feedback regulation of ligand expression is one example in which the regulatory logic has profound consequences. For contexts in which classical lateral inhibition occurs, Notch activation frequently inhibits ligand expression to polarize the signalling (negative feedback). This mechanism relies on the HES family of direct Notch targets, which antagonize the activity of proneural bHLH transcription factors, which themselves promote the expression of ligands and E3 ligases (such as Neur)¹⁵⁸. Thus, Notch activation leads to decreased ligand expression, and the fates of the

cells becomes mutually exclusive⁵. For contexts in which signalling is inductive, Notch activity promotes ligand expression (positive feedback). This feed-forward positive regulation of ligand expression occurs at the signalling boundary in the *D. melanogaster* wing disc and during the formation of sensory patches in the chick ear^{24, 31, 45, 159}. In *D. melanogaster* the evidence points to a direct regulation of Ser expression by NICD–CSL^{9, 160}. By intensifying and perpetuating ligand expression, such positive feedback can sharpen the boundaries between expressing and non-expressing regions. Alternatively, under some circumstances it could lead to a shut-down if the increased ligand levels blocked signal reception through cis-inhibition. Differences in the wiring of the regulatory network also account for opposing effects on PTEN (phosphatase and tensin homologue) that occur in response to Notch activity. In T-ALL cells, Notch activity, by directly targeting HES1, inhibits the expression of PTEN, thereby promoting proliferation¹⁶¹. By contrast, in stalk cells, PTEN is itself a direct target of Notch and is up-regulated to inhibit proliferation¹⁶².

Differences in the mode of crosstalk between the Notch and epidermal growth factor receptor (EGFR)–Ras pathways lead to them functioning antagonistically in some contexts and cooperatively in others¹⁶³. Such polarized differences partly arise from the regulatory logic of the target enhancers and partly as a consequence of whether Notch regulates the expression of Ras pathway inhibitors or activators (and vice versa). For example, in the *D. melanogaster* eye, EGFR and Notch pathways cooperate to promote the development of cone cells by converging on enhancers of key differentiation genes (for example, Pax2¹⁶⁴), but antagonize one another at the onset of ommatidial development because EGFR promotes the expression of the proneural protein Atonal whereas Notch activity inhibits *atona1* expression through HES targets¹⁶⁵. Similarly, during *C. elegans* vulval development, EGFR activity first initiates Notch activity by upregulating ligand expression to stimulate the receptor on adjacent cells, then Notch activity antagonizes EGFR–Ras signalling by promoting the expression of pathway inhibitors¹⁶⁶. These examples illustrate how differences in wiring can profoundly influence the consequences of activating the two pathways. Similar differences in crosstalk are likely to underpin many of the context-specific interactions of Notch with Wnt, fibroblast growth factor, Hippo and other pathways.

Conclusions and perspectives

The ability of the Notch pathway to carry out many tasks despite the relative simplicity of its core pathway relies on the deployment of different levels of control that adapt the pathway to each context. For example, the expression patterns of Notch and its ligands and the tissue architectures can determine both the range and strength of Notch signalling, whereas the nuclear context will shape the identity of the target genes regulated and hence the transcriptional outcome. Many of the strategies, especially in the nuclear context, will be relevant for other signalling pathways that similarly induce a diversity of tasks. For example, the context-dependent modifiers of Wnt/ β -catenin signalling that contribute to its differing effects in stem cells include the cocktail of cooperating transcription factors that are present¹⁶⁷. However, other strategies are more likely to be unique to Notch. Notably, the fact that Notch ligands are transmembrane proteins constrains the range of the signal and makes the cell architecture and tissue organization particularly important features. The one-to-one interaction between ligand and receptor places more emphasis on the precise relationship between their levels, especially as they can also inhibit one another when present in the same cells. All the levels of regulation can be modulated to enable the pathway to adapt to a changing environment. However, although we can appreciate how the regulation might occur conceptually, many aspects are still poorly understood, making it hard to predict how physiological and environmental differences will influence signalling.

Recent progress has been driven by structural studies of key complexes involved in Notch signalling; the next challenge will be to find ways to view the molecules in action, to find where on/in the cell they interact and to discover the levels, stoichiometries and dynamics of the different complexes. A more quantitative picture will aid predictions about the transcriptional and physiological outcomes. Discovering how these can be modulated by environmental factors will also be important for understanding disease susceptibilities from heterozygous mutations in Notch pathway genes.

Box 1: Mechanistic features of Notch signalling

All canonical Notch ligands are transmembrane proteins (apart from some unusual relatives in *C. elegans*) that share a largely similar structure, with an extracellular domain comprised primarily of multiple EGF repeats (EGFR, see figure, which shows *D. melanogaster* Notch and its ligands, Serrate and Delta). Serrate and its' Jagged orthologues also contain a cysteine rich domain, CRD. Binding by canonical Notch ligands involves the extracellular Delta/Serrate/Lag-2 (DSL) domain and amino-terminal (NT) domain (see figure), which contact EGF repeats 11–12 within the extracellular domain of Notch^{22, 168}. As the NT domains have phospholipid-binding characteristics¹⁶⁹, interactions with the adjacent cell membranes might also be involved. Notch receptors on the cell surface are heterodimers (see figure): the two heterodimeric portions (HDN, HDC) interact and together with the cysteine-rich Lin12/Notch repeats (LNRs) form the negative regulatory region (NRR), which occludes the cleavage site for ADAM proteases⁵⁶. The key step induced by ligand-binding is the exposure of this cleavage site, which allows access by proteases. ADAM10 is likely to be the main protease responsible for cleavage under physiological conditions¹⁷⁰. Cleavage renders the remaining transmembrane–intracellular fragment a substrate for the γ -secretase complex, which catalyses intramembrane proteolysis to release the Notch intracellular domain (NICD). NICD is characterized by a RAM (RBP-J-associated module) domain and ankyrin (ANK) repeats (see figure), both of which are required for interactions with the DNA-binding protein CSL (CBF1/Suppressor of Hairless/Lag-1; also known as RBPJ)^{3, 171-173}. Near the carboxyl terminus is a PEST domain (see figure), which regulates NICD degradation. Between the ANK repeats and PEST, NICD also contains several nuclear localization signals and a region that can confer transactivation. The association of NICD with CSL forms an interface to which the amino terminus of the co-activator Mastermind (Mam) binds, locking the complex into its active conformation and promoting gene transcription^{172, 173}.

Box 2: Players in the nuclear arena

Nuclear activation complexes

The CSL/NICD/MAM complex recruits p300/CBP, which modifies chromatin at target enhancers. Notch dependant histone modifications include wide-spread increases in H3K27ac and H3K56ac and a decrease in H3K27me3^{104, 105 107}. Other components that have been associated with the co-activator complex include the demethylase JMJD3¹⁰⁷, the RNA helicase Ddx5, the long non-coding RNA (lncRNA) steroid receptor coactivator (SRA)^{174, 175}. Chromatin modifiers that enhance Notch transcriptional activity include BRG1/Brahma complexes, BRD4, Bre1/RNF40¹⁷⁶

Nuclear co-repressor complexes: CSL-binding co-repressors for which direct interactions have been mapped include MINT/SHARP (mammals¹¹⁰) KyoT2 (mammals;¹¹¹), Hairless (*Drosophila*;¹¹⁸). SMRT/SMRTR, SIR interactions have also been detected¹¹⁹. Co-repressors recruit enzyme complexes that modify chromatin at target enhancers, including class 1 histone deacetylases (HDACs) and histone demethylases Kdm5/Lid and LSD1^{117, 119, 157, 177, 178}. Histone chaperone complexes containing CAF1, NAP1, Asf1 are also implicated in CSL dependant repression of target enhancers¹¹⁹. In some cases, co-repressor-recruitment of modifying enzymes relies on intermediaries, including Groucho/TLE, CtBP¹⁷⁹.

Additional examples of Co-operating TFs

TEAD4: a DNA binding protein that is regulated by the Hippo pathway, TEAD4 co-binds the enhancer of Cdx2 with Notch/CSL in early mouse embryos¹⁸⁰.

SoxF: Combinatorial regulation by SoxF TFs is necessary for Dll4 expression during arterial specification and co-binding is needed to give full enhancer activity ¹⁸¹.

TCF: TCF sites are present in enhancers collaboratively regulated by Notch1 and Wnt activity in intestinal crypts ¹⁸².

ETS: Ets1 binding motif and ETS1 occupancy was enriched near NICD/CSL bound regions in T-ALL cells ¹⁷⁶

Figure Legends:

Figure 1 | Ligand binding leads to exposure of the cleavage site in Notch

(a) Summary of core pathway: when canonical Notch ligands (green) bind to Notch receptors (purple; orange indicates EGF repeats 11-12, pink indicates the Negative Regulatory Region, NRR) on the adjacent cell surface they elicit two proteolytic cleavage events, the first by ADAM10 and the second by γ -secretase, that release the Notch intracellular domain (NICD). In the nucleus NICD interacts with the DNA-binding protein CSL (CBF1/Suppressor of Hairless/Lag-1; also known as RBPJ)^{3, 171-173} and the co-activator Mastermind (Mam) to promote gene transcription^{172, 173}.

(b) Schematic based on the crystal structure^{56, 57} illustrating how the Notch-NRR occludes the cleavage site for ADAM proteases until it is exposed by forces generated through ligand-binding. Left panel: the NRR comprises 3 Lin12/Notch repeats (LNR, pink) and the heterodimerization domain (HD; dark blue and light blue), which surround the recognition site making it inaccessible to ADAM proteases. Also represented are 3 EGF-repeats from the Notch ECD, the RAM (RBP-J-associated module) and first ankyrin repeat (ANK1) domain in NICD. Right panel: Ligand binding exerts a force on the receptor (right panel), which displaces the LNRs, exposing the site for cleavage by ADAM 10. This cleavage renders the residual transmembrane Notch fragment a substrate for proteolysis by the γ -secretase complex, to release NICD (see a)

Figure 2 | The consequences of *cis* inhibition and Fringe expression on Notch signalling

(a) Relative levels of ligands (green) and Notch receptors (purple) determine whether cells send or receive signals because *cis* interactions between ligands and receptors present on the same cells are inhibitory (light shading). Receiver cell, left, expresses more Notch than Delta ($N > DL$); some Notch molecules are *cis*-inhibited by DL but sufficient Notch remains available to interact with ligands from neighbouring cells, making the cell capable of receiving signals. Sending cell, right, expresses more DL than Notch ($DL > N$); all Notch molecules are *cis*-inhibited by DL and sufficient DL remains available to interact with receptors on neighbouring cells, making the cell capable of sending a signal.

(b) Fringe proteins (yellow) glycosylate the Notch extracellular domain and modulate both *cis* and *trans* interactions with ligands. In the absence of Fng (left) Jag preferentially *cis*-inhibits Notch, so that none of the ligand is available for signalling. Any uninhibited Notch is competent to interact with either DL or Jag ligands from neighbouring cells. When Fng is present (right) glycosylation of Notch interferes with Jag (turquoise) *cis* and *trans* interactions, with the result that the cell can only receive signals from DL ligands (green), but can now send Jag signals.

Figure 3 | Influence of cell contacts and tissue architecture on signalling

(a) Contact-dimensions between Delta- (green) and Notch-expressing cells (purple) could alter Notch responses. Left: Large contact region, many ligand-receptor interactions occur generating high/prolonged NICD (dark purple/Response 1). Middle: Smaller contact surface, fewer ligand-receptor interactions, less NICD and different/fewer target genes activated (mid-purple/Response 2). Right: Cell contacts via filopodia have limited/transient receptor-ligand interactions, generating low/transient NICD and response (pale purple/Response 3).

(b) Notch signalling associated with different cell architectures. Angiogenic branching and extension of blood vessels involves the formation of dynamic contacts between tip cells (green, Dll4-producing) and adjacent stalk cells (mauve, Notch expressing), during which a presumptive stalk cell might receive high levels of Notch for a brief period before losing contact. When Notch signalling is perturbed, excess tip cells

generate a densely branched, compacted network. Selection of sensory organ precursor (SOP) cells involves direct contact between signal-sending cells (green) and signal-receiving cells (purple), but might also require filipodia to transduce the signal across a longer range. Perturbations to Notch give tufts of bristles, as all neighbouring cells become SOPs. Disruptions to filopodia result in extra SOPs and altered bristle spacing. During the formation of pigmented stripes in zebrafish, melanophores are restricted to stripes via signals conveyed by long cellular protrusions extending from xanthophores (green). Perturbations to Notch result in expansion of melanophores into the inter-stripe region.

Figure 4 | Regulation of the nuclear context

Steps involved in NICD-mediated activation at target enhancers, indicating the types of regulation occurring at each; left side, Inhibitory mechanisms (grey); right side, positive mechanisms (turquoise, purple). In the absence of cooperating transcription factors, CSL motifs (orange) are obscured by nucleosomes; this may be brought about by the presence of specific repressors (e.g. Hamlet/Evi, grey) acting in combination with chromatin modifying/remodelling enzymes. Co-operating transcription factors: (e.g. Runx, cyan) promote chromatin remodelling to expose CSL motifs. In many contexts CSL binding remains transient/unstable under these conditions, possibly due to the influence of its co-repressors (dark blue). Presence of NICD: a tertiary complex containing CSL, NICD and Mam is formed and resides at the CSL binding-sites, where it recruits co-activators and stimulates transcription. Levels/extent/duration of transcription will depend on the levels/perdurance of NICD, which is regulated by post-translational modifications (PTMs). Ubiquitination directed by E3 ligases, via intermediaries such as Fbw7, promote degradation to terminate NICD activity. Note that some target genes may already be transcribed in the absence of Notch, so that Notch binding will augment rather than initiate expression.

Figure 5 | Transitions in Notch-responsive programmes

Requirement for different co-operating factors and repressors to bring about changes in the Notch response at different developmental/physiological transitions. Top: In a nucleus that contains no co-operating TFS, CSL motifs (brown) are inaccessible. Middle: when an enabling co-operating TF is expressed (Cooperating TF1, cyan), it binds to target sites in enhancers making them competent to respond to CSL-NICD complexes, yielding a specific Notch response 1. When the cell undergoes a subsequent transition (e.g. following cell division in the olfactory lineage) different TFs are expressed, some of which inhibit (grey hexagon) the response 1 class of genes by conferring a non-permissive chromatin context and others (co-operating TF2, green) opening up a new cohort of enhancers to enable a different Notch response (Notch response 2). In this way, nuclei can transition from one response to another depending on time, other signals.

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Glossary terms:

EGF-LIKE REPEATS: Protein domains, commonly found in the extracellular domain of membrane-bound proteins, that are related to a sequence in EGF and include cysteine residues involved in disulphide bonds. EGF-like domains frequently occur in numerous tandem copies in proteins as in Notch (EGF-like_dom: IPR000742)

PARALOGUES Sequences, or genes, that have originated from a common ancestral sequence, or gene, by a duplication event.

LATERAL INHIBITION The process by which a cell with a particular fate interacts with its immediate neighbours to prevent them from adopting the same fate.

GROWTH ORGANIZER Group of cells that produces signals necessary to promote growth of a tissue.

SOMITOGENESIS The process by which somites, blocks of mesoderm that give rise to axial muscles, bones and dermis in vertebrates, are formed.

PLACODE Ectodermal thickening from which a sense organ or ganglion develops.

ESCRT (endosomal sorting complex required for transport). The multiprotein ESCRT machinery (ESCRT-I, -II and -III) promotes inward vesiculation at the limiting membrane of the sorting endosome, and selects cargo proteins for delivery to the intraluminal vesicles of multivesicular bodies.

SOP CELL Sensory organ precursor cell that gives rise to all cells in a *Drosophila* sensory organ.

ADHERENS JUNCTIONS Actin-filament-associated, epithelial cell–cell junctions that have classical cadherins as their core component.

END FEET The name given to the apical membrane surface as a consequence of cortical neuroepithelial progenitors becoming very tall and thin over the course of development.

ANGIOGENIC FRONT Region at the leading edge of a vascular network where tip cells are located to initiate further growth and branching of the network

FILOPODIA Thin cellular processes containing long, unbranched, parallel bundles of actin filaments.

NOTUM structures that are part of the back of an animal, in insects the back of the thorax.

XANTHOPHORES Yellow chromatophores, pigment-containing and light-reflecting cells, of a fish, amphibian, or reptile.

MELANOPHORES Melanin containing cells, of a fish, amphibian, or reptile that appear black or dark-brown because of melanin's light absorbing qualities.

ENHANCER A DNA segment that increases transcription of a linked promoter if placed in either orientation, upstream or downstream.

BASIC HELIX–LOOP–HELIX (bHLH) GENES Genes that encode proteins that contain a basic domain adjacent to two α -helices separated by a loop (the HLH domain), which binds DNA in a sequence-specific manner.

HES gene family A family of genes related to Hairy and Enhancer of split that encode nuclear proteins that suppress transcription.

GATA-TYPE TRANSCRIPTION FACTORS A family of transcription factors that contain a zinc-finger motif that was first identified in the vertebrate GATA1 protein. These transcription factors bind the consensus sequence GATA in the regulatory regions of genes.

ZINC FINGER A motif in proteins that contains conserved cysteine residues. The sulphhydryl groups of the cysteines coordinate a Zn^{2+} ion.

BEN-SOLO PROTEINS Proteins containing only a BEN domain, a sequence-specific DNA-binding domain that has been identified in some transcription repressors.

NEUROGENIC TRANSITION Change in the competence of neural precursor cells that enable them to generate different types of neural or glial progeny

F-BOX PROTEIN A component of the machinery for the ubiquitin-dependent degradation of proteins. F-box proteins recognize specific substrates and, with the help of other subunits of an E3 ubiquitin ligase complex, deliver them to the E2 ubiquitin-conjugating enzyme.

WD40 PROTEIN A 40-amino-acid-long protein motif that contains a WD dipeptide at its carboxy terminus. This domain is found in many functionally diverse proteins and mediates protein–protein interactions.

MEDIATOR COMPLEX A multiprotein *complex* that is required for gene transcription by RNA polymerase II.

HECT Stands for homologous to E6-AP carboxyl terminus. The HECT domain is a ~350-amino-acid domain, highly conserved among a family of E3 enzymes.

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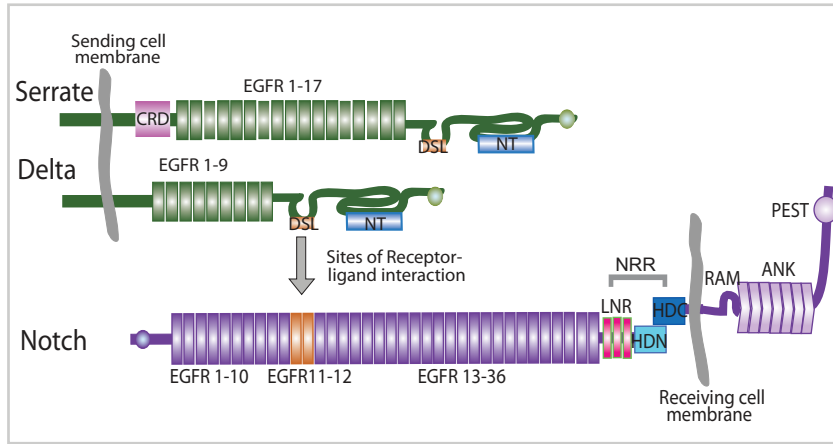
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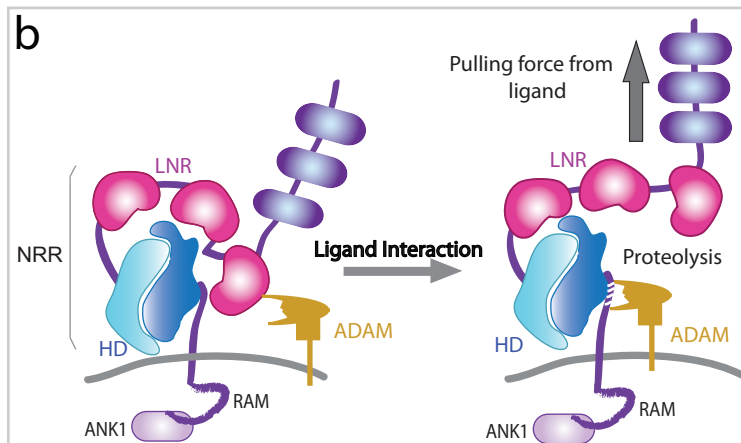
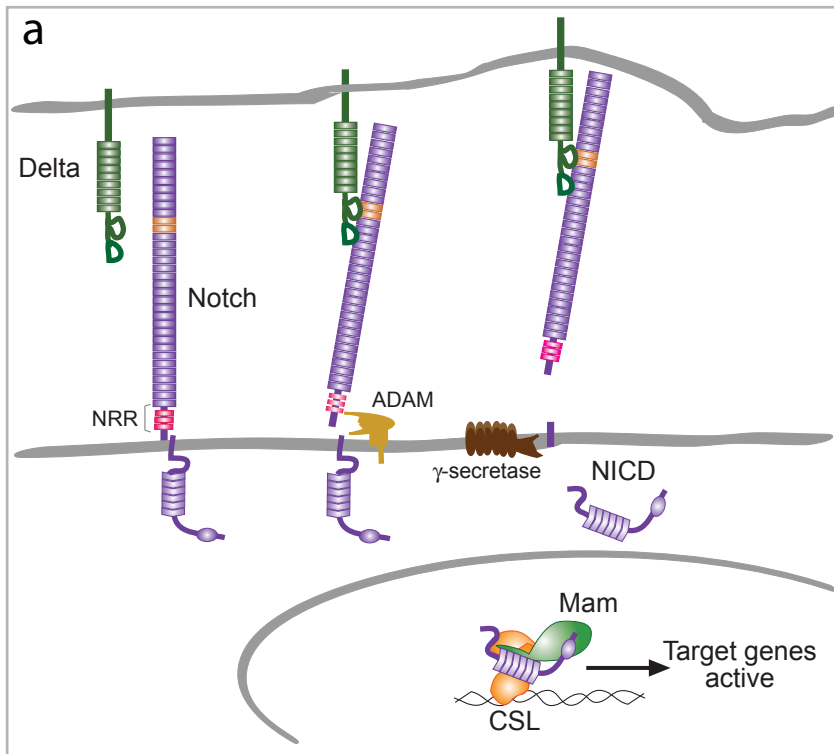
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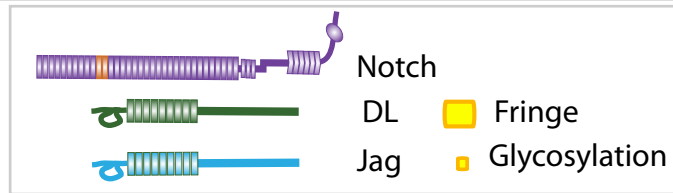
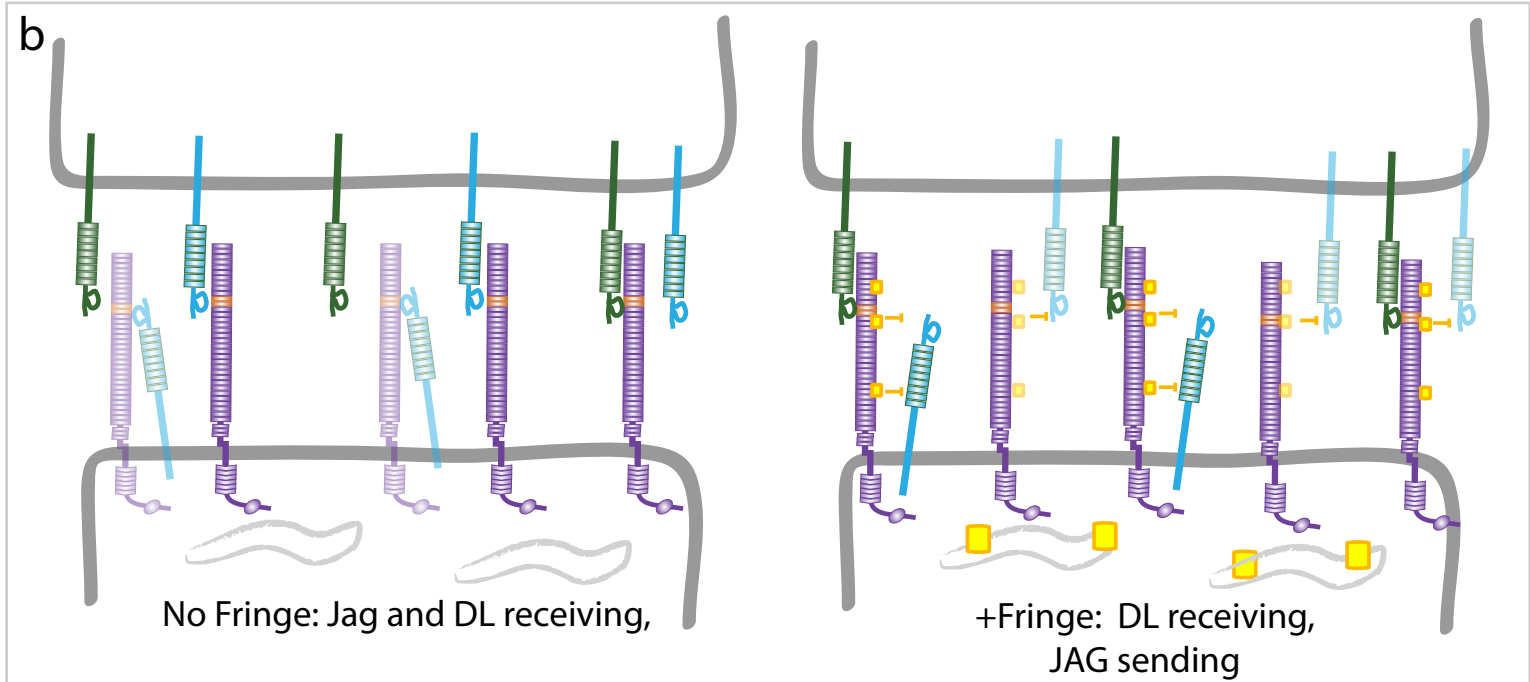
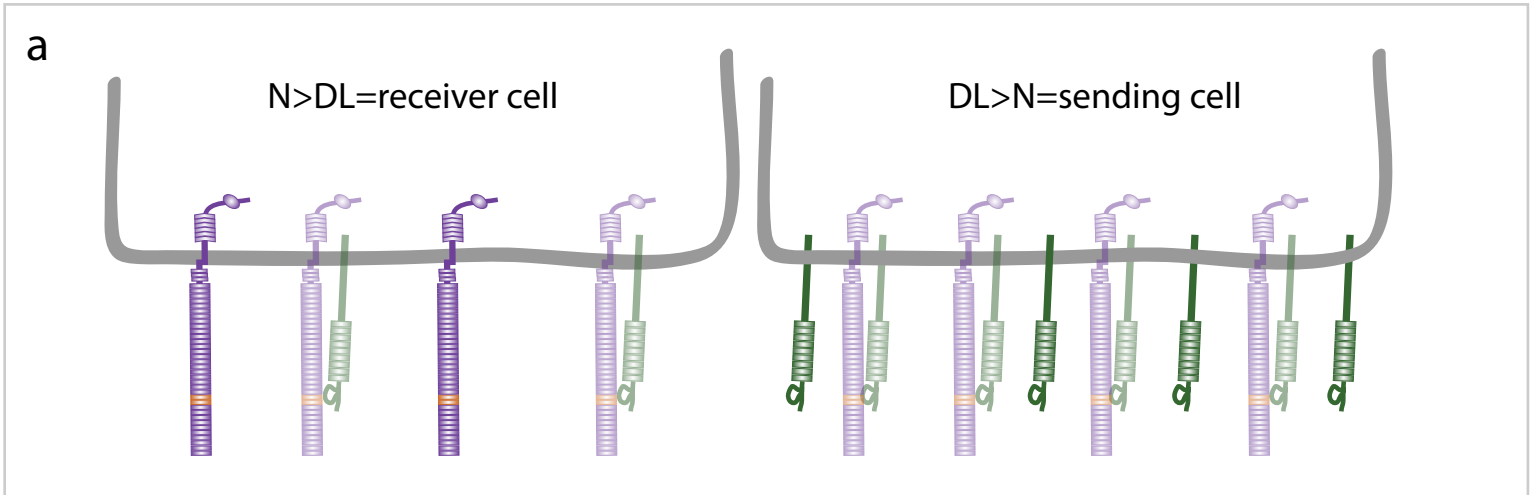
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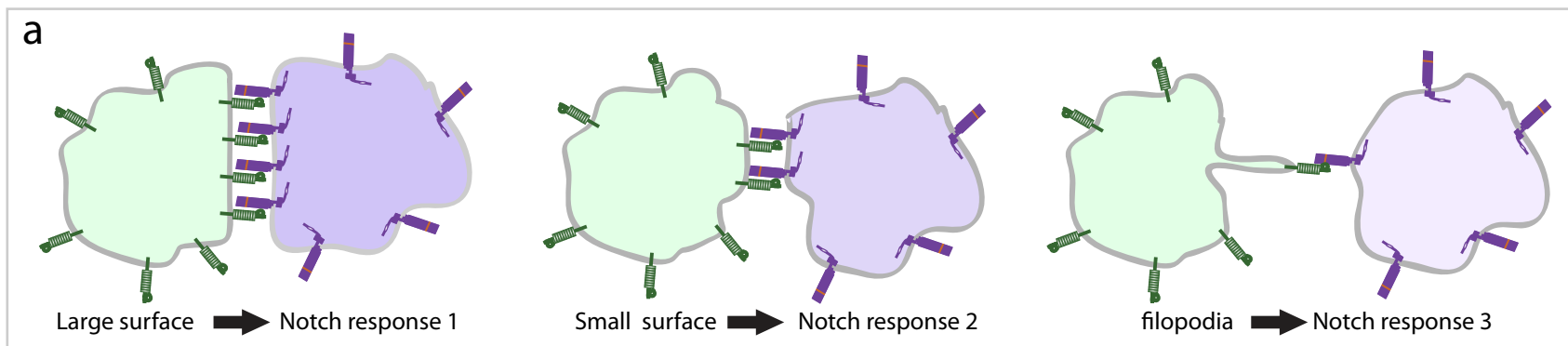
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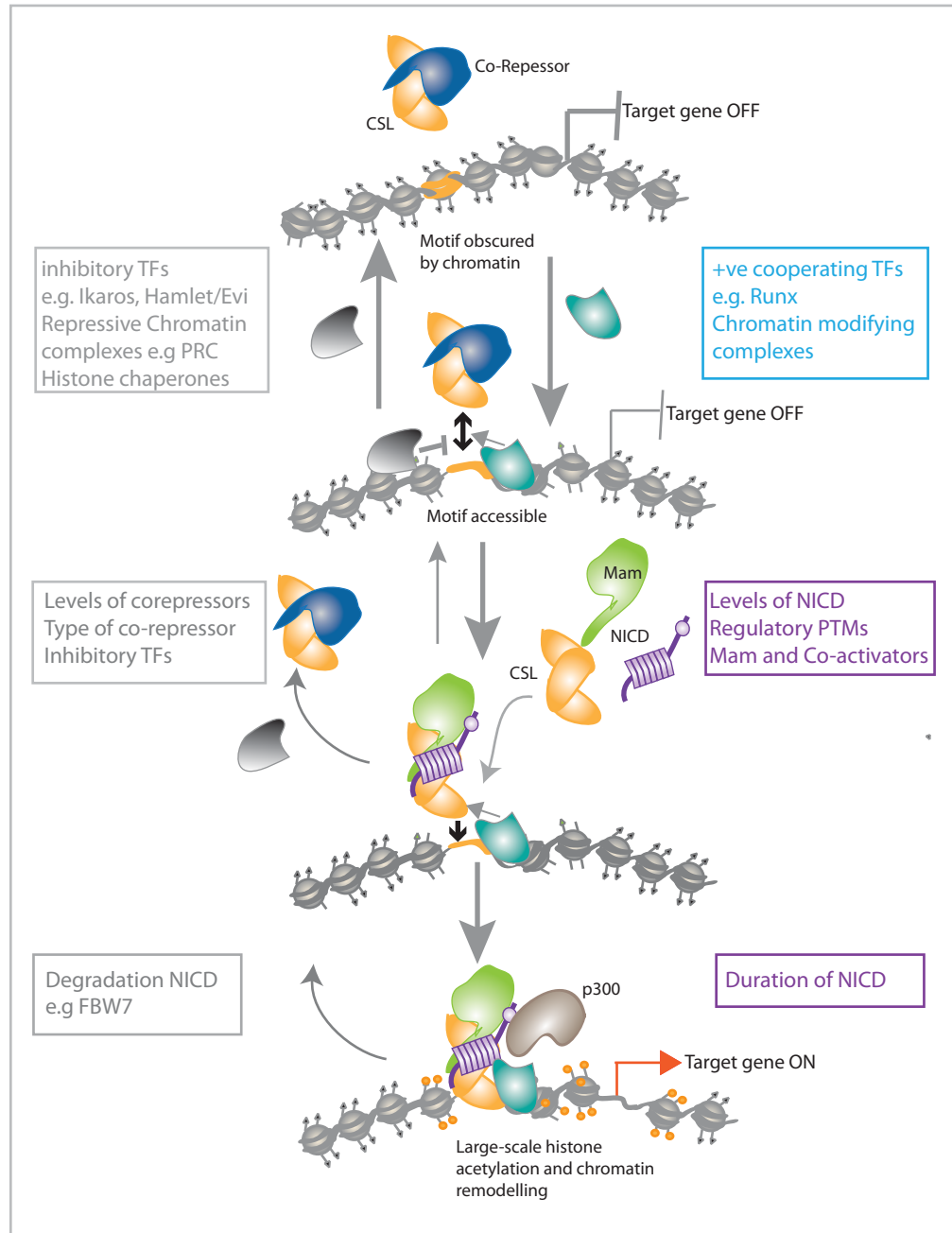






b

	Wild-type	Notch defective	
Networks of arteries			
Spaced Sensory Organs			
Pigmented Melanophore stripes			



Bray_Figure 5

