Dynamic Ligand Discrimination in the Notch Signaling Pathway

Graphical Abstract

Notch pathway can discriminate between distinct ligands
- Ligand identity controls Notch1 activity dynamics
- Notch1 dynamics control distinct target gene programs
  - Ligands can induce different cell fates

Authors
Nagarajan Nandagopal, Leah A. Santat, Lauren LeBon, David Sprinzak, Marianne E. Bronner, Michael B. Elowitz

Correspondence
melowitz@caltech.edu

In Brief
Notch ligands activate distinct targets through the same Notch receptor by triggering pulsatile or sustained activation dynamics.

Highlights
- Dll1 and Dll4 can activate distinct targets through the same Notch receptor
- Ligand identity is encoded in pulsatile or sustained Notch activation dynamics
- Dynamic encoding involves ligand-receptor clustering
- Dll1 and Dll4 induce opposite cell fates during embryonic myogenesis
Dynamic Ligand Discrimination in the Notch Signaling Pathway

Nagarajan Nandagopal, Leah A. Santat, Lauren LeBon, David Sprinzak, Marianne E. Bronner, and Michael B. Elowitz

INTRODUCTION

In metazoans, the Notch signaling pathway enables communication between neighboring cells. It plays critical roles in the development and maintenance of most tissues (Bray, 2016; Guruharsha et al., 2012), and its dysregulation has been implicated in a variety of diseases, making it an important therapeutic target (Andersson and Lendahl, 2014). In mammals, Notch signaling can be activated by four different transmembrane ligands: Dll1, Dll4, Jag1, and Jag2. When these ligands interact with Notch receptors expressed on the surface of neighboring receptor cells, they induce cleavage of the receptor. This releases the Notch intracellular domain (NICD), which translocates to the nucleus and, in complex with CSL/RBPjκ, activates Notch target genes (Figure 1A). In principle, different ligands could be used to activate distinct target programs, and thus could constitute distinct “communication channels.”

Indeed, ligand-specific effects of Notch signaling have been observed in multiple contexts and occur even with close paralogs like Dll1 and Dll4 (Figure 1A). For example, Dll4 is unable to replace Dll1 function in many tissues, leading to embryonic lethality in mice when knocked into the Dll1 locus (Preuß et al., 2015). Dll1 and Dll4 also have opposing effects on muscle differentiation: Dll1 expressed in the neural crest induces differentiation of muscle progenitors in somites (Rios et al., 2011), while Dll4 expressed in endothelial cells can revert this fate in committed skeletal myoblasts, diverting them to form pericytes instead (Cappelliari et al., 2013). Puzzlingly, although Dll1 and Dll4 can behave differently under certain conditions, they appear to function interchangeably in others. For example, when overexpressed, both ligands promote T cell differentiation of primary hematopoietic stem cells in culture, but appear to do so with different efficiencies (Mohtashami et al., 2010).

How could different ligands induce different responses in signal-receiving cells? Due to the proteolytic mechanism by which all ligands activate Notch, information regarding ligand identity must be represented in the levels or dynamics of NICD or in signal-receiving cells? Due to the proteolytic mechanism by which all ligands activate Notch, information regarding ligand identity must be represented in the levels or dynamics of NICD in signal-receiving cells. In fact, the Dll1 and Dll4 extracellular domains differ by more than 10-fold in their affinity for Notch (Andrews et al., 2013), which could lead to differences in their signaling strength (NICD levels). However, several aspects of the Notch pathway also suggest a potential sensitivity to dynamics. Cleaved NICD has a short half-life, enabling its concentration to respond rapidly to changes in Notch activation (Fryer et al., 2004; Housden et al., 2013; Ilagan et al., 2011). Similarly, the canonical Notch target genes Hes1 and Hes5 have short mRNA and protein half-lives and their levels oscillate in many contexts (Kobayashi and Kageyama, 2014). While dynamics has been shown to play critical roles in other signaling contexts (Purvis and Lahav, 2013), it has not been systematically investigated in the Notch pathway.

Here, by quantitatively analyzing Notch1 activation in individual cells, we show that Dll1 and Dll4 generate distinct patterns...
of direct target gene expression by encoding ligand identity in Notch activation dynamics. Specifically, Dll1 induces pulses of Notch activation, while Dll4 induces sustained activity. These dynamics are in turn decoded to control relative levels of Hes1 and Hey1/L target gene expression. Notch activity in receiving cells is thus inherently multi-dimensional, possessing both an activation type (pulsatile or sustained) and an activation level. This ability to respond in a ligand-specific fashion enables signal sending cells to use different ligands to activate distinct Notch target programs in receiving cells, effectively expanding the number of communication channels in the Notch pathway.

RESULTS

Dll1 and Dll4 Signal through Notch1 with Different Dynamics

In order to directly compare Notch1 signaling by Dll1 and Dll4 at the single cell level, we constructed “sender” and “receiver” cell...
lines in CHO-K1 cells (Figure 1B). Sender cells expressed either Dll1 or Dll4 along with a co-translational H2B-mCherry readout, under control of a 4epi-tetracycline (4epi-Tc)-induced promoter (LeBon et al., 2014; Sprinzak et al., 2010). We engineered receiver cells to express chimeric Notch1 receptors whose intracellular domain is replaced by the transcription factor Gal4 (Le-courtois and Schweiguth, 1998; Sprinzak et al., 2010; Struhl and Adachi, 1998), along with an H2B-3xCitrine fluorescent protein reporter that can be activated by Gal4 (Figure 1B). This “diverted” reporter system enables readout of Notch activity without activation of endogenous Notch targets, avoiding potential complications due to downstream feedback interactions.

To compare dynamics of signaling by Dll1 and Dll4, we used time-lapse microscopy of sender-receiver co-cultures. In these experiments, a minority of receiver cells were co-cultured with an excess of either Dll1 or Dll4 sender cells so that each receiver cell was in continuous contact with one or more sender cells (Figure 1C). Increases in the level of stable H2B-3xCitrine fluorescence in receiver cells reflect the activity of Gal4 released from activated receptors. Specifically, the rate of increase in total fluorescence (dCitrine/dt, “promoter activity”) is controlled by the concentration of released Gal4 (Figures S1A and S1B). We therefore estimated the Notch activity from the time derivative of each fluorescent protein trace, computed by calculating the change in total nuclear fluorescence from one time point to the next (30 min apart, see STAR Methods).

Under these experimental conditions, Dll4-expressing sender cells activated receivers in a sustained fashion. After plating, individual receiver cells activated Citrine production and continued to actively produce Citrine for the duration of the experiment (Figures 1C and 1D; Movie S1). The sustained nature of Dll4 signaling was also reflected in the average response of these cells (Figure S1C). To extract stereotyped features of the average response shape, independent of cell-cell variation in signaling amplitude and timing of activation, we normalized each trace by its maximal level and temporally aligned the resulting traces at the point of activation (Figure S1D; STAR Methods). This procedure sharpened the sustained nature of response to Dll4 (Figure 1D).

In contrast, in co-culture with Dll1-expressing senders the same receiver cells activated in discrete, transient pulses (Figures 1C and 1E; Movie S2). In each pulse, the rate of Citrine production increased transiently and then returned to baseline, displaying a characteristic shape (Figures 1E, S1C, and S1D). Pulses occurred in an unsynchronized fashion, initiating at different times in different receiver cells and could occur throughout the experiment (Figure S1E). Most cells under these conditions displayed a single pulse during the experiment (60% of traces), while two pulses could be detected in other traces (35%) (Figure S1F; Movie S3). Dll1 pulses displayed peak amplitudes comparable to the amplitude observed during the corresponding period of Dll4 signaling (Figure 1F). These results indicate that Dll1 activates Notch1 through stochastic stereotyped pulses.

In order to better understand pulsatile Dll1 signaling dynamics, we sought to estimate the duration of the underlying pulse of Notch activation, accounting for the half-lives of Gal4 protein and H2B-Citrine mRNA, which extend the duration of the observed reporter pulse. We used a mathematical model of reporter activation (STAR Methods) to analyze the decay of Citrine production rate following inhibition of Notch signaling (Figures S1G and S1H) and computed values for the half-lives of Gal4 protein (~4 hr, 95% confidence interval [CI] [3.8 hr, 4 hr]) and H2B-3xCitrine mRNA (~3.4 hr, 95% CI [3.4 hr, 3.5 hr]). Together with the measured duration (~12 hr full-width at half-maximum [FWHM]) and rise-time (~6 hr, “t_rise”) of the Dll1-induced reporter activity pulses (Figure S1I), this enabled us to estimate an upper bound of ~1 hr on the duration of the underlying signaling events (Figure S1J). Simulations showed that pulses briefer than this would produce indistinguishable reporter dynamics (Figure S1K).

As discussed more below, these brief pulses likely represent events in which multiple Notch receptors are activated (cleaved) simultaneously.

We next asked whether the apparently sustained Dll4 signaling could be explained as a series of Dll1-like pulses, occurring at an elevated rate (Figure S2A). We computationally generated pulse trains composed of pulses with the same shape and amplitude distribution observed for Dll1 pulses (Figure S2B; STAR Methods). We varied both the regularity of the pulses, using dynamic models ranging from periodic to Poisson distributed, as well as the pulse frequency (or mean interval between pulses) within each model, and analyzed the amplitude and temporal (“intra-trace”) variability of the simulated pulse trains (Figures S2C and S2D). Higher pulse frequencies lead to greater pulse overlap, increasing signaling amplitude, while reducing the temporal variability of signaling (Figure S2E). Critically, tuning pulse frequency low enough to match the observed mean Dll4 signaling amplitude generated significantly greater temporal variability than observed experimentally (Figures S2E, inset, and S2F), suggesting that the observed sustained Dll4 signaling cannot be explained as a series of Dll1-like pulses. Furthermore, the difference in experimentally observed Dll1 and Dll4 dynamics was preserved even when the time resolution of the reporter was improved from 6–12 hr (t_rise=FWHM, Figure S1G) to 2.5–6 hr by destabilizing the Citrine mRNA (Figures S2G–S2I). Taken together, these data and analysis strongly suggest that Dll1 and Dll4 activate Notch1 with distinct dynamics, Dll1 through brief pulses, and Dll4 in a sustained fashion. We note, however, that this does not rule out the possibility that Dll4 signaling originates from a series of smaller pulses (in the extreme limit, individual ligand-receptor activation events can be thought of as small, discrete “pulses”).

Dll1 Levels Modulate Pulse Frequency, while Dll4 Levels Modulate Signaling Amplitude

We next asked how the expression level of each ligand in the sender cell modulated signaling dynamics. To isolate signaling events produced by individual sender cells, we reversed the conditions of the assay, co-culturing an excess of receiver cells with a minority of sender cells (STAR Methods). We analyzed Dll1 senders across a >10-fold range of Dll1 expression levels (Figure 2A, bottom panels; Movie S4), which maintained the same stereotyped shape and duration (Figure 2B, right panels) and showed a 1.6-fold increase in amplitude (Figure 2C, right panels). At the same time, we observed a stronger increase in
the number of activated receiver cells with increasing Dll1 expression, reflecting an increase in pulse frequency (Figure 2D).

Together, these results indicate that Dll1 expression levels modulate signaling predominantly through the frequency of stereotyped signaling pulses (Figure 2E, left panel).

Unlike Dll1, Dll4 showed sustained activation in the excess receiver assay across all levels of Dll4 expression analyzed (Figures 2A, 2B, and S2K; Movie S3). We observed a systematic increase in peak (Figure 2C, left panels) and median (Figure S2L) signaling amplitude with increasing Dll4 expression level over a 10-fold range (Figure S2K). Together, these results indicate that Dll1 and Dll4 produce qualitatively different signaling dynamics across a broad range of expression and signaling levels and modulate those dynamics in distinct ways, with Dll1 mainly...
controlling the frequency of stereotyped pulses andDll4 controlling the amplitude of sustained signaling (Figure 2E).

**Pulsatile and Continuous Notch Signals Can Elicit Distinct Transcriptional Responses**

We next asked whether the different dynamics produced by Dll1 and Dll4 activation could regulate distinct sets of target genes and thereby allow cells to discriminate between the ligands. To directly test the effect of NICD dynamics on target gene expression, we took advantage of the fact that truncated Notch1 receptors lacking most of their extracellular domain (N1ΔECD) are constitutively active, but can be inhibited by DAPT (Fortini et al., 1993; Kopan et al., 1996) (Figure 3A). Cells expressing N1ΔECD can therefore be activated for different durations and to varying levels by controlling DAPT concentration in the media for corresponding time intervals (STAR Methods).

We stably expressed N1ΔECD in C2C12 cells, where the binding of the NICD-CSL complex to target gene promoters has been previously characterized using chromatin immunoprecipitation sequencing (ChIP-seq) (Castel et al., 2013). Using RNA sequencing (RNA-seq) (STAR Methods), we identified genes that were upregulated at early time points (1 hr or 6 hr) following Notch activation by DAPT removal (Figures S3A and S3B; Table S1). We focused specifically on putative direct Notch targets previously shown to bind the CSL-NICD complex in this cell line (Castel et al., 2013). Other genes that were activated were not considered for further analysis because they are not known Notch targets; several of these genes have been shown to be induced by growth factor signaling, suggesting that they could have induced by media change during DAPT removal (Allan et al., 2001; Gururajan et al., 2008; Kesarwani et al., 2017).

Interestingly, even direct Notch target genes responded to activation of the pathway at different times (Figure 3B). Hes1, but not the other target genes, was rapidly activated, showing strong (~10-fold) upregulation by 1 hr (Figures 3B and S3A; Table S1). Other Notch targets such as Hey1, HeyL, Jag1, and Nrarp responded later, showing little change at 1 hr, but strong upregulation by 6 hr (Figures 3B and S3B; Table S1). In order to follow the early and later phases of response in finer detail, we carried out a real-time qPCR time course measurement of Hes1, Hey1, and HeyL mRNA levels following DAPT removal at t = 0 hr. (D) Duration dependence of Hes1 (blue) and Hey1 (orange) response to DAPT removal for 5 min, 15 min, or 30 min followed by replenishment (“Pulse”), or no replenishment until the 1 hr or 4 hr measurement (“Sustained”). Error bars represent SEM calculated from duplicate experiments (n = 2).

These results suggested the possibility that brief (<1 hr) pulses of Notch activation could selectively activate Hes1, with the other targets requiring longer durations of Notch signaling. To test this hypothesis, we used real-time qPCR to analyze the response of Hes1 and Hey1/L to varying durations and amplitudes of Notch activation (STAR Methods). We observed that Hes1 activation was relatively insensitive to the duration of Notch activation and could be induced strongly by brief pulses (5–30 min) and by sustained activation (Figure 3D). On the other hand, Hey1/L levels did not significantly increase until the end of the Hes1 activation pulse, at 2 hr, after which they continued to rise until reaching a steady state around 4 hr.

In order to isolate the effects of signaling duration from those of signal amplitude, we compared Hey1/L expression at the same instantaneous NICD concentrations but after different durations of NICD exposure (Figures S3D–S3G). Specifically, we
compared a brief pulse of NICD generated by total DAPT removal for 15 min, with a longer (3 hr) duration of NICD activity generated by partial removal of DAPT to 0.3 μM. These two perturbations produce the same final concentration of NICD but differ in the duration of NICD activity (Figures S3D and S3E). If NICD concentration alone controlled Hey1/L expression, then the two conditions should produce similar rates of Hey1/L synthesis (Figure S3F, top). By contrast, a requirement for sustained NICD activity would lead to a greater rate of Hey1/L expression in the prolonged case (Figure S3F, bottom). For each condition, we measured the increase of Hey1/L levels in a 30-min window in order to estimate new Hey1/L expression at the corresponding time-point (Figure S3F). We observed increased Hey1/L expression only at the 3 hr time point, indicating that an extended period of activity is required for efficient activation (Figure S3G). Higher NICD concentrations were not able to overcome the requirement for extended activation, as a 30-min pulse of total DAPT withdrawal, which produced higher NICD concentrations, did not increase Hey1/L expression (Figures S3D and S3G). NICD concentration did, however, affect the maximum induction levels of the Hes/Hey genes under sustained activation (Figure S3H).

Finally, we note that the weakness of the Hey1/L response to brief activation pulses was not due to insufficient NICD, as the Notch1ΔECD system produces more NICD from DAPT withdrawal over 30 min than observed in Notch1-expressing receiver cells co-cultured with sender cells expressing maximal levels of Dll4 (Figure S3I). Together, these results indicate that pulsatile and sustained Notch dynamics are decoded into distinct gene expression patterns, with Hes1 responding strongly even to brief pulses and Hey1 and HeyL requiring sustained activation.

**Dll1 and Dll4 Induce Different Gene Responses**

Based on the different responses of Hes1 and Hey1/L to Notch dynamics, we hypothesized that Dll1 signaling could activate Hes1 without significantly inducing the Hey genes, while Dll4 could more strongly upregulate Hey1/L, even at similar Hes1 induction levels. To test this hypothesis, we used a C2C12 cell line constitutively expressing wild-type Notch1, with its endogenous Notch2 knocked down by small interfering RNA (siRNA) (STAR Methods). We first verified that the dynamic differences between Dll1 and Dll4 activation of Notch1 are preserved in this cell line, even at similar mean levels of Notch activity (Figures S4A–S4C). We then co-cultured this cell line with CHO-K1 cells expressing Dll1, Dll4, or no ligand, and measured Hes1, Hey1, and HeyL mRNA levels by real-time qPCR (Figures S4D and S4E). We found that for the same, reproducible, 1.6-fold upregulation in mean Hes1 levels, Dll4 induced ~3- to 5-fold more Hey1/L than Dll1 did (Figure S4E). This result is consistent with the different signaling dynamics of Dll1 and Dll4 inducing different Hes/Hey expression regimes. By contrast, signaling levels (amplitudes) do influence the levels of both Hes and Hey1/L expression, but do so proportionately (Figure S3H), and therefore cannot explain the disproportionate induction of these gene sets by Dll1 and Dll4.

Further, we used a complementary imaging approach to analyze the effects of single (or few) sender cells on neighboring receivers, by using plating conditions that allowed the two cell types to contact each other predominantly along a linear interface (Figure S4F; STAR Methods). Gene expression was analyzed by hybridization chain reaction-fluorescence in situ hybridization (HCR-FISH), which provides an amplified single-cell readout of specific mRNA levels (Choi et al., 2010, 2016). In these experiments, we similarly observed that Dll4 senders, but not Dll1 senders, strongly upregulated Hey1/L in neighboring receiver cells (Figures S4G and S4H). Changes in Hes1 mRNA levels were more difficult to observe at the single cell level using this technique, due to the basal expression of Hes1 (Table S1) and the stochastic, unsynchronized nature of Dll1 pulses. Nevertheless, these results further support the conclusion that Dll1 and Dll4 activate different Hes/Hey gene expression regimes, with Dll4 producing a higher expression of Hey1/L compared to Dll1 at similar Hes1 levels.

**Dll1 and Dll4 Direct Opposite Fates In Vivo**

We next sought to test the ability of Notch receiving cells to distinguish between Dll1 and Dll4 in the in vivo context of embryonic myogenesis in chick somites. In the developing embryo, it has been shown that Dll1 expressed in migrating neural crest cells signals to Notch1 expressed in the dorsomedial lip (DML) of the neighboring somite. This interaction promotes differentiation of Pax7+ progenitor cells in the DML by upregulating the muscle regulatory factors Myf5 and MyoD1, likely via Hes1 (Rios et al., 2011) (Figure 4A). Critically, in this system, transient activation of the Notch pathway enables normal muscle differentiation, while sustained activation inhibits this process (Rios et al., 2011).

Our results thus far suggest that transient and sustained Notch activation are intrinsic properties of the Dll1 and Dll4 ligands, respectively. Therefore, we predicted that the pulsatile dynamics of Dll1 would promote myogenic fate, while the sustained dynamics produced by Dll4 would inhibit myogenesis in the same cells. To test this possibility, we electroporated either Dll1 or Dll4 into the neural crest unilaterally in stage HH 12-13 chick embryos, using the other side as a negative control (Elena de Bellard and Bronner-Fraser, 2005; Rios et al., 2011). 20 hr later, we measured expression levels of Notch targets (Hes1, Hey1, or HeyL) and MyoD1 in the adjacent somites using whole-mount HCR-FISH (Figure S5A; STAR Methods). Consistent with previously published results (Rios et al., 2011), ectopic Dll1 expression in the neural crest systematically upregulated Hes1 in the somite (Figures 4B, i and ii, and quantification in S5C) and frequently increased MyoD1 in adjacent somites (Figures 4B, iii and S5C; Table 1) or maintained its levels (Figure S5C; Table 1). As expected, ectopic Dll1 expression did not significantly alter Hey1 levels (Figures 4C and S5C). On the other hand, ectopic Dll4 expression consistently increased Hey1 (Figures 4B, iv and v, and S5C) and HeyL (Figure S5B), in addition to Hes1 (Figures 4D and S5C). Importantly, Dll4 also strongly decreased MyoD1 in the majority of neighboring somites (Figures 4B, vi and S5C; Table 1). Thus, Dll1 and Dll4 induced opposite effects on cell fate in the same Notch1-expressing somite cell population that received the signal. While a role for differences in signaling levels between the two ligands in this context cannot be directly excluded, it is striking that these responses, observed in an in vivo context, matched the differences in dynamics and target specificity observed in cell culture systems.
Ligand Intracellular Domains Influence Dynamics through Differences in Transendocytosis

To gain insight into how Dll1 and Dll4 control Notch activation dynamics, we asked whether the dynamic mode was determined by the ligand intracellular domain (ICD) or extracellular domain (ECD). We constructed two chimeric Delta ligands, Dll1 ECD-Dll4ICD and Dll4 ECD-Dll1ICD, by exchanging the ICDs of Dll1 and Dll4 (STAR Methods) and stably expressed them in sender cell lines (as in Figure 1B), obtaining cell surface levels similar to those of their wild-type counterparts (Figure S5 D; STAR Methods).

We first compared Dll4ECD-Dll1ICD with Dll4 using the excess receiver co-culture assay. Unlike Dll4, the Dll4ECD-Dll1ICD ligand generated pulsatile activation, showing that the Dll1 ICD can strongly alter the activation dynamics of the Dll4 ligand (Figure 5A). The amplitude of these pulses was ~3-fold greater than signaling amplitude generated by Dll4 at the highest expression levels analyzed here, suggesting that pulsatile Dll4ECD-Dll1ICD dynamics could not be explained by a reduction in Dll4 signaling strength. In parallel, we compared Dll1 and Dll4ECD-Dll4ICD using the excess sender co-culture assay. With this chimeric ligand, most signaling occurred in a

**Figure 4. Dll1 Expression in the Chick Neural Crest Promotes Myogenesis but Dll4 Inhibits It**

(A) Developing chick embryo (dorsal view schematic). Dll1 (blue cells in 3) is expressed in a fraction of neural crest cells (gray, see 2, 3). These cells activate Notch1-expressing Pax7+ progenitor cells in the dorsomedial lip (DML, magenta) of the somite. When activated, these progenitor cells (green, 3) upregulate Hes1 and the muscle regulatory gene MyoD1.

(B–D) Representative images showing effects of Dll1 or Dll4 electroporation into the neural crest, on Hes1, Hey1, and MyoD1 expression in the DML. White arrows indicate the somites on the electroporated side. The dotted lines indicate the DMLs of somites or the central line of the neural tube.

(B) Top: Dll1-T2A-EGFP (i, blue), electroporated into the left side of the neural tube, is expressed in the neural tube and neural crest, resulting in upregulation of Hes1 (ii, red) and MyoD1 (iii, green) in the somites on the electroporated (left) side compared to the right side, which serves as negative control. Bottom: When Dll4-T2A-EGFP (iv, blue) is electroporated, Hey1 (v, red) is upregulated on the electroporated side, and MyoD1 (vi, green) expression is decreased.

(C) Dll1-T2A-EGFP (blue, left) electroporation does not affect expression of Hey1 (red, right) in adjacent somites.

(D) Dll4-T2A-EGFP (blue, left) electroporation increases expression of Hes1 (red, right) in adjacent somites.

See also Table 1 and Figure S5.
sustained fashion, but at an amplitude slightly lower than the peak amplitude of Dll1 signaling (Figure 5B). This result indicates that the Dll4 ICD can convert the dynamics of Dll1 to a more sustained behavior, even at comparable mean signaling strengths. Furthermore, consistent with the idea that dynamics strongly impact target gene expression, the Dll1ECD-Dll4ICD chimeric ligand, like Dll4, produced more Hey1/L expression than Dll1 at a similar level of Hes1 activation (Figure S5E, bottom panel inset). Additionally, it was not possible to match Dll1-induced Hes/Hey gene expression levels by varying the expression level of the chimeric ligand (thus varying signal amplitude), suggesting that this ligand produces a qualitatively distinct Hes/Hey gene expression response compared to Dll1 (Figure S5E). Together, these results indicate that the ligand ICD plays an important role in determining dynamic signaling mode of the ligand (pulsatile or sustained) and downstream gene expression.

How could ligand ICDs, functioning within sending cells, determine the dynamics of Notch activity in receiving cells? Based on previous work showing that the ligand ICD mediates receptor transendocytosis (Chitnis, 2006; Weinmaster and Fischer, 2011), we reasoned that the differences in dynamics between Dll1ICD and Dll4ICD ligands might reflect distinct modes of transendocytosis. We therefore compared transendocytosis in Dll1ICD and Dll4ICD sending cells, by immunostaining the Notch1ECD in sender-receiver co-cultures followed by confocal imaging (Figure S5F; STAR Methods).

We first compared Dll4 and Dll4ECD-Dll1ICD. At the interface between receivers and senders expressing either ligand, we observed regions of intense Notch1ECD staining, which co-localized with ligand staining (Figure S6A). This is consistent with previous observations of Notch ligand-receptor “clustering” at points of intercellular contacts (Bardot et al., 2005; Meloty-Kapella et al., 2012; Nichols et al., 2007). Within the sender cells, we observed two distinct types of staining for transendocytosed receptors: (1) dispersed, low-intensity staining that lacked apparent structure, and (2) discrete, high-intensity puncta that typically spanned >10 pixels (in three dimensions), possessed >100-fold higher cumulative intensities (Figures 5C and S6D), and co-localized with the endocytosis marker Rab5 (Figure S6B).

The generally pulsatile Dll1 ICD was strongly associated with the punctate endocytosis patterns in a signaling context. Dll4ECD-Dll1ICD senders adjacent to receivers showed a significant increase in the levels of punctate, but not dispersed, staining, relative to sender cells not adjacent to receivers (Figure S6C). Importantly, when compared at expression levels that produced similar Notch activity (Figure S6E), Dll4ECD-Dll1ICD sender cells exhibited more puncta per cell compared to Dll4 senders (Figure 5D, left). Wild-type Dll1 ligand also exhibited puncta (Figures S6F and S6G). Furthermore, the relative number of puncta per sender cell between Dll4ECD-Dll1ICD and Dll1 (Figure S6G, right) was similar to the ratio of their pulse rates (Figure S6H), while dispersed staining levels were similar. These results show that pulsatile signaling correlates with the appearance of punctate transendocytosis patterns. By contrast, Dll4 sender cells showed elevated levels of dispersed staining relative to sender cells not adjacent to receivers (Figure S6C) and also relative to Dll4ECD-Dll1ICD sender cells at the same mean signaling activity (Figure 5D, right), suggesting that dispersed staining reflects sustained signaling.

Together, these data suggest a model for how different ligands could generate different Notch activity dynamics in signal receiving cells through differences in transendocytosis patterns. In this model, the Dll1 ICD preferentially activates in the context of a ligand-receptor cluster (Figure 5E, top panel). A typical signaling event would involve the simultaneous activation of multiple receptors by interacting ligands within a single cluster, thereby releasing multiple NICDs at the same time to generate a pulse of signaling in the receiving cell (Figure S6I). In the sending cell, these events would produce transendocytic vesicles containing many receptor ECDs (punctate staining). By contrast, while the Dll4 ICD can also form clusters (Figures 5C and S6A), it would not require clustering for activation. It could thus predominantly activate in the context of smaller complexes, or individual ligand-receptor pairs (Figure 5E, bottom panel). This would enable Dll4 ICD to generate sustained Notch signal in the receiver cell (Figure S6I), consisting of a relatively steady “trickle” of receptor transendocytosis events, each generating a transendocytic vesicle containing a smaller number of receptor ECDs, leading to more dispersed staining in the sending cell.

**DISCUSSION**

The use of multiple channels is a fundamental aspect of engineered communication systems and could similarly provide powerful capabilities for intercellular communication. We find here that Dll1 and Dll4 can function as distinct communication channels in the Notch pathway by activating Notch1 with distinct dynamics (Figures 1 and 2) that can then be decoded into...
different patterns of Hes and Hey target gene expression (Figure 3) and cell fate (Figure 4).

While ligands differ in their mean amplitude of signaling, several lines of evidence show that downstream programs are particularly sensitive to dynamics. First, direct manipulation of signaling dynamics through the Notch1 ΔECD system (Figure 3) demonstrates that even at pulse amplitudes larger than those occurring during intercellular signaling in co-cultures, the duration of NICD pulses strongly affect target gene activation patterns, with Hey1/L activation occurring only after a delay. This time-dependence cannot be explained by a slow ramp-up in NICD levels (Figures S3D–S3G). The role of dynamics is further supported by analysis of gene expression induced by Dll1 andDll1ECD-Dll4ICD, which share the same extracellular domain, and therefore the same affinity for Notch1, but differ in their intracellular signaling dynamics (Figures 5B and S5E).

Overall, while amplitude undoubtedly plays an important role, these results are consistent with dynamic encoding and strongly argue against an exclusively amplitude-based scheme for ligand discrimination.

Dynamic encoding can be explained by a simple model based on previous observations that Notch ligands and receptors spontaneously assemble into ligand-receptor clusters at cell-cell interfaces. In the model, a Dll1-mediated pulse occurs when receptors in the cluster activate in a coordinated manner, releasing a burst of NICD (Figures 5E and S6I). The key requirement of the model is that the Dll1 ICD does not efficiently initiate transendocytosis until clusters reach a critical size, ensuring that most signaling occurs in pulses. By contrast, Dll4 may cluster, but would not require clustering for activation, and therefore be able to generate sustained signaling through activation of individual ligand-receptor complexes or smaller clusters (Figures 5E and S6I). Future studies should provide a more complete understanding of the molecular and biophysical basis of encoding by directly testing the sensitivity of transendocytosis to ligand-receptor clustering and elucidating the mechanism and dynamics of the clustering process (Seo et al., 2017).

Decoding of Notch dynamics is evident in the distinct responses of Hes and Hey Notch target genes to different durations of Notch activation (Figure 3). Known features of the
Hes/Hey system, including the short half-life and negative autoregulation of Hes1 (Hirata et al., 2002), and negative cross-regulation between Hes1 and Hey1/L could play roles in decoding (Fischer and Gessler, 2007; Heisig et al., 2012; Kobayashi and Kageyama, 2014). The homologous Drosophila Hairy/E(spl) Notch target genes also show differential responses to different durations of Notch activation (Housden et al., 2013; Krejci et al., 2009), suggesting that dynamic ligand discrimination could have existed ancestrally. A more complete and quantitative understanding of Hes/Hey interactions, including dimerization and cross-regulation, could provide insight into the decoding of Notch dynamics.

The ability of the Notch pathway to either promote or inhibit somite myogenesis, depending on the activating ligand (Figure 4), challenges the view that Notch activity promotes a single fate in any given context and shows that a seemingly minor change in ligand usage (i.e., from Dll1 to Dll4) can have dramatic consequences. Such contrasting roles for Notch ligands have also been reported in other contexts (Gama-Norton et al., 2015). The distinct effects of different ligands on cell responses could have implications for therapeutic interventions targeting Notch signaling and for directed differentiation applications that require control of Notch-dependent cell fate decisions (Andersson and Lendahl, 2014; Behar et al., 2013; Dahlberg et al., 2011; Mohtashami et al., 2010). We note that, despite their intrinsic differences, there are cases where Dll1 can partially compensate for Dll4 (Mohtashami et al., 2010). This may be because at high expression levels, Dll1 pulses from multiple sender cells effectively ‘merge’ and thereby become indistinguishable from sustained activation produced by Dll4.

The use of dynamics to transmit multiple signals through the same pathway occurs in other systems (Purvis and Lahav, 2013) including p53 (Batchelor et al., 2011; Purvis et al., 2012), NFAT (Noren et al., 2016; Yissachar et al., 2013), nuclear factor κB (NF-κB) (Cheong et al., 2008; Covert et al., 2005), growth factor signaling (Marshall, 1995; Santos et al., 2007), and yeast stress response (Hansen and O’Shea, 2016; Hao and O’Shea, 2011), suggesting it is a broadly useful strategy. Dynamic encoding could be particularly beneficial when the amplitude of signaling is difficult to control precisely, due to variability in expression or cell contact. Signaling pathways such as transforming growth factor β (TGF-β), bone morphogenetic protein (BMP), and Wnt, like Notch, also utilize multiple ligands capable of interacting with multiple receptors (Antebi et al., 2017). This raises the question of whether these different ligands can be discriminated by signal-receiving cells and, if so, whether this discrimination involves dynamics. Finally, pulsatile and sustained signaling could also provide different patterning capabilities in highly dynamic Notch-dependent patterning processes such as neurogenesis (Imayoshi and Kageyama, 2014), lateral inhibition (Barad et al., 2010; Cohen et al., 2010), and somitogenesis (Oates et al., 2012; Pourquier, 2011). Ultimately, the discovery that the Notch pathway can transmit more and different types of information than previously suspected should help to explain how it enables such an extraordinary range of outcomes, in development and physiology.
Caltech provided essential technical assistance. This work was supported by the Defense Advanced Research Projects Agency (HR0011-16-0138), by the NIH (R01 HD075335), and the NSF (EFRI 1137269). N.N. was a Howard Hughes Medical Institute International Student Research fellow.

AUTHOR CONTRIBUTIONS


DECLARATION OF INTERESTS

The authors declare no competing interests.

Received: January 13, 2017
Revised: August 10, 2017
Accepted: January 3, 2018
Published: February 1, 2018

REFERENCES


Barad, O., Rosin, D., Hornstein, E., and Barkai, N. (2010). Error minimization in the Defense Advanced Research Projects Agency (HR0011-16-0138), by the NIH (R01 HD075335), and the NSF (EFRI 1137269). N.N. was a Howard Hughes Medical Institute International Student Research fellow.

**AUTHOR CONTRIBUTIONS**


**DECLARATION OF INTERESTS**

The authors declare no competing interests.

Received: January 13, 2017
Revised: August 10, 2017
Accepted: January 3, 2018
Published: February 1, 2018

**REFERENCES**


**REFERENCES**


# STAR★METHODS

## KEY RESOURCES TABLE

<table>
<thead>
<tr>
<th>REAGENT or RESOURCE</th>
<th>SOURCE</th>
<th>IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Antibodies</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mouse anti-hNotch1 (extracellular domain)</td>
<td>BioLegend</td>
<td>Cat# 352014; RRID:AB_10899408</td>
</tr>
<tr>
<td>Rabbit anti-hN1ICD (V1744) monoclonal [D3B8]</td>
<td>Cell Signaling Technology</td>
<td>Cat# 4147; RRID:AB_2153348</td>
</tr>
<tr>
<td>Mouse anti-mGAPDH [8C5]</td>
<td>Abcam</td>
<td>Cat# ab8245; RRID:AB_2107448</td>
</tr>
<tr>
<td><strong>Chemicals, Peptides, and Recombinant Proteins</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Human Dll1 ext-IgG</td>
<td>Sprinzak et al., 2010</td>
<td>N/A</td>
</tr>
<tr>
<td>Mouse N1ext-mFc</td>
<td>R&amp;D systems</td>
<td>Cat# 5267-TK</td>
</tr>
<tr>
<td>Hamster Fibronectin</td>
<td>Oxford Biomedical Research</td>
<td>Cat# CT30</td>
</tr>
<tr>
<td>DAPT</td>
<td>Sigma Aldrich</td>
<td>Cat# D5942</td>
</tr>
<tr>
<td>4-epi tetracycline Hydrochloride</td>
<td>Sigma Aldrich</td>
<td>Cat# 37918</td>
</tr>
<tr>
<td><strong>Critical Commercial Assays</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>RNeasy mini kit for RNA extraction</td>
<td>QIAGEN</td>
<td>Cat# 74106</td>
</tr>
<tr>
<td>iScript cDNA synthesis kit</td>
<td>Bio-Rad</td>
<td>Cat# 1708890</td>
</tr>
<tr>
<td>iQ SYBR Green Supermix</td>
<td>Bio-Rad</td>
<td>Cat# 1708880</td>
</tr>
<tr>
<td>DNA HCR kit</td>
<td>Molecular Instruments</td>
<td>N/A</td>
</tr>
<tr>
<td><strong>Deposited Data</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Raw and analyzed RNaseq data</td>
<td>This paper</td>
<td>GSE72847</td>
</tr>
<tr>
<td><strong>Experimental Models: Cell Lines</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CHO-K1</td>
<td>ATCC</td>
<td>Cat# CCL-61; RRID:CVCL_0214</td>
</tr>
<tr>
<td>CHO TREx</td>
<td>Invitrogen</td>
<td>RRID:CVCL_D586</td>
</tr>
<tr>
<td>C2C12</td>
<td>ATCC</td>
<td>Cat# CRL-1772; RRID:CVCL_0188</td>
</tr>
<tr>
<td>CHO-K1-TREx + pcDNA3-CMV-hN1ECD-Gal4esn + pEV-UAS-H2B-3xCitrine</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pcDNA3-CMV-hN1ECD-Gal4esn + pEV-UAS-H2B-3xCitrine + pGK-H2B-mCherry</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pcDNA5-CMV-TO-rDll1-T2A-H2B-mCherry</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pcDNA5-CMV-TO-rDll1-T2A-H2B-mCherry + pLenti-CAG-H2B-Cerulean</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pcDNA5-CMV-TO- hDll4-T2A-H2B-mCherry</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pb-CMV-TO-hDll4-T2A-H2B-mCherry + pb-CMV-H2B-Cerulean</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pb-CMV-TO-Gal4esn-T2A-H2B-mCh</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pEF-hN1ECD-Gal4esn + UAS-H2B-3xCitrine-3Hes1UTR</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pcDNA5-CMV-TO-rDll1 ECD + Dll4ECD-^</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pcDNA5-CMV-TO-hDll4 ECD-^</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>CHO-K1-TREx + pb-CMV-TO- hN1-T2A-H2B-mCherry</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>C2C12 + pb- CMV7-hN1-myc-T2A-H2B-mCherry</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>C2C12 + pb- CMV7-hN1-FLAG</td>
<td>This paper</td>
<td>N/A</td>
</tr>
</tbody>
</table>

(Continued on next page)
<table>
<thead>
<tr>
<th>REAGENT or RESOURCE</th>
<th>SOURCE</th>
<th>IDENTIFIER</th>
</tr>
</thead>
<tbody>
<tr>
<td>C2C12 + pb/hN1ECDeGal4esn-ANK-T2A-H2B-Cer + pEV/UAS-dm-H2B-3xCitrine</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>C2C12 + pb/TO-hDll1-T2A-H2B-mCh-P2A-Hyg</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>C2C12 + pb/TO-hDll1-T2A-H2B-mCh-P2A-Hyg</td>
<td>This paper</td>
<td>N/A</td>
</tr>
<tr>
<td>Experimental Models: Organisms</td>
<td>Stage 12-13 chicken embryos</td>
<td>This paper</td>
</tr>
<tr>
<td>Recombinant DNA</td>
<td>pcDNA3/CMV-hN1ECDeGal4esn</td>
<td>Sprinzak et al., 2010</td>
</tr>
<tr>
<td></td>
<td>pEV/UAS-H2B-3xCitrine</td>
<td>LeBon et al., 2014</td>
</tr>
<tr>
<td></td>
<td>pcDNA5/CMV-TO-rDll1-T2A-H2B-mCherry</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pcDNA5/CMV-TO-hDll4-T2A-H2B-mCherry</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pb/CMV-TO-Gal4esn-T2A-H2B-mCherry</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pEV/UAS-H2B-3xCitrine-3' Hes1 UTR</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pcDNA5/CMV-TO-Dll1ECD-Dll4ICD</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pcDNA5/CMV-TO-Dll4ECD-Dll1ICD</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pcDNA5/CMV-TO-Dll1-FLAG</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pcDNA5/CMV-TO-Dll4ECD-Dll1ICD-FLAG</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pcDNA5/CMV-TO-Dll4-FLAG</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>pb/CVM-TO-hN1ECDeGal4esn-ANK-T2A-H2B-Cer + pEV/UAS-dm-H2B-3xCitrine</td>
<td>This paper</td>
</tr>
<tr>
<td></td>
<td>mouse Hes1 primer set 1 (Figure 3C) - Forward, 5'-CAACACGACACCGGACAAA -3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>mouse Hes1 primer set 1 (Figure 3C) - Reverse, AAGAATAAATGAAAGTCTAAGCCAA -3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>Mouse Hes1 primer set 2 (Figure 3D, S4, 5) – Forward, 5'-AAGAATAAATGAAAGTCTAAGCCAA -3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>Mouse Hes1 primer set 2 (Figure 3D, S4, 5) – Reverse, 5'-TTCTTGCCCTTCGCCCTTCTC -3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>mouse Hey1 primers – Forward, 5'-GCCGAAGTTTGCCGGATCATC-3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>mouse Hey1 primers – Reverse, 5'-CGCTGGATGTGCTTATCTC-3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>mouse HeyL primers – Forward, 5'-GAGCTGACGCGCTGC-3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>mouse HeyL primers – Reverse, 5'-GAGCTGACGCGCTGC-3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>mouse SdhA primers - Forward, 5'-AGTGGGCTGTTGCTTTGCTTAC-3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td></td>
<td>mouse SdhA primers - Reverse, 5'-GGATTGCTTCTGGTGG-3'</td>
<td>IDT DNA</td>
</tr>
<tr>
<td>Oligonucleotides</td>
<td>siRNA targeting mouse Notch2 5'-UGAACUUGCGAGGUGAAGGUC-3'</td>
<td>Invitrogen</td>
</tr>
<tr>
<td>Software and Algorithms</td>
<td>Segmentation, tracking, and fluorescence analysis software</td>
<td>This paper</td>
</tr>
</tbody>
</table>
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Michael Elowitz (melowitz@caltech.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Gene constructs
All constructs used in this paper were assembled using standard restriction enzyme-based cloning and/or Gibson cloning (Gibson et al., 2009). pcDNA3-hNECD-Gal4 (Figures 1, 2, and 5) has been described previously (Sprinzak et al., 2010). The H2B-3xCitrine fluorescent reporter (Figures 1, 2, and 5) was constructed by cloning 3 repeats of mCitrine in frame with H2B, downstream of a UAS promoter. The mRNA destabilized version of this reporter was constructed by fusing the 3’UTR of mouse Hes1 downstream of the STOP codon. Ligand constructs were cloned into pcDNA5 or piggyBac plasmids (System Biosciences Inc.) by fusing the complete rat Dll1 (kind gift from G.Weinmaster) or human Dll4 cDNA in frame with T2A-H2B-mCherry, downstream of a previously described inducible pCMV-TO promoter (Sprinzak et al., 2010). We note that hDll1 shows the same pulsatile behavior described here for rDll1. Chimeric ligands (Figure 5) were constructed by exchanging the intracellular domains of Dll1 (aa 561 – 714) and hDll4 (aa 551 – 685). The hN1ΔECD gene (Figure 3) was cloned from hN1 (kind gift from J. Aster) by removing residues 22-1716 and fused in frame with myc-T2A-H2B-mCherry, downstream of the CMV-TO promoter in a piggyBac construct. Constructs used for in ovo electroporation (Figure 4) were made by cloning rDll1 or hDll4 cDNA (minus stop) upstream of, and in frame with, T2A-EGFP in a pCI-CAGG plasmid.

Tissue culture and Cell lines
CHO-K1 (Hamster cells, RRID:CVCL_0214, ATCC Catalog No. CCL-61) or CHO-TREx (RRID:CVCL_D586, Invitrogen) cells and their derivatives were grown on tissue-culture grade plastic plates (Thermo Scientific) in Alpha MEM Earle’s Salts (Life Technologies), supplemented with 10% Tet System Approved FBS (ClonTech), 100 U/ml penicillin, 100 ug/ml streptomycin, 0.292 mg/ml L-glutamine (GIBCO).

C2C12 cells (Mouse cells, RRID:CVCL_0188, ATCC Catalog No. CRL-1772) were grown in DMEM (Life Technologies), supplemented with 20% Tet System Approved FBS (ClonTech), 100 U/ml penicillin, 100 ug/ml streptomycin, 0.584 mg/ml L-glutamine (GIBCO). C2C12 media was used for CHO-K1 + C2C12 co-culture assays (Figure S4). All cells were grown at 37°C in 5% CO2 in a humidified atmosphere. Cells were passaged every 2-3 days, depending on confluency, using 0.05% or 0.25% Trypsin-EDTA (Life Technologies).

Cell line engineering
All cell lines used in this paper contained stable integrations of transgenes, and were typically clonal populations. To create each stable cell line, the following steps were followed: 1) Cells were first transfected with 800-1000 ng of plasmid DNA using Lipofectamine 2000 or Lipofectamine LTX. 2) 24 h later, cells were transferred to selection media containing 600 ug/ml Geneticin, 500 ug/ml Hygromycin, 400 ug/ml Zeocin, or 10 ug/ml Blasticidin as appropriate. 3) After selection for 1-2 weeks, the resulting polyclonal populations stably expressing the transgene were allowed to recover for ~1 week. 4) Single clones were isolated through the technique of limiting dilution. 5) Single clonal populations were screened for desired behavior, usually high expression (for constitutive genes) or low background expression of the transgene and large dynamic range (for inducible genes and reporter genes). Cell lines incorporating multiple transgenes were constructed by sequential rounds of this process. For piggybac constructs, the initial transfection comprised of the target plasmid along with the construct expressing the piggybac transposase, typically in a 1:1 or 2:1 molar ratio.

Chicken embryos
Fertile chicken (Gallus gallus) eggs, purchased from commercial sources, were incubated at in a humidified 37 C incubator, and staged by the criteria of Hamburger and Hamilton (HH) (Hamburger and Hamilton, 1992). Embryos were electroporated at stage 12-13, replaced in the incubator, and dissected 20h later.

METHOD DETAILS

Co-culture assays and time-lapse microscopy
Used in Figures 1, 2, 5, S1, S2, and S5

Surface treatment
In preparation for plating of cells, glass-bottom multi-well plates (MatTek, No. 1.5 glass, 10 mm radius) were coated with 5 ug/ml Hamster Fibronectin (Oxford Biomedical Research) diluted in 1x Phosphate-Buffered Saline (PBS) for 1h at room temperature.
Cell culture
After trypsinization, sender cells (pre-induced for > 48h with 4-epiTc, Sigma) or CHO-K1 cells were mixed in suspension with similarly trypsinized receiver cells at a ratio of 100:1 or 1:100, for excess sender or excess receiver assays, respectively. A total of 8x10^4 cells (60% confluence) were plated for each experiment, with continued 4-epiTc induction when appropriate. Imaging commenced 2-4h post-plating.

Time-lapse microscopy
Movies were acquired at 20X (0.75 NA) on an Olympus IX81 inverted epi-fluorescence microscope equipped with hardware autofocus (ZDC2) and an environmental chamber maintaining cells at 37°C, 5% CO2. Automated acquisition software (METAMORPH, Molecular Devices) was used to acquire images every 30 min in multiple colors (YFP, RFP, CFP) or differential interference contrast (DIC), from multiple stage positions.

Plate-bound Dll1 assay
Used in Figures S1D and S1E
Recombinant human Dll1ext-Fc fusion proteins (kind gift from I. Bernstein) were diluted to 1 ug/ml in PBS, and the solution was used to coat the tissue-culture surface. After 1h incubation at room temperature, the solution was removed, and cells were plated for the experiment.

Image segmentation, tracking, and single-cell fluorescence calculation
Used in Figures 1, 2, 5, S1, S2, and S5
Custom MATLAB code (2013a, MathWorks) was used to segment cell nuclei in images based on constitutive CFP/RFP fluorescence or background YFP fluorescence. The segmentation procedure uses edge detection, adaptive thresholds, and the Watershed algorithm to detect nuclear edges. Nuclear segments were then matched in pairs of images corresponding to consecutive time frames, and thus tracked through the duration of the movie. Single-cell tracks were subsequently curated manually. In particular, there were periods where any given cell could not be automatically segmented (typically due to high density) but could be visually followed. In such cases, the tracks corresponding to the cell prior to and after such time frames were manually linked if fewer than ~5 frames were missing.

Fluorescence data was extracted from nuclear segments by calculating the integrated fluorescence within the segment and subtracting a background fluorescence level estimated from the local neighborhood of the segment. This fluorescence was linearly interpolated across time frames where nuclei could not be segmented automatically. Division events were detected automatically, and fluorescence traces were corrected for cell division by adding back fluorescence lost to sister cells. The resulting ‘continuized’ traces were smoothed and the difference in fluorescence between consecutive time frames was calculated. A smoothed version of this difference was used as the rate of change or promoter activity of the fluorescence.

Analysis of single-cell traces
Used in Figures 1, 2, 5, S1, S2, and S5
Alignment
For each receiver cell trace, including those of cells in control conditions (showing background fluorescence levels) an average rate of fluorescence increase (‘average slope’) was calculated by dividing the change in total fluorescence of the reporter by the duration of the trace. Traces showing activation were automatically selected for further analysis based on their average slopes surpassing a threshold value, chosen to be higher than average slopes observed in receiver cells under control conditions. Activating traces were aligned at the point of activation, defined as the time point when their promoter activity crosses an absolute threshold level, chosen based on typical promoter activities corresponding to background activity. Note that activations occurring during the first 15h of the movie were typically not considered, to eliminate transient effects produced by cell transfer to imaging conditions. The same thresholds were always used when direct comparisons were made between ligands or conditions, and we verified (by varying threshold levels) that qualitative results did not depend strongly on the choice of threshold.

For C2C12 dynamics (Figure S4) promoter activity could not be reliably used to align traces due to the low levels of reporter activity and resulting noise in the promoter activity data. These traces were instead aligned based on when the total fluorescence levels increased a threshold level.

Double-pulse alignment
In order to align traces showing two pulses in response to Dll1 (Figure S1D) at the second pulse, the following procedure was used: the first activation was determined using the usual procedure (see above). Traces were then normalized by the peak activity (‘Peak1’, 95th percentile) in the 0-7.5h window during which the first pulse is expected to reach maximum levels. Starting at 7.5h, i.e., after the peak of the first pulse, traces were re-aligned at the point when the subsequent promoter activity values cross Peak1, and re-normalized to the 90th percentile of values in the period from 7.5h (relative to the first activation point) to the end of the trace.

Normalization
When applied, the object of normalizing the response trace by its amplitude is to demonstrate its stereotyped features, such as relative rise time and duration. Un-normalized averaging would distort the shape of the response because higher-amplitude signals are also prolonged, since the timescales of the reporter are fixed by the half-lives of its components (Gal4 protein, H2B-3xCitrine mRNA)
and do not scale with amplitude. Traces were typically normalized to the 90th percentile value during the analysis time window, except in Figure S2H, where traces were normalized to the 90th percentile value occurring within 15h after activation.

**Amplitudes**

While normalized traces were used to make comparisons of the stereotyped shapes of responses (see above), absolute values of promoter activity, calculated from non-normalized promoter activity, are reported in all amplitude comparisons. Except in Figure 2C, this amplitude represents the 95th percentile of (absolute, non-normalized) promoter activity values between 0 and 7.5h (after alignment) in the traces. This time window is chosen to simultaneously estimate the promoter activity at the peak of pulses and at steady-state levels of sustained signaling. In Figure 2C, the amplitude represents the 95th percentile of promoter activity values during the 25h after activation (the period over which activities are averaged).

**Trace filtering**

In Figure 1D, traces were included in the Dll1 alignment if the median promoter activity between 20-25h fell below 50% of the peak activity (95th percentile) in the 0-7.5h period (after alignment). This criterion was designed to automatically detect single pulses in the data. In Figure 2B traces were only included in the Dll1 alignment if the normalized value at 20h fell below 0.7. This filter eliminates traces consisting of multiple pulses, especially in the high Dll1 cases. A similar filter applied to Dll4 traces reveals a small fraction of cells activated transiently, but displaying qualitatively different behavior, such as a systematic increase in duration and amplitude with increasing Dll4 levels in senders. For C2C12 experiments in Figures 3G and 3H, activating cells were identified based on an increase in total fluorescence levels above a threshold.

**Estimating Gal4 and mRNA half-lives, Related to Figure S1H**

For this model, we assume that the free Gal4 protein produced due to cleavage of N1ECD-Gal4 degrades with first-order kinetics with rate $\gamma_{Gal4}$ after inhibition of the pathway using DAPT, at time 0h.

$$\frac{dGal4}{dt} = -\gamma_{Gal4}Gal4$$

Reporter mRNA $m$ is produced through non-cooperative binding of Gal4 to the promoter, with dissociation constant $K$ and maximum rate $\beta_m$. $m$ is degraded with rate constant $\gamma_m$.

$$\frac{dm}{dt} = \beta_m \frac{Gal4}{K + Gal4} - \gamma_mm$$

The parameters $\gamma_{Gal4}$, $K$ and $\gamma_m$ were calculated by fitting the Citrine mRNA $m$ to the experimentally measured decay in Citrine fluorescence rate using the *lsqnonlin* function in MATLAB. The fit was constrained using bounds for $\gamma_{Gal4}$ and $\gamma_m$ of $\log(2)/5h$ – $\log(2)/3h$, based on Sprinzak et al. (2010) and Bintu et al. (2016). Bootstrapped 95% confidence intervals were computed from 100 iterations of fitting 30 points, chosen randomly with replacement, out of a total 50 measured time points.

**Mathematical model for estimating duration of Notch activation, Related to Figure S1J**

For this model, we assume that Gal4 is produced at a rate $\beta_{Gal4}$ for a duration $\tau_{act}$, and degrades with first-order kinetics with rate $\gamma_{Gal4}$.

$$\frac{dGal4}{dt} = \begin{cases} \beta_{Gal4} - \gamma_{Gal4}Gal4, & t \leq \tau_{act} \\ -\gamma_{Gal4}Gal4, & t > \tau_{act} \end{cases}$$

Reporter mRNA $m$ is produced through non-cooperative binding of Gal4 to the promoter, with dissociation constant $K$ and maximum rate $\beta_m$. $m$ is degraded with rate constant $\gamma_m$.

$$\frac{dm}{dt} = \beta_m \frac{Gal4}{K + Gal4} - \gamma_mm$$

For the results of Figure S1, $\beta_{Gal4} = 1$, $\beta_m = 1$, and $K = 6.6$ (also fitted in Figure S1E), and estimated mean values from Figure S1E were used for the Gal4 and mRNA degradation rates.

**Simulations of Dll1 pulse trains and analysis, Related to Figures S2A–S2F**

This model constructs pulse-trains composed of Dll1-like pulses occurring at various frequencies and regularities based on each of three underlying pulse models, and analyzes the features of the resulting simulated signaling traces.

**Pulse train construction (Figure S2B)**

For each simulation we construct 200 pulse trains. Each pulse train is constructed from a series of pulses with the average Dll1 promoter activity pulse shape (Figure S1B), scaled by an amplitude randomly sampled from the empirically measured distribution of Dll1 pulse amplitudes (from the Figure 1D dataset). The first pulse occurs at 0h, representing activation at time 0 in the aligned Dll4 traces. Subsequently, new pulses are introduced after successive time intervals chosen based on one of the underlying pulse models (see below), and the composite signal is constructed until it extends at least 10h beyond the 25h time period averaged in Figure 1D.
Feature analysis (Figure S2D)

For each trace, two features are analyzed:

1) Amplitude: The amplitude of each constructed trace is its median value over 25h.
2) Intra-trace variability: After calculation of the amplitude, each trace is normalized to its 90th percentile value. For each point \( t \) in this trace, the local temporal variability is estimated by the standard deviation of values in a 10h window starting at \( t \). The overall intra-trace variability calculated for each trace is the median of the local variability value at each point, calculated by moving a 10h time window through the trace.

For each simulation (200 constructed traces), the medians of the calculated amplitudes and intra-trace variability are tabulated, and the SEM calculated.

Pulse models (Figure S2C)

Three models are considered for the underlying pulsing process:

1) Periodic model: In this model, the interval \( \tau \) between adjacent pulses is fixed at a value \( T_{\text{period}} \), that can range from 1h to 8h. Since theDll1 pulse decay becomes apparent after 7.5h (Figure 1D), intervals greater than 8h will result in pulse trains in which the individual pulses can be clearly discerned in each trace, and the average behavior will show oscillations. Since neither individual Dll4 traces, nor the average shape display overt oscillatory features, values for \( T_{\text{period}} \) greater than 8h are not considered in the simulation.

2) Poisson model: In this model, the interval between successive pulses \( i \) and \( i+1 \), \( \tau_i \), represents the inverse of a pulse rate, \( r_i \), drawn from a Poisson distribution with parameter, \( \lambda \), ranging from 1/h-1/15h.

3) Mixed model: In these models, the interval \( \tau \) between adjacent pulses is drawn from a normal distribution with mean \( T_{\text{period}} \) (range 1h - 15h) and standard deviation \( \sigma \) (2.5h or 5h). This model therefore combines the regular pulsing inherent to the periodic model with the trace-to-trace variability of the Poisson model (thus preventing ‘constructive interference’ of pulse peaks, which would lead to apparent oscillations in the average signal shape).

Bootstrapped analysis of variability in measured Dll4 signaling trace (Figure S2F)

Finally, for direct comparison to simulation data, the Dll4 dataset of traces (200 traces in total) was subsampled 30 times (50 traces per sample) to generate a bootstrapped distribution of measured median intra-trace variability, and a corresponding median value was calculated. This bootstrapped median is compared to simulation data in Figure S2F.

Sender cell categorization in excess receiver assays

Used in Figures 2 and S2

Dll1- and Dll4-T2A-H2B-mCherry sender cells were induced with different 4epi-Tc concentrations, to access their full dynamic range of ligand expression. Following co-culture with receiver cells and timelapse analysis, individual sender cell nuclei were automatically segmented, and mCherry levels were calculated. At the same time, each receiver cell response was automatically associated with the closest sender cell. All data, across 4epi-Tc induction levels, were then pooled, and sender cells re-categorized into ‘low’, ‘medium’, or ‘high’ expression along with their associated receiver cell responses. This process of pooling and recategorization was necessary because of the broad, overlapping distributions in mCherry expression produced by 4epi-Tc treatment.

Detection of surface ligand

Used in Figure S5D

Recombinant mouse Notch\(^{\text{ext}}\)-Fc chimeric protein (R&D Systems) was used for surface-detection of ligands at a concentration of 10 ug/ml, based on a previously described protocol (LeBon et al., 2014). Sender cells were first cultured and induced with 4epiTc for 48h, then transferred from media to blocking solution (2% FBS in Phosphate Buffered Saline, PBS) for 30 min at room temperature (RT). Cells were then incubated with recombinant mouse Notch\(^{\text{ext}}\)-Fc protein in binding solution (blocking solution containing 100 ug/ml CaCl\(_2\), R&D Systems) for 45 min at RT. Following this, cells were washed 3x with binding solution, then incubated with anti-mouse secondary antibody conjugated to AlexaFluor-488 (1:1000 dilution, Life Technologies) for 30 min. Cells were then trypsinized and analyzed using flow cytometry.

C2C12 N1\(\Delta\)ECD activation assays

Used in Figures 3 and S3.

The procedure for activating the Notch pathway in C2C12-hN1ECD cells was as follows: Cells were cultured in 10 \(\mu\)M DAPT (Sigma-Aldrich) until the experiment. In order to wash out DAPT, cells were washed quickly twice and a third time for 5 min with media at room temperature. Finally, cells were incubated in medium containing the appropriate activating DAPT concentration (0, 0.3, or 0.5 \(\mu\)M) at 37 C for the required activation duration (5 min, 15 min, 30 min, or until RNA extraction, i.e., sustained). In order to generate a pulse of activation, medium was then replaced with fresh 10 \(\mu\)M DAPT medium.
RNaseq

Used in Figures 3 and S3.

RNA was prepared using the RNeasy kit (QIAGEN) and submitted to the Caltech sequencing core facility, where cDNA libraries for RNaseq were prepared according to standard Illumina protocols. 100 base single-end read (100SR) sequencing was performed on a HiSeq2500 machine at the same facility. Reads were assembled, aligned, and mapped to the mouse genome (mm9 assembly) on a local instance of the Galaxy server, using Tophat. Cufflinks was used to calculate FPKM values.

In the analysis, we focused first on genes that showed > 5 fold-changes in their FPKM values (highlighted in Table S1). We further narrowed our subsequent analyses to the transcription factors Hes1, Hey1, and HeyL, because their promoters were shown to directly bind NICD by ChiP-Seq, they show early and strong (> 10-fold) responses to NICD, and they are key factors mediating Notch responsive behaviors in many contexts. These are also the only Hes and Hey family genes that activate in response to Notch in C2C12 cells (Castel et al., 2013). The RNaseq experiment did show upregulation of other genes, but we did not focus on them either because they were not transcription factors (such as Jag1 or Nrarp), or were not direct NICD targets based on the ChiP-Seq data.

RT-qPCR

Used in Figures 3 and S3.

RNA was prepared using the RNeasy kit (QIAGEN). cDNA was prepared from 500ng RNA using the iScript cDNA synthesis kit (Bio-Rad). 0.5 μL cDNA was used per 10 μL RT-qPCR reaction mix containing 1X iQSYBR Green Supermix (Bio-Rad) and 450 nM total forward and reverse primers. Reactions were performed on a BioRad CFX Real-Time PCR Detection System using a 2-step amplification protocol, with the following thermocycling parameters: 95°C, 3 min followed by 40 cycles of 95°C, 10 s (melting) and 55°C, 30 s (annealing + extension). All reactions were performed in duplicate.

Western blot analysis of NICD

Used in Figure S3

For this analysis, 0.5x10^6 - 1x10^6 cells were trypsinized after treatment, spun down in excess PBS, and lysed using Lithium Dodecyl Sulfate (LDS) buffer also containing reducing agents (DTT + 2-Mercaptoethanol) and Protease Inhibitors (Roche). Standard procedure was used for LDS-PAGE gel electrophoresis and transfer to nitrocellulose (iBlot, Thermo Fisher Scientific). Cleaved NICD (1:1000, Cell Signaling Technology, Catalog # D3B8) and GAPDH (1:5000, Abcam, Catalog #6C5) were detected using monoclonal antibodies. The blots were subsequently stained using HRP-conjugated secondary antibodies and detected using the Enhanced Chemiluminescence system (Pierce).

CHO-C2C12 co-culture assay

Used in Figure S4.

In preparation for the co-culture, C2C12-hN1 cells (4-6x10^4 cells in 12 well multi-well plate wells) were transfected with 60 pmol siRNA directed against mouse Notch2 (5'-UGAACUGCGAGGUGUAAAGGUC-3'), using Lipofectamine RNAiMAX (Life Technologies). 24h later, 3x10^4 CHO-K1 based Dll1- and Dll4- sender cells (pre-induced for > 48h) were plated within the two chambers of ibidi culture inserts (ibidi USA) on hamster fibronectin-treated (5 mg/ml in PBS, incubated for 3-5h at RT) surfaces of 24-well glass bottom plate wells. Once cells had attached to the surface (< 6h), inserts were removed and previously prepared C2C12-hN1 cells were plated, in 5 μM DAPT media, at high density so as to cover the gaps on the surface. After 12h, DAPT was washed out and cells were allowed to signal for 6h, after which the cultures were fixed in 4% formaldehyde at room temperature for 10 mins.

in ovo Electroporation

Used in Figures 4 and S4.

Batches of eggs were selected at random for electroporation with either Dll1 or Dll4, and the final data represents experiments conducted on at least two separate batches. The neural tubes of HH stage 12-13 embryos were injected with plasmid DNA (5 mg/ml) and electroporated by applying a series of current pulses (25V, 5x, 30 ms pulses separated by 100 ms) at the level of the pre-somitic mesoderm. 20h post-electroporation, embryos were screened for GFP fluorescence. Healthy embryos showing strong fluorescence in the neural crest were dissected (to remove extra-embryonic tissue) in Ringer’s solution and transferred to freshly prepared 4% paraformaldehyde, on ice. Embryos were fixed overnight at 4 C.

Hybridization Chain Reaction Fluorescence In Situ Hybridization

Used in Figures 4 and S4.

The hybridization chain reaction fluorescence in situ hybridization (HCR-FISH) protocol was based on a previously described protocol (Choi et al., 2016). Briefly, in situ HCR-FISH detection involves the following steps: 1. Dehydration and rehydration of embryos in MeOH, 2. Overnight hybridization with probes at 45 C, 3. Removal of unbound excess probes through washes at 45 C, 4. Overnight amplification at room temperature, and 5. Removal of excess amplifier. Each gene of interest was detected using 6 probes. At most three genes were detected simultaneously, typically EGFP, MyoD1, and Hes1, Hey1, or HeyL. After HCR processing, portions of the embryos anterior to the forelimbs were removed. Embryos were then mounted on glass-bottom multiwell plates in 1% agarose, with the dorsal surface in contact with the glass.
Confocal laser-scanning microscopy of embryos

*Used in Figures 4 and S4.*

Samples were imaged on a Zeiss LSM700 or using a 20x (0.8 NA) dry objective. For embryos, Z stacks were acquired using Zen software (ZEISS) and 3D-reconstructed in Imaris 8.0 (Bitplane). Optical slices in Imaris were used to remove obscuring auto-fluorescence from residual extra-embryonic tissue in the reconstructed images, without affecting signal in the areas of interest. For cell-culture Z stacks, the sum was projected in 2D using ImageJ.

Quantification of fold-changes in MyoD1, Hes1, and Hey1 gene-expression (*Used in e8 Cell* b8

Quantification of fold-changes in MyoD1, Hes1, and Hey1 gene-expression (*Used in Table 1*)

3D images of transverse optical sections of the interlimb region of the trunk (containing 3-5 pairs of somites per image), were sorted randomly, and then scored blindly for differences in somite MyoD1 levels between the electroporated and control sides of the embryo. The scoring procedure was as follows: any features that might reveal the specific experimental perturbation (Dll1 or Dll4 ectopic expression), such as image filenames, differences in pseudo-color attributes, or information from secondary channels, were removed before the files were re-ordered using a pseudorandom sequence. Subsequently, images were scored blindly, comparing MyoD1 signal in somites on the electroporated side with signal in the corresponding somites on the control side, as long as the two somites were level with each other. This requirement minimizes imaging artifacts. Finally, sample images were re-matched with the perturbation type and scores were tallied. The number of embryos scored per condition (11 Dll1 expressing embryos, 10 Dll4 expressing embryos, 61 somites for each perturbation) is standard for this type of quantification (Rios et al., 2011).

Quantification of fold-changes in MyoD1, Hes1, and Hey1 gene-expression (*Used in Figure S5C*)

The DML regions of the somites on the electroporated and control sides were manually identified in Z-projections of 3D-reconstructed confocal images (see above), and the maximal HCR-FISH staining intensities (90th percentile values within identically-sized areas on both sides) were calculated. The reported fold-changes represent the ratio of these values for electroporated versus control DMLs.

Immunofluorescence detection of transendocytosed Notch in co-cultures

*Used in Figures 5 and S5*

Sender cells and receiver cells were co-cultured on glass-bottom dishes, in the excess sender configuration, as described above. After 24h of co-culture, cells were fixed in 4% formaldehyde (diluted in PBS). All subsequent steps were carried out in blocking solution (2% Bovine Serum Albumin diluted in PBS). Following 1h of incubation at room temperature, samples were incubated overnight at 4 C with 1:250 mouse anti-hNotch1 (Biolegend Catalog No. 352014, RRID AB_10899408). Samples were then washed and incubated in an anti-mouse secondary antibody conjugated to Alexa Fluor 488 (Life Technologies). After room temperature washes, samples were permeabilized in 0.3% Triton X-100 (Sigma-Aldrich) for 1h. Samples were then again incubated in 1:250 anti-hNotch1 overnight at 4C, following which they were incubated in Alexa Fluor 647 conjugated anti-mouse antibody (Life Technologies).

Confocal imaging and quantification of transendocytosed Notch

*Used in Figures 5 and S5*

Immunostained cultures (see above) were imaged as Z stacks (0.8 µm intervals) on an LSM800 inverted confocal microscope using a 100x (1.3 NA, oil-immersion) objective. Sender cells abutting receiver cells (or distant from them, for background estimation) were manually segmented in ImageJ software, and stacks composed of 5 slices each were exported to MATLAB. In MATLAB, pixels within the stacks were categorized as being either intracellular, or belonging to the cell surface, based on the intensity of pre-permeabilization stain. Only cells that showed mean dispersed staining intensities higher than the median of the background staining levels were included in further analysis. This selected cells that were likely to be active senders (especially in the Dll1 case); we verified that none of the cells eliminated at this step displayed puncta. Next, in order to identify puncta, thebwconncomp function in the Image Processing Toolbox was used to assess 3-D connectivities of intracellular pixels possessing intensities above a fixed threshold and to group them into puncta of sizes >6 pixels. Several threshold/puncta size combinations were tested; one pair of values that returned puncta numbers most consistent with visual estimation was chosen. Qualitative conclusions remained the same for a range of threshold/size values. Pixels with intensities below the threshold, or failing to be included in such puncta were deemed part of the ‘dispersed’ staining.

QUANTIFICATION AND STATISTICAL ANALYSIS

Statistics

The non-parametric two-sided KS-test was typically used to compare the distributions of receiver activation amplitudes in response to different sender cell lines. All pairwise comparisons between samples fulfilled the criterion n1*n2/(n1 + n2) ≥ 4, where n1 and n2 represent the number of data points in two samples. Under this condition the KS-statistic is greater than the twice the inverse of the Kolmogorov statistic, and the calculated P-value is accurate. The non-parametric nature of the KS-test obviates the need to make assumptions regarding the shape of the distributions being compared. Furthermore, since the KS-test compares the distributions directly, and not the mean values, it is sensitive to differences in variance. Where the distribution itself is not shown, variance in
The distribution is displayed as standard deviations or s.e.m. The number of samples (‘n’) used for calculating statistics is indicated in the Figures or accompanying legends.

**DATA AND SOFTWARE AVAILABILITY**

**C2C12 hN1ΔECD transcriptomic data**

*Used in Figures 3 and S3*

The accession number for the raw sequencing reads and processed FKPM data reported in this paper is Gene Expression Omnibus (GEO): GSE72847.

**Code availability**

Image segmentation and cell tracking code used can be accessed at https://github.com/nnandago/cell2017-segtrack. Datasets and processing code is available upon request.
Figure S1. Characterization of Diverted Notch Reporter System and Dll1 Pulses, Related to Figure 1

(A) Simulation showing how the derivative of total fluorescence ($d/dt$Citrine = ‘Promoter Activity’) can recover underlying Notch activation dynamics. In response to a simulated pulse of Gal4 (orange trace), H2B-Citrine reporter fluorescence increases (‘Total Citrine’, green). The first derivative of total Citrine fluorescence (blue trace) provides a reconstructed estimate of the active Gal4 concentration (compare ‘Reconstructed Pathway Activity’ and ‘Actual Pathway Activity’).

(B) Correlation between Gal4 levels and promoter activity of the fluorescent reporter gene. (Inset) The cell line used for experiment expresses Gal4 (white, with a co-translationally expressed H2B-mCherry cassette, red) under control of a 4epi-Tetracycline (‘4epi-Tc’) inducible promoter. These cells also contain a fluorescent H2B-Citrine reporter gene (purple) that is responsive to Gal4 protein. (Main plot) Scatterplot of maximal (Gal4-T2A)-H2B-mCherry production rate (95th percentile value of promoter activity ($d/dtmCitrine$) versus the maximal promoter activity of the H2B-Citrine reporter gene rate (95th percentile value of promoter activity $d/dt$Citrine), for different levels of 4epi-Tc induction. Each circle represents the response in a single cell. The line connects median values within equally-spaced bins of H2B-mCherry levels, and error bars indicate 25th to 75th percentile levels of the distribution of values within each bin. Cell-to-cell variability in induction of the reporter is partly due to extrinsic noise and can be observed in the promoter activity of a co-expressed constitutive H2B- Cerulean gene. Note that since the units of total fluorescence are arbitrary, the resulting derivatives are also in arbitrary units (AU).

(C) Average of unprocessed total fluorescence traces (representing cumulative Notch activity) from the same set of activating cells plotted in Figures 1D and 1E. No alignment, normalization, or time derivative was applied to the data. A pulse of activity would lead to an initial rise in fluorescence followed by a plateau (as for Dll1, blue), while sustained activity would result in a continuous increase in total fluorescence (as for Dll4, red). Solid colored lines indicate median values, shaded regions indicate s.e.m. ‘n’ corresponds to number of traces included in the average.

(D) (Left) Median of total Citrine (top) and promoter activity (bottom) after alignment at the point of activation, defined as t = 0 (see STAR Methods). (Right) Corresponding plots after individual traces were not only aligned, but also individually normalized to the 90th percentile value within the averaging time window (STAR Methods). Note that these plots are the same as Figures 1D and 1E.

(E) Distribution of time points at which cells first activate during the Dll1 (top) or Dll4 (bottom) excess sender co-culture. Note that the time period prior to 15h was not considered in the analysis in order to eliminate contributions from transient effects on the reporter after transfer of cells to imaging conditions. Activation events in sender-receiver co-cultures are also delayed relative to the time of plating because cells are plated at sub-confluence (STAR Methods). Error bars indicate s.e.m (n = 2 experiments).

(F) (Left) Representative traces of cells displaying two pulses during co-culture with Dll1 (same dataset as in Figures 1D and 1E). Arrowheads indicate peak times. See Movie S3. (Right) Median response profile of receiver cells activated by Dll1 that could not be classified as single pulses, aligned after a period corresponding to the first pulse (7.5h, see STAR Methods). Each trace is aligned at a point when the promoter activity subsequent to the initial phase of activation (0-7.5h) again reaches the peak activity of the initial phase (n = 151 traces, 35% of all traces, cf. Dll1 response in Figure 1D). Each trace is individually normalized to the 90th percentile of promoter activity value between the alignment point and the end of the trace. Solid line indicates the median of the normalized promoter activity, light shaded areas show s.e.m. and the light gray shaded area indicates the standard deviation. The dark gray box indicates time period prior to the alignment time point, and the dashed horizontal line indicates the median level of (normalized) promoter activity at the alignment point. The median response in Dll1 returns to this level after 25h, consistent with a systematic second pulse of activity. (Inset) The same procedure applied to the Dll4 data (from Figure 1D) does not reveal a second pulse.

(G) Median promoter activity of receiver cells (same as Figure 1B) cultered on plate-bound Dll1 ECD-Fc. Also shown are standard errors of the mean (light colored lines) and standard deviations (gray areas).

(H) (Top, schematic) Expected Gal4 response (orange trace) in cells allowed to activate for 7h, then inhibited (indicated by gray shading) by treatment with DAPT (t = 0h). (Bottom) Median promoter activity in receiver cells (blue line) after DAPT treatment. Red line shows simulated response using fitted parameters for Gal4 protein half life (‘Gal4 t1/2’ = 4h, 95% bootstrapped confidence interval [3.8h, 4h]) and H2B-Citrine mRNA half life (‘Citrine t1/2 = 3.4h, 95% bootstrapped confidence interval [3.4h, 3.5h]). See STAR Methods for fitting procedure and Supplementary Information for model.

(I) Median (blue line) and standard error of the mean (lighter blue region) of normalized Dll1 pulse promoter activities (same as Figure 1E). ‘trise’ denotes time from 10% to 90% of the peak promoter activity, and FWHM denotes the full-width at half-maximum of the peak promoter value. ‘95% CI’ indicates bootstrapped confidence intervals.

(J) Dependence of FWHM (blue circles) and trise (purple circles) values on duration of underlying Notch activation (‘Activation pulse duration’) based on a mathematical model for pulse-like activation, using values calculated in panel H (see Supplemental Information). The horizontal gray line indicates measured values for these quantities.

(K) Simulated time-course of Gal4 protein (orange, dashed) and H2B-Citrine mRNA (blue) for 15 min Notch activation, using values for half-lives calculated in panel H. Note similarity with measured pulse shape in panel I.
Figure S2. Pulse Train Model and Analysis of Increased Time-Resolution Reporter, Related to Figures 1 and 2

(A–F) Comparison of simulated Dll1 pulse trains to observed Dll4-induced responses.

(A) (Schematic) Can sustained Dll4 signal (red) be composed of a series of Dll1-like pulses (blue)?

(legend continued on next page)
(B) Schematic diagram illustrating the pulse train simulation procedure (see also Supplementary Information). Each pulse train is constructed from a series of pulses with the averageDll1 pulse shape (Left schematic, median of averaged traces in Figure 1D), scaled by an amplitude randomly sampled from the empirically measured distribution ofDll1 pulse amplitudes (Center schematic, measured amplitude distribution shown in Figure 1F). Each pair of adjacent pulses (shades of blue) is temporally separated by an interval \( t \) chosen based on one of the underlying pulse models shown in panel C, and combined to generate a particular pulse train (Right schematic, gray line).

(C) Schematics illustrating the underlying pulsing models. In each case two example traces are shown (top and bottom). (Left) In the Periodic model, the interval \( t \) between adjacent pulses is fixed at a value \( T_{\text{period}} \) that can range from 1h to 8h. Periods greater than 8h result in oscillating pulse trains in which the individual pulses can be clearly discerned. (Right) In the Poisson model, the interval between pulses \( i \) and \( i+1 \), \( t_i \), represents the inverse of a pulse rate, \( r_i \), drawn from a Poisson distribution with parameter, \( \lambda \), ranging from 1/h - 15/h. (Center) The mixed models interpolate between fixed and random intervals. In these models, the interval \( t \) between adjacent pulses is drawn from a normal distribution with mean \( T_{\text{period}} \) (range 1-15h) and standard deviation \( s = 2.5h \) or 5h). See Supplemental Information for further details.

(D) For each simulation, 200 pulse trains were simulated (\( n' = 1 \) to 200). For each simulated trace, the median amplitude (Left) and the median temporal variability (Intra-trace variability, Right) were tabulated (see Supplemental Information).

(E) Relationship between median amplitude and median intra-trace variability for the different models. Each point corresponds to a different mean interval, and represents the median of 36 simulations. The black marker shows measured values of median amplitude and median intra-trace variability forDll4 traces (using the data in Figures 1D and 1E). (Inset) Zoom into the region (red box) of theDll4 data point (black). Error bars onDll4 marker represent s.e.m. (n = 200 traces). The gray shaded area delineates region of the plot that has median amplitude within the measured error in theDll4 median amplitude. Filled circles are points in the mixed (\( \sigma = 2.5h \), \( T_{\text{period}} = 10h \)) and Poisson (\( \lambda = 1/15h \)) models closest to the experimental measurement.

(F) Boxplot comparison between median intra-trace variability in the simulations highlighted in panel E (inset) and the intra-trace variability measured forDll4. For simulations, each point represents a single simulation (comprising 200 pulse trains). The colored horizontal lines (blue or green) represent the median value of these simulations (n = 36 simulations), while boxes delimit 25th - 75th percentile values. The black horizontal line represents a bootstrapped average of intra-trace variability values calculated from measuredDll4 data. See Supplemental Information for further details. \( P \)-value calculated by two-sided KS-test.

(G) (Schematic) Improved time-resolution reporter cell line, expressing the NotchECD (green)-Gal4 (orange) receptor and a Gal4-responsive H2B-3xCitrine (chartreuse) fused to 3'UTR (blue) derived from the mouse Hes1 gene (cf. Figure 1B, see STAR Methods).

(H) Median normalized promoter activity of reporter cells co-cultured with eitherDll1- (blue) orDll4- (red) expressing sender cells. Also shown are s.e.m. (light colored lines) and standard deviations (gray areas). See STAR Methods for alignment and normalization procedure. Note the decreased values of rise time ('\( t_{\text{rise}} \)') and full width at half max ('\( \text{FWHM} \)') of theDll1 pulse compared to S1I, which shows corresponding values for the original reporter cell line.

(I) Boxplots of maximal promoter activities of reporter cells in response toDll1 (blue) orDll4 (red), calculated for the traces averaged in H. Colored lines represent median values, and boxes delimit 25th to 75th percentile values. \( P \)-value calculated using two-sided KS-test.

(J) Distribution of mCherry levels inDll1-T2A-H2B-mCherry sender cells used in Figures 2B and 2C, with 'Low', 'Medium', and 'High' fractions delineated. 'n' values correspond to the number of sender cells in the experiment (cf. Figure 2B).

(K) Distribution of mCherry levels inDll4-T2A-H2B-mCherry sender cells used in Figures 2B and 2C, with 'Low', 'Medium', and 'High' fractions delineated. 'n' values correspond to the number of sender cells in the experiment (cf. Figure 2B).

(L) Comparison of median promoter activities in activated receiver cells adjacent to sender cells expressing Low (light blue), Medium (navy blue), or High (dark blue) levels ofDll4 (same designations as used in Figure 2B). Grey circles represent individual responses, solid horizontal lines represent medians, while the boxes delineate 25th - 75th percentile values. \( P \)-values calculated by two-sided KS-test.
Figure S3. Duration Dependence of Gene Expression in C2C12 Cells, Related to Figure 3

(A) Transcript levels in C2C12-N1ΔECD cells in DMSO-treated cells versus cells activated for 1h by DAPT removal. Circled genes are putative direct Notch targets, upregulated by > 5-fold (Table S1).

(B) Transcript levels in C2C12-N1ΔECD cells in DMSO-treated cells versus cells activated for 6h by DAPT removal. Circled genes are putative direct Notch targets upregulated by > 5-fold (Table S1).

(C) Response of HeyL (orange) to complete DAPT removal for 5 min, 15 min, or 30 min followed by replenishment (‘Pulse’), or removal without replenishment (‘Sustained’). Error bars represent s.e.m. for duplicate experiments.

(D) (Schematic) For the experiment, DAPT is either washed out completely (final concentration 0 μM DAPT) or partially (final concentration 0.3 μM) for 3h (green line). (Main panel) Western blot analysis of cleaved NICD levels in cells after different DAPT washout treatments. NICD is detected using an antibody that detects the cleaved version (with N-terminal Val at amino acid position 1744, see STAR Methods). GAPDH levels represent the loading control. Note similar levels of NICD at 15’ and 3h samples. The vertical line indicates splicing together of two parts of the same western blot (removing intervening lanes).

(E) (Schematic) Expected time evolution of NICD concentration within cells for complete (green line) or partial (chartreuse line) DAPT washout. Black markers indicate points at which NICD levels are measured in D.

(F) Predictions from Amplitude-only model and Duration-based models for control of Hey expression. (Top) In the amplitude-only model, the rate of Hey1/L expression is determined by the concentration of NICD. Therefore, the 15’ complete DAPT washout (green line) and 3h partial DAPT washout (chartreuse line), which result in comparable NICD levels, should result in similar increases in Hey1/L expression levels (Δ) during a given time window at the end of treatment. (Bottom) In the duration based model, the rate of Hey1/L expression depends on the duration of NICD activation. Specifically, a short period of Notch activation (15’) leads to a smaller increase in Hey1/L levels (Δ) during a given time window than a longer period of Notch activation (3h), despite similar NICD levels at the two time points.

(G) Median expression level of Hey1 and HeyL, (each normalized to its expression level at 3.25h with complete DAPT removal) 15 min prior to (‘T-15 min’, empty bars) or 15 min after (‘T+15 min’, filled bars) a 15 min, 30 min, or 3h period of Notch activation with complete (dark green) or partial (chartreuse) DAPT removal. Error bars represent s.e.m. calculated from duplicate experiments.

(H) Maximum Hes1 (blue), Hey1 (burnt orange), and HeyL (orange) mRNA levels measured during the [0h, 1h, 4h] time-course, in response to sustained reduction of DAPT concentration to indicated levels (0, 0.3, or 0.5 μM). Error bars represent SEM calculated from duplicate experiments.

(I) Western blot analysis of cleaved NICD levels in N1ΔECD cells (see Figure 3A) and receiver cells expressing full-length human Notch1 (Figure S4D). N1ΔECD cells (co-cultured with CHO-K1 control cells to match total number of cells in co-culture) were maintained in DAPT (‘–’) or DAPT was washed out completely for 30 min (‘30 min’), Notch1 receiver cells were co-cultured with CHO-K1 control cells (‘–’) or sender cells expressing maximal levels of Dll4 for 8h. NICD is detected using an antibody that detects the cleaved version (with N-terminal Val at amino acid position 1744, see STAR Methods). GAPDH levels represent the loading control.
Figure S4. Dynamic Ligand Discrimination in C2C12 Cells, Related to Figure 3

(A–C) Dll1 and Dll4 activate Notch1 using pulsatile and sustained dynamics, respectively, in C2C12 cells. (A) (Top) Engineered C2C12 sender cell lines contain stably integrated constructs expressing Dll1 (blue) or Dll4 (red), each with a co-translational (T2A, gray) H2B-mCh readout (purple). (Bottom) Receiver cell lines stably express a chimeric receptor combining the Notch1 extracellular domain (N1ECD, green) with a Gal4 transcription factor (orange) in place of the endogenous intracellular domain. Cells also contain a stably integrated H2B-3xCitrine fluorescent reporter (green), destabilized at the mRNA level by fusing it with a 3'UTR derived from the Hes1 cDNA (light blue).

(B) Median normalized promoter activities in receiver cells co-cultured with an excess of Dll4 (red) or Dll1 (blue) senders. Traces were aligned (see STAR Methods for special alignment procedure) and normalized to their 90th percentile value prior to averaging. Also shown are standard error of the mean (light colored lines), and standard deviations (gray areas).

(C) Boxplots of maximal promoter activities of the responses averaged in panel B. Each gray circle represents the response of a single cell to Dll1 (blue) or Dll4 (red). Colored lines represent the median, while boxes delimit 25th to 75th percentile values. P value calculated using two-sided KS-test.

(D) (Schematic) RT-qPCR based measurement of Hes1/Hey1/HeyL response in C2C12-Notch1 + CHO-Dll1/4 co-cultures. C2C12-Notch1 receiver cells, expressing full length Notch1 (green) are co-cultured with CHO-K1 sender cells expressing either Dll1 or Dll4 (gray). The target genes (shown as brown gene target and corresponding brown mRNA) in C2C12-Notch1 receiver cells are specifically amplified using mouse-specific primers.

(E) RT-qPCR expression levels of Hes1, Hey1, and HeyL in C2C12-Notch1 receiver cells co-cultured with CHO-K1 control cells (black), Dll1- (blue), or Dll4-expressing (red) sender cells for 8h. Error bars represent standard error of the mean (n = 2 replicates). F–H, Characterization of Hey transcriptional response at the single-cell level using HCR-FISH.

(F) CHO sender cells (highlighted with blue nuclei) and C2C12-Notch1 receiver cells (gray cells) were co-cultured such that they interface along a line (see STAR Methods).

(G) HCR-FISH detection of Hey1+HeyL mRNA (red) in C2C12-Notch1 cells co-cultured with sender cells (blue nuclei shown) expressing low or high levels of Dll1 (left panels) or Dll4 (right panels). Arrowheads indicate cells showing clear HCR-FISH signal.

(H) Quantification of panel G images. Values indicate number of pixels in the left half of images that lie above a threshold intensity value.
Figure S5. Quantitation of Effects on MyoD1, Hes1, and Hey1 in Ovo and Gene Expression Response to Dll1ECD–Dll4ICD Ligand, Related to Figure 4 and Table 1

(A) Schematic for chick electroporation experiments. The neural tubes of HH stage 12–13 embryos were injected in ovo with CAG-Dll1/4-T2A-EGFP plasmid and a rightward current was applied at the level of the pre-somitic mesoderm. 20h later embryos were dissected, fixed, and subjected to 3-color whole-mount HCR-FISH to detect GFP (blue), MyoD1 (green), and Hes1/Hey1/HeyL (not shown). Inter-limb somites were then imaged using scanning laser confocal microscopy.

(B) Representative image showing an increase in HeyL (red, right) in the DML of somites when the neural tube and crest were electroporated with Dll4-T2A-EGFP (blue, left). Arrows indicate the electroporated side.

(legend continued on next page)
(C) Fold-changes in expression of MyoD1, Hes1, and Hey1 within somites (see STAR Methods), in response to electroporation ofDll1 (blue bars) or Dll4 (red bars), calculated relative to the control side. ‘n’ values correspond to number of somites analyzed.

(D) Cell surface levels of natural and chimeric ligands, detected using soluble Notch1ECD-Fc (see STAR Methods). (Left) Dll1 (blue bars) and Dll1ECD-Dll4ECD (purple bars) are detected at similar surface levels before (‘-’, light shading) and after (‘+’, dark shading) maximal induction with doxycycline (‘dox induction’). (Right) Dll4ECD-Dll1ICD (magenta bars) and Dll4 (red bars) are detected at similar surface levels before (light shading) and after (dark shading) maximal induction. Error bars indicate standard deviation. Note that Notch1ECD has a higher affinity for the Dll4ECD than the Dll1ECD (Andrawes et al., 2013); hence ligand levels detected after induction in the left plot are higher than in the right plot.

(E) (Top, schematics) Expected dose-response behavior of Hes1 and Hey1 gene expression to varying Dll1 (blue) or Dll1ECD-Dll4ICD (purple) ligand levels if (Left) ligands varied only in the amplitude of signaling (‘Amplitude-only’ model) and if (Right) ligands produced qualitatively different gene expression (‘Qualitative-difference’ model). In an amplitude-only model, it is possible to simultaneously match the Hes1 and Hey1 gene expression levels produced by one ligand by varying levels of the other. On the other hand, if ligands produced qualitatively different Hes1/Hey1 expression patterns, there exist Hes1/Hey1 expression levels induced by one ligand (for e.g., Dll1) that cannot be matched by varying the levels of the other ligand. (Bottom) Hes1 and Hey1 expression levels in N1 receiver cells co-cultured with CHO-K1 cells (control, black marker), Dll1 senders (blue marker), or sender cells expressing three different levels of Dll1ECD-Dll4ICD (purple marker, increasing marker sizes indicate increase levels of level induction). Error bars represent SEM from duplicate experiments. (Inset) Comparison of fold-changes in Hes1, Hey1, and HeyL expression levels (relative to CHO-K1 control co-culture) in receiver cells co-cultured with Dll1 senders (blue bars, same experiment as indicated by blue marker) or Dll1ECD-Dll4ICD senders (purple bars with orange outlines, same experiment as indicated by purple marker with orange outline).

(F) Schematic of types of staining observed in co-culture NotchECD immunostaining assay and the molecular species they are expected to represent. Intracellular Notch1 will only be labeled only after cell-permeabilization and should only be detected in a single channel (shown here as red). White circles within cells represent endosomes. All cell-surface Notch1, including unbound and ligand-bound forms, will be labeled both before and after cell-permeabilization; it should thus be detected in two channels (shown here as red + green = yellow). Receiver cells can be distinguished from sender cells based on expression of nuclear H2B-mCherry, which will also be detected in the red channel. Ligand-NotchECD complexes within the sender cell could exhibit two types of staining: large, bright puncta, corresponding to endocytosed ligand-receptor clusters, and low intensity staining, corresponding to individual or few endocytosed ligand-receptor complexes.
Figure S6. Role of Ligand Intracellular Domains in Receptor Transendocytosis, Related to Figure 5
(A) Representative image showing Dll4 co-localization with Notch in clusters at the cell interface between the Notch1 receiver cell (‘R’) and a Dll4-FLAG expressing sender cell (‘S’) shown in Figure 5B. Cells are immunostained with antibodies for Notch1ECD (red, green, left panels, same as Figure 5B) and FLAG (blue, right panels). 

(legend continued on next page)
right panels). Cell-surface receptors appear as yellow because they are labeled in two channels (red + green). White arrowheads indicated co-clustered ligands and receptors at the interface between the sender and receiver cell.

(B) Representative image showing co-localization of transendocytosed NotchECD puncta with the early endocytic marker Rab5 within a Dll4ECD-Dll1ICD expressing sender cell (‘S’) next to a Notch1 receiver cell (‘R’). Cells are immunostained for Notch1ECD (red, green) and the early endocytic marker Rab5 (blue). Cell-surface receptors appear as yellow because they are labeled in two channels (red + green).

(C) Transendocytosis staining patterns in Dll4ECD-Dll1ICD senders (Left) or Dll4 senders (Right) adjacent to or not adjacent to receiver cells within the same co-culture. (Top Left) Median values of the number of puncta detected (see STAR Methods) in Dll4ECD-Dll1ICD sender cells that are adjacent to (magenta bar) or not adjacent to (black bar) receiver cells. (Top Right) Corresponding median values of number of puncta detected in Dll4 cells. (Bottom Left) Median values of the mean pixel intensity of dispersed signal in Dll4ECD-Dll1ICD sender cells that are adjacent to (magenta bar) or not adjacent to (black bar) receiver cells. (Bottom Right) Corresponding median values of dispersed signal in Dll4 cells. Error bars represent standard error of the mean. *P value calculated using the two-sided KS-test. ‘n’ indicates number of sender cells considered in the analysis.

(D) Median values for intensities of pixels categorized as ‘disperse’ staining (light shading) and for cumulative intensities of ‘puncta’ (dark shading) in immunostained co-cultures of receiver cells with Dll1 (blue bars), Dll4ECD-Dll1ICD (magenta bars), or Dll4 (red bars) sender cells. Error bars represent standard deviations from the mean.

(E) Flow cytometry histograms of activation levels in a Notch-Gal4 receiver cell line (same as used in Figures 1, 2, and 5A; background activity indicated by gray histogram) co-cultured with Dll1 (blue), Dll4ECD-Dll1ICD (magenta) or Dll4 (red) cells expressing ligand levels similar to that used in Figures 5C and 5D.

(F) Representative image of co-culture of Notch1 receiver cells (‘R’) and Dll1 sender cells (‘S’), immunostained for NotchECD. The red channel shows staining for intracellular receptors. Cell-surface receptors appear as yellow because they are labeled in two channels (red + green).

(G) Median values of the background subtracted mean pixel intensity of dispersed signal in Dll1 (blue) or Dll4ECD-Dll1ICD (magenta) sender cells that abut receiver cells (see STAR Methods). (Right) Median values of the number of puncta detected (see STAR Methods) in Dll1 (blue) or Dll4ECD-Dll1ICD (magenta) sender cells that abut receiver cells. Error bars represent standard error of the mean. *P value calculated using the two-sided KS-test.

(H) Median values of the rate at which receiver cells are activated by Dll4ECD-Dll1ICD (magenta) sender cells or Dll1 (blue) sender cells in excess receiver conditions. Error bars indicate standard error of the mean.

(I) (Schematic) Molecular basis of pulsatile and sustained signaling. Top, (1) Dll1 ligands (blue) do not activate Notch1 receptors (green) efficiently in single ligand-receptor complexes, but (2) ligand-receptor complexes can assemble into larger clusters, leading to (3) efficient transendocytosis and coordinated activation of constituent receptors in the receiver cell. (4) This releases a burst of NICD (green ovals) which is reflected in a strong pulse of downstream response in receiver cell (green shading, green trace in plot). (5) The system resets (locally) and awaits another clustering event. Bottom, Dll4 ligands (red) transendocytose Notch1 (green) receptors efficiently even when bound in single ligand-receptor complexes (1, 2), or in small clusters (not shown). (3-5) This leads to a steady rate of NICD cleavage (green ovals), and a sustained downstream response in the receiver cell (green shading, green trace in plot).