Introduction

The Notch signaling pathway mediates cell-to-cell communication in several developmental contexts [1–4]. This communication occurs through the binding of the Notch receptor in a cell membrane to its ligand (e.g., Delta) in a neighboring cell [5], what herein we refer as trans-interactions. The bound complex is then claved and its intracellular domain (NICD) targets gene expression within the cell that harbors the bound receptor [6]. In the case of neural development, Notch signaling mediates lateral inhibition, which drives the selection of cells that express high levels of proneural genes and ultimately become neurons [7–11]. Notch signaling activity inhibits the proneural genes, which in turn activate Delta expression. As a result, a cell committed to the neural fate with high proneural gene and Delta levels inhibits its neighboring cells from adopting the same fate (i.e., it performs lateral inhibition) [12]. When all precursor cells are initially equivalent and signal similarly, mutual lateral inhibition arises and can drive a spontaneous spatially periodic selection of precursors by amplifying the initial small differences between them [13].

Different experimental evidences show that the Notch receptor can bind as well to the ligands when they are both in the same cell, what we refer as cis-interactions [5,14–18]. Cis-interactions drive a reduction of Notch signaling by sequestering the receptor and impeding its signaling [14,16,18–28]. This is known as cis-inhibition and its effects have started to be theoretically and computationally addressed too [25,29–38]. These studies have revealed that cis-inhibition can facilitate patterning by promoting faster responses, enhancing robustness and precision, and relaxing the constraints required for patterning [25,29,32,33,35,37,38].

Although cis-interactions commonly inhibit Notch signaling [14,16,18–28], Notch activity coming from cis-interactions has been proposed for specific scenarios [39–43]. For instance, Coumaillé et al. (2009) pointed to cell-autonomous Delta-dependent active Notch in Sara endosomes of Drosophila bristle precursor cells. Guy et al. (2013) indicated that cell-autonomous Notch-mediated activation of the cell-cycle regulator c-Myc in mouse T-cells is impaired when ligand-receptor cis-binding is prevented. The mechanism by which cis-driven signaling can occur in these scenarios is still missing. According to the
mechanism proposed by Fürthauer and González-Gaitán (2009), ligand-receptor binding within multivesicular endosomes could drive the release of Notch intracellular domain, driving Notch activation. This binding would occur in anti-parallel configurations (like for trans-interactions), as opposed to the parallel binding that is commonly associated to cis-interactions when occurring in the cell membrane and which is believed to prevent signaling. The interplay between Notch and the endocytic routes is starting to be uncovered and may shed light on this issue.

In the present work we take advantage of computational and mathematical modeling to address the question of which would be the effect expected from cis-signaling when another signaling source that also uses the Notch receptor is acting (Fig. 1). Since both signaling sources use the Notch receptor, both can compete for it. The additional source of signaling, hereinafter referred to as primary signaling source, can be associated with trans-interactions. Yet, our modeling approach is not exclusive for such trans-interactions. The primary signal can be also driven by additional alternative mechanisms. In this context, recent work has shown that ligand-independent Notch activity can arise from the binding of Notch to other factors and from impaired endocytic regulation [39,40,44–54]. For instance, a recent study in Drosophila has detected a ligand-independent Notch signal that has a significant role in their development [52,53]. Our results show that when acting together with a primary signal, cis-signaling can act as cis-activation or as cis-inhibition. Competition between signaling sources underlies this switch. We establish under which conditions each regime arises. An extensive analysis of the parameter space shows that cis-inhibition enriches patterning, as opposed to cis-activation. Cis-inhibition promotes pattern multistability and modulates the selection of precursors. In addition, cis-inhibition facilitates cell-autonomous bistability.

Results

A simple model for lateral inhibition with cis-signaling

We set a mathematical phenomenological model that includes two sources of Notch signaling: a primary signaling source and signaling driven by cis-interactions between the Notch receptor and its ligand within the same cell. The primary signal can be driven either by trans-interactions between the Notch receptor and its ligand in an adjacent cell (Fig. 1A) or through ligand-independent mechanisms (Fig. 1B). We assumed that both the primary signaling source and cis-interactions drive the same type of signal. We set primary Notch signaling to occur in a graded non switch-like fashion and to saturate to a maximal value, as recently experimentally reported for trans-interactions [25]. We assumed that cis-interactions can drive a graded increasing production of Notch signal activity with ligand up to saturation like the primary signaling source does (Methods). When both signaling sources are acting, the production of signal that each of them drives depends on the other source since both sources use the Notch receptor to signal. Accordingly, these productions in cell i read (Methods):

\[ P_{li} = \frac{\mu}{1 + \mu + r_{cli}} \]  

(1)

\[ P_{2i} = \frac{cr_{li}}{1 + \mu + r_{cli}} \]  

(2)

where the primary source produces Notch signal activity at rate \( P_{li} \) whereas cis-interactions produce it at rate \( P_{2i} \). In the above equations \( li \) stands for the ligand activity in cell i and \( \mu \) is a measure of the amount of primary signaling source. Non-dimensional units are used with maximal signal production being 1 for the primary signal without any loss of generality.

We considered that each source can drive Notch activity production at a different rate such that it results into different values of the stationary saturated Notch activity. Parameter \( \epsilon \) accounts for the ratio of the stationary saturated Notch activity driven by cis-interactions over that one driven by the primary signaling source. A more complex biochemical reaction-based scheme indicates that \( \epsilon \) can be understood as the ratio between the signaling efficiency of the two sources, which depends on the signaling rate and the stability of each source (Methods). Therefore, we refer to \( \epsilon \) as relative signaling efficiency. \( \epsilon = 0 \) corresponds to the well-known cis-inhibition, in which cis-interactions titrate the receptor and drive no signaling. When \( \epsilon < 1 \), cis-interactions drive signaling less efficiently than the primary signaling source.

When the primary signaling source is driven by trans-interactions we have \( \mu = r_{i} \langle L_i \rangle \), where \( \langle L_i \rangle \) stands for the weighted average of ligand activity in cells adjacent to cell i (Methods). Parameters \( r_{i,\epsilon} \) arising for each Notch signaling source are the trans and cis-interactions strengths, which are related to the trans and cis-binding and unbinding rates of the receptor and ligand complexes (see Methods for their definition in a more biochemical reaction-based approach). The trans and cis-interactions strengths parameters \( r_{i,\epsilon} \) set the threshold values of ligand for signal activation.

Figure 1. A model for Notch signaling driven by a primary signaling source and by cis-interactions. Cartoons of the Notch signaling components under study for (A) two adjacent cells that interact and for (B) an isolated cell. Black arrows stand for activation while red blunt arrows denote inhibition. (A) The ligand (red) in a cell binds the Notch receptor (blue) in a neighboring adjacent cell (trans-interactions). This elicits a Notch signal (NICD) that inhibits ligand production in the adjacent cell. The ligand can also bind the receptor within the same cell (cis-interactions) and drive Notch signaling at a different strength (dashed arrow). (B) A primary signaling source that is ligand-independent is depicted as well as signaling driven by cis-interactions. In both panels, the Notch signal inhibits the ligand through the proneural genes.

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We set production of ligand to be inhibited by Notch signaling and took linear degradation for both signal and ligand activities (Methods). Taken together, the dynamics in cell *i* of the Notch signal *s* and the ligand *l* activities read:

\[
\frac{ds_i}{dt} = P_{1i} + P_{2i} - s_i \tag{3}
\]

\[
\frac{dl_i}{dt} = v \left\{ \frac{1}{1 + hs_i^m} - l_i \right\}, \tag{4}
\]

where time *t* is non-dimensional, \(v = v_l/v_s\), being \(v_l\) and \(v_s\) the degradation rates of the ligand and Notch signal respectively, *h* stands for the strength of ligand inhibition through Notch signaling and *m* represents an effective cooperativity of such an inhibition. When cis-interactions are not present \((r_c = 0)\), Eqs. 1–4 reduce to the model for Delta/Notch-mediated lateral inhibition dynamics early proposed by Collier et al. [13].

### A switch from cis-activation to cis-inhibition

We first evaluated whether the total Notch signaling increases (cis-activation) or decreases (cis-inhibition) when cis-interactions that drive Notch signaling are included. From Eqs. 1–3, it is obtained that cis-inhibition always occurs when cis-interactions do not drive a signal \((\epsilon = 0)\) as expected (Fig. 2A). In contrast, either cis-inhibition or cis-activation can arise when cis-interactions drive signaling on their own (Fig. 2B,C).

We explored under which conditions each cis-signaling regulatory role (cis-inhibition versus cis-activation) arises and found that cis-inhibition occurs for (Fig. 2D, Methods)

\[
\epsilon < \frac{\mu}{1 + \mu^2}, \tag{5}
\]

Otherwise, cis-interactions drive cis-activation (Fig. 2D). According to the above relation, cis-inhibition requires cis-signaling to be less efficient than the primary signaling source \((i.e., \epsilon < 1)\). Yet, less efficient cis-signaling does not ensure cis-inhibition. The above relation shows that the regulatory role of cis-signaling depends on the amount of the primary signaling source \((\mu)\) as well. When the primary signaling source comes from trans-interactions, the regulatory role of cis-signaling depends on the trans-interactions strength \((r_t)\). For \(\epsilon < 1\), the regulatory role switches from cis-activation to cis-inhibition as the primary source becomes more abundant \((\mu\) or \(r_t\) increases) (Fig. 2B).

The above result indicates that whether cis-signaling is acting as cis-activation or cis-inhibition does not depend on the strength of cis-interactions \((r_c; \text{Fig. S1})\). This is because the qualitative change (decrease or increase) in Notch signaling driven by the addition of ligand within a cell with primary Notch signal activity is independent of the amount of ligand being added. In contrast, the quantitative change depends on \(r_c\) and on the amount of added ligand (Fig. S1C).

**Cis-signaling can drive two distinct effective circuit architectures**

The above analysis only took into account the dynamics of the signal when two sources of signaling (primary and cis-driven) are competing for the Notch receptor, Eqs. 1–3. We then asked which is the effect of this switch between cis-inhibition and cis-activation on the overall signal and ligand dynamics. To this end, we first

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**Figure 2. A switch between cis-activation and cis-inhibition.** (A–C) Stationary Notch signal in a cell \((s_i)\) versus the amount of ligand in that cell \((l_i)\) and the amount of primary signaling source \((\mu)\) for (A) \(\epsilon = 0\), (B) \(\epsilon = 0.1\) and (C) \(\epsilon = 1\). Red lines show the Notch signal dependence on \(l_i\) in the absence of the primary signaling source \((\mu = 0)\) and for a primary signaling source with \(\mu = 1\). Decreasing curves indicate cis-inhibition and increasing curves show cis-activation. In B, cis-interactions drive cis-activation at low \(\mu\) values, whereas they drive cis-inhibition at higher \(\mu\). (D) Parameter space showing where cis-activation (gray region) and cis-inhibition (white region) occurs, according to inequality 5. (E–F) Effective circuit architectures of the model when cis-interactions drive cis-activation (top) and cis-inhibition (bottom) for (E) isolated cells with a primary signaling source (straight arrow) and for (F) two adjacent cells that interact through trans-binding. Black arrows stand for activation, while red blunt arrows for inhibition. Parameter values: \(r_c = 10\) in panels A–C, \(h = 4\) and \(r_c = 1\) in panel D.

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evaluated the effective genetic circuits that arise when the dynamics of the ligand, Eq. 4, is included. By “effective” we mean that the circuit does not describe the individual interactions per se but their resulting regulatory role, which takes into account the context in which they occur. Therefore, we considered which circuit architecture arises when cis-interactions perform cis-activation and when they drive cis-inhibition. It can be readily seen that cis-interactions, coupled to a primary signal, give rise to two different effective genetic circuits (Fig 2E). When cis-interactions drive cis-activation, a negative intracellular transcriptional feedback loop arises. In contrast, a positive feedback loop emerges when cis-interactions drive cis-inhibition. It is worth stressing that the amount of primary signal (μ) and the relative cis to primary signaling efficiencies (ɛ) control which of the two effective architectures is acting by setting which is the regulatory role of cis-signaling.

When the primary source is trans-interactions, these intracellular feedbacks are coupled to the intercellular mutual inhibition loop that is characteristic of lateral inhibition (Fig. 2F). Positive and negative feedback loops are well known to drive different dynamics, like bistability for the former and homeostasis for the latter (see for instance [55]). Accordingly, we can expect different roles of cis-signaling on patterning and cell fate choices, depending on whether it is in the cis-inhibition or cis-activation regime.

Cis-activation inhibits fine-grained pattern formation, while optimal values of cis-inhibition promotes it

From theoretical arguments it has been shown that the intercellular positive feedback mediated by trans-interactions is a sufficient mechanism for spontaneous pattern formation [13]. This feedback amplifies small differences in ligand and signal levels between cells and drives a mostly periodic lateral inhibition pattern composed of two cell types (Fig. 3A) [13]. This type of periodic pattern arises spontaneously for a large range of trans-interactions strengths (r_1) and above a minimal ligand inhibition strength (b) (Fig. S2 and Methods) [13]. The pattern solution exists and is stable in an even larger region of the parameter space, with a minimal yet lower ligand inhibition strength (Fig. S2 and Methods) [10,56].

From the effective circuit architectures (Fig. 2F), we may propose that cis-activation can inhibit pattern formation. Cis-activation drives a negative feedback loop within cells which could dampen the amplification of differences between precursor cells driven by trans-interactions. In contrast, we may expect that cis-inhibition can promote patterning as it can enhance amplification of precursor differences by driving an additional positive feedback loop within cells (Fig. 2F). This latter expectation is in agreement with previous computational studies on cis-interactions that do not trigger a signal [25,29,33,35].

We first evaluated the case of efficient cis-signaling (ɛ = 1) driving cis-activation whatever the strength and amount of trans-interactions. We explored extensively the parameter space to characterize where lateral inhibition patterning occurs (Methods). We chose a regime for which the lateral inhibition pattern can arise for a wide range of trans-interactions strengths (r_1) in the absence of cis-interactions (r_c = 0; Fig. 3B). When cis-interactions are added (r_c > 0), pattern formation becomes forbidden (Fig. 3B).

At very weak cis-interactions strengths, pattern formation is still possible albeit in a reduced range of trans-interactions strengths. We conclude that cis-activation inhibits patterning. Following the same procedures, we evaluated whether pattern formation is promoted by cis-inhibition. To this end, we analyzed the well-known case of cis-interactions which do not elicit signaling (ɛ = 0), driving always cis-inhibition. In this case, it is known that cis-inhibition facilitates patterning by allowing it for graded trans-signaling and graded ligand inhibition (h = 1) [25] (Fig. S3A). We found that cis-inhibition facilitates spontaneous patterning as well for low ligand inhibition cooperativities (Fig. S3B). At higher cooperativities, the analysis showed that cis-inhibition promotes patterning too by reducing the minimal ligand inhibition strength (b) required for patterning (Fig. 3C). Cis-inhibition can have a detrimental effect as well (Fig. 3C). When the strength of cis-interactions is too high compared to trans-interactions strength (r_c >> r_1), the coupling between cells becomes less relevant, impeding patterning. Yet, if the trans-interactions strength increases, patterning is enabled (Figs. 3C and S3A–C).

We finally evaluated a more complex scenario in which cis-signaling switches from cis-activation to cis-inhibition when trans-interactions strength (r_1) increases (Fig. 3D). Our results show that in the cis-activation regime, pattern formation is inhibited, since an increase in cis-interactions impedes pattern formation. In contrast, in the cis-inhibition regime, pattern formation is facilitated since a minimal value of cis-interactions strength (r_c) enables spontaneous patterning. These results confirm the existence of distinct regulatory roles of cis-signaling (cis-activation versus cis-inhibition) as a function of the amount of trans-interactions as well as the different effect each of them has on lateral inhibition patterning. Taken together, the results suggest

Figure 3. Cis-activation inhibits patterning and cis-inhibition facilitates it. (A) Stationary lateral inhibition pattern formed in an array of irregular cells in the absence of cis-interactions (r_c = 0). Grayscale is used to denote the ligand level (black for high ligand, l_1 = 1, and white for low ligand, l_1 = 0). (B–D) Regions of patterning for cis-interactions and trans-interactions strengths r_c,r_1 for (B) ɛ = 1 (cis-activation), (C) ɛ = 0 (cis-inhibition) and (D) ɛ = 0.4. The black dot-dashed line in D divides the parameter space into the cis-activation region, on its left, and the cis-inhibition region, on its right. Blue regions show where the pattern grows spontaneously (LSA in Methods). Green dashed lines enclose the regions where lateral inhibition pattern solutions exist and are stable (Exact periodic solutions in Methods). In B, the patterning region is the one below the green dashed line. B and D show that patterning becomes forbidden as r_c increases when cis-activation is acting. C and D show that patterning is enabled above a minimal cis-interactions strength r_c when there is cis-inhibition. Other parameter values: h = 4 for all panels, r_c = 1 and r_1 = 0 for A, h = 1000 for A–B and D, b = 10 for C.

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that the effect of cis-signaling on patterning can be simplified to that of the regime in which it is acting.

Cis-inhibition can modulate patterning and enhance multistability

We next evaluated which patterning features arise in the cis-inhibition regime. Cis-inhibition can make cells worse receivers of inhibition [14,20,26,57]. This is confirmed in our model by evaluating the change in the threshold level of ligand activity required to drive ligand inhibition in an adjacent cell when cis-interactions are added in the receiving cell (Methods, Fig. S4). When cells become worse receivers of inhibition we can expect the ratio of high-ligand expressing cells to increase. Simulations results confirm cis-inhibition can increase the ratio of selected precursor cells (Fig. 4A for \( \epsilon = 0 \) and Fig. S5 for \( \epsilon > 0 \)). The strength of cis-interactions \( r_c \) and the fraction between cis and trans-interactions strengths \( r_c/r_t \) become a control parameter for this ratio (Figs. 4B,C). When precursor cells exhibit large random initial variability between them in ligand and signal levels, the strength of cis-interactions can increase more gradually the ratio of high-ligand expressing cells (Figs. 4B–D).

These results showed that cis-inhibition can enable a new regular salt-and-pepper pattern with 66% of cells highly expressing the ligand (Figs. 4A, S5). This pattern has the periodicity of the lateral inhibition pattern. However, the ratio of selected high-ligand expressing cells is complementary to it. Since our results show that cis-inhibition can modulate the threshold for lateral inhibition and thereby the ratio of selected precursors, we wondered whether it can enrich patterning and drive additional periodic patterns. By using a combined analytical-computational approach (Methods), we searched across the parameter space of cis-interactions \( r_c \) and trans-interactions strengths \( r_t \) whether and where different periodic patterns composed of two cell types were stable solutions of the dynamics. We chose to search for three different types of patterns that involve different numbers of selected precursors and spatial organizations (Fig. S6). One of them is the salt-and-pepper pattern of Fig. 4A with a 66% ratio of selected precursors (Fig. S6A). Another one is a salt-and-pepper pattern too but with a different periodicity and a 75% ratio of selected precursors (Fig. S6B). The third chosen pattern is stripped with 50% of cells expressing high-ligand levels (Fig. S6B). Our study showed that high enough cis-interactions strengths \( r_c \) enable the emergence of these patterns with high numbers of precursor cells (Figs. 5, S7, S8 and S9). In the absence of cis-interactions and for low cis-interaction strengths \( r_c \rightarrow 0 \), the salt-and-pepper spatial organizations can persist but with much lower

![Figure 4. Cis-inhibiting interactions increase the ratio of high-ligand expressing cells.](https://example.com/figure4.png)

(A) Stationary patterns of ligand levels arising from precursor cells with small initial variability between them for different inhibiting cis-interactions strengths \( r_c \). Color code as in Fig. 3A. (B) Ratio of stationary high-ligand fated cells as a function of the cis-interactions strength \( r_c \) when precursor cells show small (red triangles) and large (blue circles) initial variability between them. (C) Density plot representing the ratio of high-ligand cells in a tissue arising from precursor cells exhibiting large initial variability. Solid and dashed lines as defined in Fig. 3B–D. White vertical line is drawn for indicating the value \( r_t \sim 3 \) along which simulations are performed in panels A, B and D. (D) Stationary patterns of ligand levels arising from precursor cells with large initial variability between them for different inhibiting cis-interactions strengths \( r_c \). In B–C panels, cells are considered high-ligand fated cells when its ligand level is over the threshold of 0.5. Parameter values: \( \epsilon = 0, b = 4, b = 1000 \) and \( v = 1 \) for all panels. Similar results are found for \( \epsilon = 0.4 \) (Fig. S5). In B, each point comes from the average of 10 numerical integrations of the dynamics on a lattice of \( 12 \times 12 \) irregular cells starting at different initial conditions. In C, the results correspond to numerical integration of the dynamics performed over a lattice of \( 30 \times 30 \) perfect hexagonal cells.

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numbers of selected precursors (33% and 25%, Figs. S7, S8). In contrast, cis-inhibition with high enough cis-interactions strengths ($r_c$) enable the spatial organization of precursor cells within stripes (Figs. 5, S9).

Additionally, we found that for high cis-interactions strengths ($r_c$) all these patterns are stable, i.e., there is multistability of pattern states (Figs. S7–S9). Hence, precursor cells could potentially become organized in any of them and should choose which pattern to form (Palau-Ortín et al., unpublished).

**Cis-inhibition allows localized patterning highly sensitive to the precursor state**

As shown in Fig. 4D, stable patterns without an obvious periodicity can arise too for high strengths of cis-inhibiting interactions ($r_c$). This occurs when precursor cells show large initial random variability between them, in agreement with [25].

We evaluated whether a high sensitivity to the initial state of precursor cells was causing the absence of periodicity. The results show that the finally formed stable pattern has strong similarity to the initial state of precursor cells (Fig. 6A–B). Simulations across the parameter space using precursor cells with large initial random variability between them confirmed that the pattern being formed is quite random as the state of precursors cells is for high cis-interactions strengths (Fig. S10). In contrast, large initial random variability between precursor cells dynamically evolves to more regular and periodic patterns of two cell fates for lower cis-interactions strengths (Fig. S10).

Simulations results indicated that the patterns exhibiting high sensitivity to the initial random state of precursor cells keep spatially localized without spreading to the rest of the tissue (Fig. 6C). This absence of spreading is in sharp contrast with the dynamics of nucleating patterns driven only by trans-interactions. Nucleating patterns invade the rest of the tissue that is under lateral inhibition Notch dynamics through a traveling wave [58,59]. The localized patterns we find remain where they arise performing cis-inhibition enrich patterning from precursor cells that show large initial variability between them. The arising patterns keep localized within the tissue and are reminiscent of the initial states of precursor cells.

**Cis-inhibition can drive cell-autonomous bistability**

We reasoned that cis-driven dynamics at the cell-autonomous level could be relevant for the phenomenon of localized patterning. Specifically, we wondered whether cis-inhibition could drive bistability of distinct ligand and signal level states in isolated cells. To evaluate it, we considered the role of cis-inhibition in the dynamics of single isolated cells that have a primary source of Notch signal ($\mu = \mu_0$, being $\mu_0$ a constant, see Text S1). This primary signaling source could be ligand-independent.

Our results show that cis-inhibition can drive cell-autonomous bistability when a primary signaling source is present (Fig. 7A,B, Methods). This bistability drives similar cell types to the ones found in lateral inhibition patterning: cells are expected to be either on a high-ligand expression state or in a low-ligand expression state with opposite signaling state (Fig. 7A,B). Bistability of cell fates requires a minimal amount of cis-interactions ($r_c$) and of primary signaling source ($\mu$; Fig. 7C).

We evaluated the existence of this cell-autonomous bistability when the amount of primary signal corresponds to the signal that trans-interactions drive for the homogeneous state of equivalent cells (Methods). For this primary signal, bistability arises for high cis-interactions strengths ($r_c$) and encloses the region where patterns keep localized (Fig. 6D). This result suggests that cell-autonomous bistable dynamics arising from cis-inhibition may be relevant for the phenomenon of non-periodic pattern localization.

**Discussion**

**Competition for signaling: a switch from cis-activation to cis-inhibition**

Several experimental evidences support the existence of signaling driven by cis-interactions [39–43]. In this work we have theoretically characterized the effect of cis-signaling in different contexts. We found that a switch from cis-activation to cis-inhibition (or vice versa) arises. The switch can occur by quantitatively changing the signaling sources; either by changing the amount of the primary signaling source (e.g., trans-interactions), or by modulating the ratio between the signaling efficiencies of each source. As a result, phenotypes involving a reduction of Notch signaling when the ligand is increased within a cell (cis-inhibition) can be compatible with cis-signaling.

The results show that cis-signaling can drive opposed capabilities to the patterning process, each arising on the different regulatory regimes of cis-activation and cis-inhibition. Cis-signaling acting as cis-activation creates a negative intracellular feedback loop that inhibits pattern formation. On the other side, cis-signaling acting as cis-inhibition creates a positive intracellular feedback loop facilitating patterning. This regime promotes patterning as cis-inhibition driven by null cis-signaling does [25,33,35].

The switch from cis-activation to cis-inhibition exemplifies a case of competition leading to a complex dynamical output: two signaling sources – the primary and the cis sources – with different efficiencies and competing for the same substrate – the Notch receptor – can result in a cis-productive signaling that drives cis-
inhibition. Such effect is reminiscent of the behavior of full and partial pharmacological agonists, where a partial agonist can act as a competitive inhibitor of a full agonist [60]. It is also an example of complex regulation in which the interplay between different components changes the regulation performed by one of them [61].

This notion can be extrapolated to any competition between signaling sources that share the same receptor. When different ligands (canonical or not) bind the same type of Notch receptor but drive signaling with different efficiencies, one can expect competition between them and switches of regulatory roles [62] (Jelena et al., unpublished). This approach could also help to decipher the roles of different canonical factors that bind to Notch and present both activatory and inhibitory effects on Notch signaling. For instance, this is the case of the Dll1/2 non-canonical ligands [3,63] and the proteins MAGP1/2 [45,64]. Different competition events have already been shown to drive significant regulatory effects in other signaling pathways (see for instance [65–70]).

In addition, our approach provides a definition for the ratio of signaling efficiencies coming from different sources that share the same receptor. This ratio is defined by the signaling rates and by the strength of the signaling sources. This definition is relevant to determine the regulatory role each signaling source drives on the overall signaling.

**Cis-inhibition as a modulator of the ratio of selected precursor cells.** From experimental grounds it has been already pointed out that cis-inhibition can drive cells to become worse receivers [14,20,26,57]. This effect can be obtained from our model too and enables cis-interactions to increase the number of high-ligand expressing cells. Cis-inhibition is expected also to drive cells to become worse signal senders by sequestering the ligand. Albeit this aspect is not considered in our simplified model, this effect should increase the ratio of cells reaching the high-ligand fate too. We have checked that a more complex model involving ligand sequestration similar to Sprinzak et al. (2010) (Methods and Text S1) also shows an increase in the ratio of high-ligand expressing cells. Consistent with our simplified model, competition between signaling sources in the complex model also yields switches of the regulatory role of cis-interactions (Fig. S11, Methods). The results confirm the increase in the ratio of high-ligand fated cells with the strength of cis-interactions in the cis-inhibition regime (Fig. S12). This complex model indicates that this increase occurs too when there is no cooperativity in the inhibition of the ligand (i.e. for h=1 in Eq. S4a of Text S1, Fig. S12).

It has been reported that Lunatic Fringe knockdown produces an increase in the neurogenesis ratio in the hindbrain of zebrafish embryos [71]. This could be an example of the cell-type ratio modulation we find in our simulations, since Fringe potentiates Delta-Notch trans-interactions [72] and could inhibit the cell-autonomous association of Delta and Notch [20].

The strength of cis-inhibiting interactions is not the only potential modulator of the ratio of high-ligand expressing cells. Specifically, different theoretical approaches that do not take cis-interactions into account have reported other components that can modulate this ratio through changes in the level of the threshold that drives inhibition of the ligand [73,74]. Also, it has been shown that higher rates of Delta production can drive a graded increase of high-Delta cells, which was validated experimentally [75].

Noteworthy, Notch signaling dynamics in *in vivo* can select a different ratio of high-ligand cells in different contexts [76–81]. Our results suggest that cis-inhibition could underlie the selection processes that involve high ratios of selected cells.

**Cis-inhibition potentiates multistability and enriches patterning.** Multistability can enable the change of fate of cells and it has been widely evaluated in the context of single cells, specially in bacteria and stem cells. A recurrent circuit topology that shows multistability and participates in stem cell renewal and differentiation is the toggle switch with auto-activation [82–84]. Auto-activation in this dynamics facilitates multistability [82]. Herein we show that lateral inhibition Notch dynamics with cis-

![Figure 6. Cis-inhibition allows pattern localization.](image)
inhibition can be described with an effective topology that corresponds to a toggle switch with auto-activation (Fig. 2F). In this effective topology, cis-inhibition drives auto-activation and facilitates multistability, in agreement with the effect of auto-activation in cell-autonomous toggle switches. This is reminiscent to the reported multistability in the ommatidia formation in Drosophila eye, which has been proposed to be driven by an auto-activatory feedback loop due to Atonal [85].

At the cell-autonomous level, we find that cis-signaling can drive bistability of ligand and signaling states when a basal cell-autonomous activity of Notch is present and cis-inhibition is acting. In this case, the effective circuit topology corresponds to a positive feedback loop that involves a mixed-feedback loop [86,87]. The bistability regime is confirmed and becomes more prominent in the more complex model (Fig. S13, Methods). The prediction of cell-autonomous bistability due to cis-inhibition could shed light to new functions of Notch in single cells. Recently, cell-autonomous bistability in Notch has been identified in the context of colon cancer stem cells [88]. In particular, it has been shown that the sequestering of mRNA Notch1 by the tumor suppressor microRNA miR-34a drives cells with Notch signal bimodality [88].

Simulation results show that cis-inhibition enables the spatial localization of patterns, which do not propagate spontaneously on the entire tissue. This could correspond to a wave-pinning phenomenon [58]. Typically, wave-pinning arises in discrete dynamical systems when the coupling between the discrete units is below a critical strength [99]. In our scenario, the critical coupling would be related to cis versus trans-interactions strengths (r_c/r_t ratios). Moreover, the most disordered patterns, with high-ligand cells adjacent to each other [25], appear in the region where localized patterning occurs. In such regions of the parameter space, the final pattern strongly depends on the initial precursor state. This dependence on the precursor state is reminiscent of the directionality provided by cis-inhibition in the differentiation of R1/R6/R7 precursor photoreceptor cells in the Drosophila eye [26]. Together, these results suggest that cis-inhibition can enrich patterning by enabling additional modulations of cell fate decisions.

**Methods**

Model formulation within an irregular cellular array layout

The simple model phenomenologically includes the competition between cis and the primary signaling sources for the Notch receptor and the inhibition of Notch signaling on the ligand. It is based on the approach introduced by Collier et al. [13] for lateral inhibition dynamics through trans-interactions. We included competition such that it is in agreement with a more biochemical reaction-based approach (see Complex model below). The simplicity of the simple model strongly facilitates the vast exploration of different patterning regions in the parameter space.

The weighted average of non-dimensional ligand concentration, 〈li〉, appearing in Eq. 1 through μ = r_t 〈li〉 due to trans-interactions, describes the interactions between adjacent cells on a two-dimensional irregular array of cells:

\[
\langle l_i \rangle = \sum_{j \neq \text{cell}(i)} x_{ij} l_j,
\]

with \(x_{ij} = e_{ij} / \sum_{j \neq \text{cell}(i)} e_{ij}\), being \(e_{ij}\) the length of the cell membrane edge shared by adjacent cells \(i\) and \(j\), and the summation involves all cells adjacent to cell \(i\) (\(j \neq \text{cell}(i)\)) [90]. We constructed an irregular two-dimensional array of cells with periodic boundary conditions as in [10] with irregularity parameter \(\gamma = 1.33\) (see Fig. S3 in [10]). First, we generated an irregular distribution of points on a plane starting from a perfect triangular lattice and considered periodic boundary conditions by surrounding the array of points with equivalent arrays. Second, a Voronoi tessellation was created around these points using Mathematica’s Computational Geometry Package (Wolfram Research, Inc. (2008), Mathematica, Version 7.0, Champaign, IL, USA).

Figure 7. Cis-inhibition with a primary Notch signaling source creates cell-autonomous bistability. (A) Stationary ligand level as a function of the cis-interactions strength \(r_c\), for \(\mu = \mu_0 = 10\) and \(\epsilon = 0\). Solid lines denote linearly stable solutions, dashed lines indicate linearly unstable solutions. Black dots refer to the stationary ligand levels for \(r_c = 100\). (B) Nullclines diagram showing the three possible solutions at \(r_c = 100\). The blue and red lines represent the nullclines. The continuous black line is a separatrix, which divides the parameter space into two basins of attraction of the two stable solutions. Percentages indicate the distance of the parameter space to these basins of attraction for\(\epsilon = 0\). (C) Phase diagram showing the cell-autonomous bistability region zone where two states are linearly stable. The gray area is the theoretically computed region, and the blue circles correspond to simulation results (Methods). Parameter values: \(h = 4, \ b = 1000, \ \epsilon = 0\) and \(v = 1\) for all panels. These results can also be obtained for \(\epsilon > 0\) in the cis-inhibition regime (data not shown).

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Complex model

The Complex model takes into account the dynamics of the Notch receptor and of the complexes formed by receptors and ligands (see Text S1 for all reactions, model equations and further details). It includes receptor and ligand inactivation through proteolytic cleavage [25,35] and it does not make assumptions regarding the capability of sending and receiving signals (e.g., when cis-interactions are acting, signal sending cells are not necessarily refractory to receive inhibitory signals from its neighbors). In this Complex model, the dynamics of the free receptor \( (N_i) \), trans and cis-formed receptor-ligand complexes \((C_i \text{ and } B_i \text{ respectively})\), and Notch signal \( S_i \) in cell \( i \) when the primary signaling source is due to trans-interactions read:

\[
\frac{dN_i}{dt} = -k_b N_i \langle L_i \rangle - k_{-b} N_i C_i + k'_{-b} B_i + \beta_n - v_n N_i \tag{7}
\]

\[
\frac{dC_i}{dt} = k_b N_i \langle L_i \rangle - k_{-b} C_i - k_s C_i - v_s C_i \tag{8}
\]

\[
\frac{dB_i}{dt} = k'_s S_i L_i - k'_{-b} B_i - v_b B_i \tag{9}
\]

\[
\frac{dS_i}{dt} = k_s C_i + k'_s B_i - v_s S_i \tag{10}
\]

where the variables are in dimensional units and \( L_i \) is the free ligand in cell \( i \), whose dynamics are detailed in Text S1. \( \langle L_i \rangle \) is given by Eq. 6 applied on species \( L \). \( \tau \) is the dimensional time. The binding and unbinding dynamics of Notch receptors with its ligand in trans and in cis have rates \( k_b, k_{-b}, \) and \( k'_s, k'_{-b}, \) respectively. Notch production \( \beta_n \) and degradation \( v_n \) are also taken into account. Trans and cis complexes, \( C_i \) and \( B_i \), have degradation rates \( v_s \) and \( v_b \) respectively. The model does not detail the overall mechanism by which trans-interactions drive signal activity. Instead, it assigns a rate \( k'_s \) to the proteolytic cleavage of the trans complex and the ultimate release of Notch signal \( S_i \). The model also considers the case in which cis-interactions drive Notch signaling. We implemented it by taking into account the argued mechanism for cis-signaling [40], so that the release of Notch intracellular domain would also occur for cis complexes \( (B_i) \). We set this step to occur at rate \( k'_s \). By taking \( k'_s = 0 \), the above equations account for the usual scenario of cis-interactions that sequester the receptor and drive no signaling.

Notice that the stationary solution of Eqs. 1–3 is the same function of \( L_i \) and \( \langle L_i \rangle \) as the stationary solution of Eqs. 7–10, which reads \[(dN_i/dt = 0, dC_i/dt = 0, dB_i/dt = 0, dS_i/dt = 0)\]:

\[
S_i^* = S_i^* \frac{r_i \langle L_i \rangle}{1 + r_i \langle L_i \rangle + r_l L_0} + S_i^* \frac{cr_i L_0}{1 + r_i \langle L_i \rangle + r_l L_0}, \tag{11}
\]

with \( S_i^* = k_b \beta_n / (v_b + v_s) \), \( r_l = k_b / v_b \), \( k_s + k'_{-b} + r_l \), \( 0 \leq r_l < 1 \), \( L_0 \) is a characteristic dimensionless concentration of ligand \( \langle L_i \rangle \). The first term on the right-hand side corresponds to the stationary primary signaling driven by trans-interactions whereas the second term is the stationary signaling driven by cis-interactions.

We define the efficiency of each source as the ratio of success to signal of the receptor-ligand complexes. This efficiency corresponds to \( k_s / (k_s + v_s) \) for the primary signaling source and to \( k'_s / (k'_s + v_b) \) for cis-interactions. Therefore, \( \epsilon \) parameter \( (\epsilon = k'_s / k'_s + v_b) \) is the relative efficiency of the cis-driven source compared to that of the primary signaling source. From Eq. 11 it can be seen that \( \epsilon \) corresponds as well to the ratio of maximal saturated stationary Notch activity driven by cis-interactions over that one driven by the primary signaling.

The equations of the Complex model for single isolated cells with ligand-independent and cell-autonomous primary signaling sources are detailed in Text S1.

Evaluation of the regulatory role of cis-interactions

We defined the regulatory role of cis-interactions (cis-inhibition or cis-activation) through the (negative or positive, respectively) change in Notch signal dynamics within a cell when its ligand content increases, \( R_{ni,i} \).

\[
R_{ni,i} = \frac{\partial S_i}{\partial L_i} = r_i (\epsilon + \mu (\epsilon - 1)) \quad (1 + \mu + r_i) \tag{12}
\]

where the result of the derivative for the model described by Eqs. 1–3 is indicated. Cis-inhibition is defined as a decrease in Notch signaling when the ligand content increases within the same cell, \( r_i < 0 \), whereas cis-activation corresponds to an increase in Notch signaling, \( r_i > 0 \). Based on the above expression for \( R_{ni,i} \), cis-interactions drive cis-inhibition when \( \epsilon + \mu (\epsilon - 1) < 0 \). Therefore, the condition for cis-inhibition to happen can be rewritten as inequality 5:

\[
\epsilon < \frac{\mu}{1 + \mu} \frac{r_i \langle L_i \rangle}{1 + r_i \langle L_i \rangle} \tag{13}
\]

where \( \mu = r_i \langle L_i \rangle \) for trans-interactions has been introduced in the last right-hand side term. This inequality states that cis-inhibition occurs when the maximal (saturated) signaling driven by cis-interactions is lower than the signaling driven by the primary source when acting alone (Fig. S1). Notice that this criterion for the cis-regulatory role is independent of \( r_i \) (Fig. S1C).

Eq. 13 gives the regulatory role of cis-interactions in the Complex model at the steady state as well \( R_{Si, L_0} = \frac{S_i^*}{S_i^*} \), with \( \epsilon = k'_s / k'_s + v_b \) and \( r_i = k_s / v_s \), \( k_s + k'_{-b} + r_l L_0 \). Accordingly, for no cis-signaling \( k'_s = 0 \) cis-interactions perform cis-inhibition, whatever the context and additional parameter values. For \( k'_s / k_s > v_b / v_s \), \( r_i > 1 \), cis-interactions perform always cis-activation. For \( 0 < k'_s / k_s < v_b / v_s \) \( r_i < 1 \), a switch from cis-activation to cis-inhibition can occur as the amount of trans-interactions increase. Noteworthy, when the receptor-ligand complex formed by cis-interactions is more unstable than the complex formed by trans-interactions \( v_b / v_s > 1 \), it can drive cis-inhibition even if it signals faster than the trans complex \( 1 < k'_s / k_s < v_b / v_s \).

From the model equations, it can be readily seen that the switch of regulatory role can only take place in the non-linear regime of the signaling function (Eqs. 1, 2 and 11). Notice that this regime does not require saturation of the Notch receptors. When the
primary source is acting in the linear regime (i.e. \( \mu < 1 \)), the addition of ligand within the cell does not reduce the primary signaling since there is no competition for the Notch receptor. As a result, cis-signaling always drives cis-activation in this linear regime.

The inequality arising for isolated cells with cis-signaling and a primary signaling source is detailed in Text S1 (Fig. S11B).

**Linear stability analysis (LSA)**

Linear stability analysis [15,56] has been applied to Eqs. 1–4 to determine in which regions of the parameter space spontaneous patterning occurs. We defined spontaneous patterning as the process that drives pattern formation from a linear instability of the homogeneous initial state through small non-homogeneous perturbations (i.e. when small initial variability between precursor cells becomes amplified) [91]. LSA enabled us to make analytic predictions of how the pattern formation capabilities of the system would be changed by cis-interactions. LSA indicated that cis-interactions do not change the fastest growing mode, and hence the periodicity of the pattern is expected to be the same as in the absence of cis-interactions for spontaneous patterning (Text S1). LSA indicated that spontaneous patterning in a regular hexagonal array of cells with periodic boundary conditions would happen when (see Text S1 for details and Fig. S14)

\[
1 < \mathcal{C} < \mathcal{A}/2,
\]

being

\[
\mathcal{B} = \frac{1}{\mathcal{C}} \frac{\partial l_i}{\partial t}_{|_{l_0}} , \quad \mathcal{C} = \frac{\partial s_i}{\partial t}_{|_{l_0}} , \quad \mathcal{A} = \frac{\partial s_i}{\partial t} \bigg|_{l_0} ,
\]

where \( l_i \) is the ligand level in a neighboring cell to cell \( i \), \( s_0 \) and \( l_0 \) are the homogeneous steady states for the multicellular system (i.e. the solutions of \( dl/dt = 0 \), \( ds/dt = 0 \) for \( \langle l_i \rangle = l_0 \)) and \( \omega \) is the number of nearest neighbors in a hexagonal cellular array (\( \omega = 6 \)). \( \mathcal{B} \) measures the strength of ligand repression at the homogeneous stationary state and verifies \( \mathcal{B} < 0 \). \( \mathcal{A} \) measures the strength of trans-activation and verifies \( \mathcal{A} > 0 \). Notice that \( \mathcal{C} = R_{h Bj} \) computed at the homogeneous steady state. Therefore, when cis-inhibition (\( R_{h Bj} < 0 \)) is acting at such homogeneous state then we have \( \mathcal{C} < 0 \). \( \mathcal{A} \) measures the strength of cis-inhibition (when \( \mathcal{C} < 0 \)) and of cis-activation (when \( \mathcal{C} > 0 \)).

According to inequality 14, cis-inhibition \( \mathcal{C} < 0 \) facilitates patterning (Fig. S13). In contrast, cis-activation \( \mathcal{C} > 0 \) inhibits patterning (Fig. S15). Inequality 14 was also used to evaluate where spontaneous patterning can emerge in the \( r_i-r_c \) parameter space. These results are depicted by solid lines in Figs. 3, 4 and 6. The use of the simple model of Eqs. 1–4 enabled a vast exploration across the parameter space. We checked several of the regions obtained by LSA with numerical simulations (Text S1, and Fig. S16 as example).

**Exact periodic solutions**

We evaluated the \( r_i-r_c \) parameter space regions where the lateral inhibition pattern is a stable solution of the dynamics defined by Eqs. 1–4. We also evaluated whether other periodic patterns are stable solutions of these dynamics. This analysis was strongly facilitated by the use of the simple model. To this end, we extended a method we previously introduced [56] to our system and to new periodic patterns. We considered periodic patterns composed of only two different cell types: cell type \( A \) and cell type \( B \). According to dynamics given by Eqs. 1–4, the stationary state values \( (dl/dt = 0, ds/dt = 0) \) of these two cell types are:

\[
l_A = \frac{1}{1 + b s_A^a} , \quad s_A = \frac{r_A \langle l_A \rangle + r_c l_A}{1 + r_i \langle l_A \rangle + r_c l_A} ,
\]

\[
l_B = \frac{1}{1 + b s_B^a} , \quad s_B = \frac{r_B \langle l_B \rangle + r_c l_A}{1 + r_i \langle l_B \rangle + r_c l_B} .
\]

Based on the periodicity of the pattern, we imposed which is the neighborhood of cell types each cell type interacts with:

\[
\langle l_A \rangle = m_A l_A + (1 - m_A) l_B , \quad \langle l_B \rangle = m_B l_B + (1 - m_B) l_A ,
\]

where \( m_{AB} \) is the ratio of \( A,B \)-like cells neighboring to the \( A,B \)-cell type. For the common lateral inhibition pattern we have \( m_{AB} = 0 \) and \( m_{BA} = 1/2 \). The \( r_i-r_c \) parameter boundary regions enclosing the region where this lateral inhibition pattern is a stable solution of the dynamics are depicted with dashed lines in Figs. 3, 4 and 6. The \( m_{AB} \) values for other periodic patterns are as follows. For an additional salt-and-pepper pattern (Fig. S6B) we have \( m_{AB} = 2/3 \). For the stripped pattern \( m_{AB} = 1/3 \) (Fig. S6B).

Solutions of Eqs. 16–18 were found using NSolve from Mathematica and also through custom made programs using the bisection method. Stability of solutions was evaluated computationally by numerical integration of the dynamics with patterned initial conditions (see below and Text S1). Together, these results show that the parameter space region where the pattern with the periodicity of the lateral inhibition pattern is stable is the largest one and contains the regions where the other patterns are stable (Figs. S7–S9).

**Threshold for lateral inhibition**

We evaluated how cis-interactions within a cell \( i \) change its capacity to receive the inhibition from adjacent cells. We termed \( \mathcal{I} \) the threshold for lateral inhibition and defined it as the ligand activity in adjacent cells that drives the inhibition of the ligand in cell \( i \). This inhibition of the ligand was defined as having a production rate of ligand activity \( \times \) times \( 0 < x < 1 \) the maximum production rate, which is 1 for Eq. 4. Therefore, according to Eq. 4, a production rate \( x \) of ligand occurs for \( s_i' = (1 - x)/(1 + b s_A^a) \). We then computed which ligand activity in adjacent cells \( \mathcal{I} \) is required to drive a stationary signaling \( s_i' \) within cell \( i \) if this cell has a ligand activity \( l_i \) which drives cis-interactions (i.e. we isolated \( \mathcal{I} \) from \( s_i' = \frac{s_i}{1 + \frac{l_i}{l_i}} \) according to Eqs. 1–3). Taken together we obtain that the threshold for lateral inhibition is:

\[
\mathcal{I} = \mathcal{I}_\text{trans} \left[ 1 + r_i l_i \left( \frac{1 - s_i'}{s_i'} \right) \right] ,
\]

where \( \mathcal{I}_\text{trans} = \frac{s_i}{r_i} \left( 1 - s_i' \right) \) is the threshold for lateral inhibition in the absence of cis-interactions. The above equation indicates that cis-interactions increase this threshold (\( \mathcal{I} > \mathcal{I}_\text{trans} \)) making cells worse receivers of inhibition when cis-inhibition is taking place (Fig. 4A–C).
Cis-driven cell-autonomous bistability

We evaluated whether the positive feedback generated by cis-inhibition with a competing primary signaling source in isolated cells (Fig. 2E) is sufficient to drive cell-autonomous bistability. To this end, we computed the steady state solutions \( \frac{dx_i}{dt} = 0 \), \( \frac{dP_i}{dt} = 0 \) for Eqns. 1-4 when \( \mu = \mu_0 \). Linear stability of these solutions was given by

\[
1 < \mathcal{R} \mathcal{E},
\]

where \( \mathcal{R} \) and \( \mathcal{E} \) are defined as in the LSA section in Methods but are evaluated at the steady state solutions. Analysis of the nullclines \( \frac{dx_i}{dt} = 0, \frac{dP_i}{dt} = 0 \) indicates that cis-inhibition is required to have bistability (both nullclines need to be decreasing functions). We extensively explored in the \( \mu_0-r_c \) parameter space where bistability occurred with LSA, and corroborated it with numerical simulations (Fig. 7C). In Fig. 6D we used \( \mu_0 = r_c \mu_0 \) being \( \mu_0 \) the homogeneous steady state for the multicellular system. All these simulations were performed for 400 cells with random initial conditions. The region of bistability in the phase space was delimited similarly than the LSA regions (Text S1). The stability of the bifurcation branches in Fig. 7A was checked with simulations of 900 cells with small fluctuations around the different branches.

Cell-autonomous bistability in the Complex model was analyzed through nullclines analysis and numerical integration of the dynamics. In this case, bistability arises too in the absence of cooperativity in ligand inhibition \( [h=1] \). Fixed points were computed with Mathematica with the NSolve function and with custom made programs with a bisection method.

Numerical integration of the dynamics

We integrated Eqns. 1–4 with custom made programs using a Runge-Kutta fourth-order algorithm [92] with a time step of \( dt = 0.1 \). The Complex model for the multicellular and single cell system was integrated with Mathematica by using the NDSolve function.

To evaluate which patterns were formed and their stability we used three types of initial conditions. (1) Precursor cells (i.e., cells at their initial condition) show small variability between them: for each molecular species \( x \), \( x_i(t=0) = \sigma_i (1 + \delta U^i - 0.5) \) being \( \sigma_i \) the homogeneous steady state for the species \( x \), \( U^i \) a uniform random number between 0 and 1 and \( \delta = 0.1 \). (2) Precursor cells show large random variability between them: \( x_i(t=0) = U^i \), unless otherwise stated. (3) Precursor cells already form a regular pattern, with small variability: \( x_i(t=0) = \sigma_i (1 + \delta U^i - 0.5) \) being \( \sigma_i \) the steady state pattern solution and \( \delta = 0.1 \). We performed numerical integration of the dynamics for different parameter values defined on a logarithmic mesh across the \( r_c-r_t \) parameter space. Simulations were stopped when the steady state was reached.

Supporting Information

Figure S1 The regulatory role of cis-interactions when acting together with trans-interactions. (A–C) Signal production rate in cell \( i \) due to (left) trans \( P_1 \) and (middle) cis-interactions \( P_2 \), and (right) total signal production rate \( P_1 + P_2 \) as a function of the ligand level \( l_i \) within the cell. (Left) \( P_1 \) decreases with \( l_i \) for all parameter values because cis-interactions drive competition for the Notch receptor. \( \langle l_i \rangle \) value is set at the homogeneous fixed point. (Right) \( P_2 \) always increases with \( l_i \) since we impose that cis-signaling on its own activates Notch signaling. (Right) \( P_1 + P_2 \) can be either a decreasing or an increasing function of \( l_i \). This indicates the regulatory role of cis-interactions. Cis-inhibition occurs when \( P_1 + P_2 \) decreases with \( l_i \), whereas cis-activation is acting when \( P_1 + P_2 \) increases with \( l_i \). (A) Signal production rates for different cis/trans signaling efficiencies \( \epsilon \): \( \epsilon = 0.1 \) (gray solid line), \( \epsilon = 0.5 \) (black solid line) and \( \epsilon = 0.9 \) (dashed line). The regulatory role changes from cis-inhibition to cis-activation as \( \epsilon \) increases. (B) Signal production rates for different trans-interactions strengths \( r_c \); \( r_c = 50 \) (gray solid line), \( r_c = 100 \) (black solid line) and \( r_c = 1000 \) (dashed line). The regulatory role changes from cis-activation to cis-inhibition as \( r_c \) increases. (C) Signal production rates for different cis-interactions strengths \( r_s \); \( r_s = 10 \) (dashed line), \( r_s = 100 \) (black solid line) and \( r_s = 250 \) (gray solid line). The regulatory role does not change with \( r_s \). This is because neither \( P_{2r} \) at \( l_i = 0 \) \( P_{2r} (l_i = 0) = r_s \langle l_i \rangle / (1 + r_s \langle l_i \rangle) \) nor the saturated value of \( P_{2r} \) \( P_{2rsat} = \epsilon \) depend on \( r_s \). Parameter values are \( h = 2 \), \( b = 1000 \), \( r_t = 250 \), \( r_c = 500 \) if not indicated otherwise. (B) \( \epsilon = 0.5 \) and (C) \( \epsilon = 0.9 \). (TIFF)

Figure S2 Results in the absence of cis-interactions \( (r_c = 0) \). (A) Scheme of interactions as in [13] of two cells that inhibit each other through Notch-mediated lateral inhibition. Black (blunt red) arrows denote activation (inhibition). Notice the positive intercellular feedback loop. (B–C) Phase diagrams in the parameter space of ligand inhibition strength \( b \) and trans-interactions strength \( r_t \) for (B) high \( [h=4] \) and (C) low \( [h=2] \) cooperativity in ligand inhibition. The blue region in (B) is where the homogeneous state is linearly unstable. This is the region of spontaneous patterning, where the lateral inhibition pattern can arise from the amplification of small differences between precursor cells, as described in [13]. The region above the dashed line is where the pattern solution (with the periodicity shown in Fig. 3A) is an exact stable solution of the dynamics [56]. Above the dashed line and below the blue line in panel B both the homogeneous state and the lateral inhibition pattern are stable solutions of the dynamics (i.e., it is a bistable region). The continuous and dashed lines in (B) have been shown in [10]. Spontaneous patterning does not occur at low cooperativities \( [h=2] \). (TIFF)

Figure S3 Phase diagrams in the \( r_c-r_t \) parameter space for cis-inhibition with different cooperativities in ligand inhibition. (A–C) Phase diagrams in the \( r_c-r_t \) parameter space for (A) no cooperativity \( [h=1] \), (B) low \( [h=2] \) and (C) high \( [h=4] \) cooperativity. (A) In the absence of cooperativity \( [h=1] \), a minimal amount of cis-interactions is required to create a pattern for any \( b \) value, being consistent with Sprinzak et al. (2010) [25]. (B) At low \( [h=2] \) cooperativity, cis-interactions enable spontaneous patterning. (C) At high cooperativity \( [h=4] \) cis-interactions can promote the bistable regions. In all panels, \( b = 1000 \) and \( \epsilon = 0.1 \). Color codes and line types as in Fig. S2. (TIFF)

Figure S4 cis-interactions in the cis-inhibition regime make cells worse receivers of inhibition. (A) Threshold for lateral inhibition \( P^* \) (Eq. 19 with \( x = 0.01 \) and \( l_i = 0.5 \)) as a function of the trans-interactions strength \( r_t \) for \( \epsilon = 0 \). Results for different cis-interactions strengths are depicted: \( r_c = 0 \) (solid line), \( r_c = 10 \) (gray dashed line) and \( r_c = 100 \) (dotted-dashed gray line). The vertical line is a guide to the eye for a particular trans-interactions strength value, to better appreciate the rise of \( P^* \) due to cis-interactions strength. (B–C) Contour lines for different \( P^* \) values are depicted across the \( r_t-r_c \) parameter space for (B) \( \epsilon = 0 \) and (C) \( \epsilon = 0.4 \). Lines are depicted for \( P^* = 0.1 \) (long-dashed), 0.5, 0.7, 1.0 (short-dashed). As a guide to the eye, the spontaneous pattern
Cis-inhibiting interactions increase the ratio of high-ligand expressing cells at $\epsilon = 0.4$. Simulation results showing patterns of ligand levels from precursors with large initial variability between them for different cis-interactions strengths ($r_1$). Grayscale is used to denote the ligand level (black for the highest ligand activity, $l_1 = 0.035$, and white for no ligand activity, $l_1 = 0$). Other parameter values are $r_1 = 1000$, $b = 1000$, $h = 4$ and $\nu = 1$.

Figure S5 Cis-inhibiting interactions increase the ratio of high-ligand expressing cells at $\epsilon = 0.4$. Simulation results showing patterns of ligand levels from precursors with large initial variability between them for different cis-interactions strengths ($r_1$). Grayscale is used to denote the ligand level (black for the highest ligand activity, $l_1 = 0.035$, and white for no ligand activity, $l_1 = 0$). Other parameter values are $r_1 = 1000$, $b = 1000$, $h = 4$ and $\nu = 1$.

Figure S6 Representation of periodic patterns composed of two cell types on a regular hexagonal array. (A) Salt-and-pepper patterns with the periodicity of the fastest growing mode ($\Omega = -1/2$, see LSA in Text S1). The P and I patterns have the same periodicity but 33% and 66% of cells, respectively, are high-ligand expressing cells (black). (B) Patterns with the periodicities of the secondary fastest growing modes ($\Omega = -1/3$, see LSA in Text S1). P2 and I2 are salt-and-pepper patterns too with 25% and 75% of high-ligand expressing cells respectively. The pattern of stripes (S) has 50% of cells with high ligand levels. On the right of each row of patterns, two groups of 6 cells neighboring a central cell illustrate how many neighboring cells are like the central one and how many are different. Each group has a different cell type on the center (cell type B in violet and cell type $A$ in green). Notice that cell types $A$ and $B$ are defined by the $m_A$ and $m_B$ values (Methods) and not by their ligand level. These illustrations facilitate the computation of $m_A$ and $m_B$ values of Eqs. 16–18 for each pattern.

Figure S7 Cis-inhibiting interactions enable the salt-and-pepper pattern with 66% of cells highly expressing the ligand. (A) Phase diagram showing where patterns (green $P$ and red $I$) are each a stable solution of the dynamics to small perturbations. Green dashed and blue lines as in Fig. S3C. (B) Bifurcation diagrams for each cell type, $A$ and $B$, for $r_1 = 3$. The periodic solutions are shown in blue. The homogeneous solution is shown in gray. Solid (dashed) lines correspond to linearly stable (unstable) states. At low cis-interactions strengths, the stable branches correspond to $P$ (light blue) and at higher cis-interactions strengths to $I$ (dark blue). Note that there is a large parameter region in which both patterns are stable. Solutions for patterns were found by solving Eqs. 16–18 with $m_A = 0$ and $m_B = 2/3$. Stability of solutions was evaluated through numerical simulations of the dynamics (Text S1). Parameter values: $\epsilon = 0$, $h = 4$, $b = 1000$ and $\nu = 1$.

Figure S8 Cis-inhibiting interactions enable the salt-and-pepper pattern with 75% of cells highly expressing the ligand. (A) Phase diagram showing where patterns (green $P_2$ and red $I_2$) are each a stable solution of the dynamics to small perturbations. Green dashed and blue lines as in Fig. S3C. (B) Bifurcation diagrams for each cell type, $A$ and $B$, for $r_1 = 3$. The periodic solutions are shown in blue. The homogeneous solution is shown in gray. Solid (dashed) lines correspond to linearly stable (unstable) states. At low cis-interactions strengths, the stable branches correspond to $P_2$ (light blue) and at higher cis-interactions strengths to $I_2$ (dark blue). Note that there is a large parameter region in which both patterns are stable. Solutions for patterns were found by solving Eqs. 16–18 with $m_A = 0$ and $m_B = 2/3$. Stability of solutions was evaluated through numerical simulations of the dynamics (Text S1). Parameter values: $\epsilon = 0$, $h = 4$, $b = 1000$ and $\nu = 1$.

Figure S9 Cis-inhibiting interactions enable the stripped pattern with 50% of cells highly expressing the ligand. (A) Phase diagram showing where pattern (green) $S$ (as defined in Fig. S6) is a stable solution of the dynamics to small perturbations. Green dashed and blue lines as in Fig. S3C. (B) Bifurcation diagrams for each cell type, $A$ and $B$, for $r_1 = 3$. The periodic patterned solution is shown in blue. The homogeneous solution is shown in gray. Solid (dashed) lines correspond to linearly stable (unstable) states. Since type $A$ and type $B$ cells are equivalent for this pattern, there is bistability of the stripes solution. In all the parameter region where the stripped pattern is stable there are several patterns ($P$, $I$, $P_2$ or $P_2$) that are stable too (Figs. S7, S8). Solutions for patterns were found by solving Eqs. 16–18 with $m_A = 1/3$ and $m_B = 1/3$. Stability of solutions was evaluated through numerical simulations of the dynamics (Text S1). Parameter values: $\epsilon = 0$, $h = 4$, $b = 1000$ and $\nu = 1$.

Figure S10 Sensitivity to initial conditions occurs at high cis-interactions strengths in the cis-inhibition regime. (A–C) Stationary patterns of ligand level emerging from precursors with large initial random variability between them. (D–F) Structure function (Text S1) of the patterns in A–C respectively, without the homogeneous mode ($\Omega = 1$). (A,D) The disordered pattern that emerges at high cis-interactions strengths ($r_1 = 1000$), where the homogeneous solution is linearly stable. (B,E) Regular pattern that emerges at lower cis-interactions strengths ($r_1 = 200$), where the homogeneous solution is linearly stable. In these panels we set $r_1 = 50$, (C,F) Salt-and-pepper periodic pattern that emerges within the spontaneous pattern formation region, for very low cis-interactions strengths ($r_1 = 0.1$ and $r_1 = 100$). Other parameter values are $\epsilon = 0$, $h = 4$, $b = 1000$ and $\nu = 1$.

Figure S11 The switch between cis-activation and cis-inhibition regulatory roles also occurs in the Complex model. Stationary Notch signal in cell $i$, $E_{i1}$, versus the amount of free ligand in the cell, $L_i$, and the primary signaling source for (A–C) the multicellular system ($K_i (L_i)$ with $K_i = k_b / k_{-b}$) and (D–F) the single cell system ($K_i = k_0 / \mu_{-a}$) for (A,D) null ($K_i = 0$), (B,E) slow ($k_0 = 0.1k_b$ in $B$, and $k_0 = 0.05k_b$ in $E$), and (C,F) fast $k_0 = k_b$ cis-signaling. The value of $\epsilon$ is (A,D) $\epsilon = 0$, (B) $\epsilon = 0.4$, ($E_1$) $\epsilon = 0.24$ and (C,F) $\epsilon = 1$. Red lines show the dependence of $S_i$ on $L_i$ when there is no primary source and when it is maximal on the plot. An increasing function denotes cis-activation, while a decreasing function corresponds to cis-inhibition. A,D ($\epsilon = 0$) show cis-inhibition; B,E (\epsilon = 0.4) and (\epsilon = 0.24) show a switch from cis-activation to cis-inhibition as the primary source increases; D,F (\epsilon = 1) show cis-activation.

Figure S12 Cis-inhibiting interactions promote higher ratios of high-ligand expressing cells in the Complex model. Stationary patterns reached by numerical integration of the dynamics for different cis-interactions strengths as measured through the cis-binding rates $k_b$ values below the panels (in au$^{-1}$ hr$^{-1}$ units). Ligand levels are represented in grayscale (black for
250 and white for 0). Lower cis-binding affinities (\(k^b_0 = 400\)) allow high-ligand expressing cells next to each other [25]. Higher cis-affinities drive a gradual increase of the ratio of ligand-positive cells in the tissue. Herein this phenomenology occurs even in the absence of cooperativity (\(h = 1\)). Parameter values are in the cis-inhibition regime: \(k^b_1 = 0\), \(\beta^b_0 = 20\) au hr\(^{-1}\), \(\beta^b_1 = 5\) au hr\(^{-1}\), \(k^b = 25\) au hr\(^{-1}\), \(k^b_0 = 0.1\) au hr\(^{-1}\), \(v^b_1 = 0.1\) hr\(^{-1}\), \(v^b_0 = 0.031\) hr\(^{-1}\), \(v^b = 2\) hr\(^{-1}\), \(v^b_k = 0.1\) hr\(^{-1}\), \(b^b_k = 1000\) au, \(h = 1\). hr refers to hours and au refers to arbitrary concentration units. Precursor cells (initial conditions) were set as \(L^b_i(t=0) = U^b_i/\beta_i/v^b_i\) and \(N^b_i(t=0) = U^b_i/\beta_i/v^b_i\) where \(U^b_i\) is a uniform random number between 0 and 1, and the remaining variables were set to 0.

**Figure S13** Cis-inhibition with a primary Notch signaling source can create cell-autonomous bistability in the Complex model. Representation of relations S9a–S9b in the materials/analysis tools: PFJ MI. Wrote the paper: PFJ MI. Conceived and designed the experiments: PFJ MI. Performed the experiments: PFJ MI. Ares and Siheru Kondo for fruitful discussions.

**References**

1. Artavanis-Tsakonas S, Rand M, Lake R (1999) Notch signaling: cell fate control and signal integration in development. Science 284: 770.

2. Schwanbeck R, Martin S, Bernoth K, Just U (2011) The notch signaling pathway: molecular basis of context dependency. Eur J Cell Biol 90: 572–81.

3. Andersson ER, Sandberg R, Lundahl U (2011) Notch signaling: simplicity in design, versatility in function. Development 138: 3593–612.

4. Hor i, Sen A, Artavanis-Tsakonas S (2013) Notch signaling at a glance. J Cell Sci 126: 2133–40.

5. Fedor R, Koch P, Rebay I, Regier C, Xu T, et al. (1996) Molecular interactions between the protein products of the neurogenic loci notch and delta, two egf-homologous genes in drosophila. Cell 61: 523–534.

6. Kopan R, Ilagan MXG (2009) The canonical notch signaling pathway: unfolding the activation mechanism. Cell 137: 216–33.

7. Chithis A, Henrique D, Lewis J, Ish-Horowicz D, Kintner C (1995) Primary neurogenesis in Xenopus embryos regulated by a homologue of the drosophila neurogenic gene delta. Nature 375: 761–766.

8. Hadden C, Smithers I, Schneider-Maunoury S, Coche T, Henrique D, et al. (1998) Multiple delta genes and lateral inhibition in zebrafish primary neurogenesis. Development 125: 359.

9. Kageyama R, Ohtsuka T, Shimojo H, Imayoshi I (2008) Dynamic notch signaling in neural progenitor cells and a revised view of lateral inhibition. Nat Neurosci 11: 1247–1251.

10. Formosa-Jordan P, Baetes M, Ares S, Frade JM (2012) Regulation of neuronal differentiation at the neurogenic wavefront. Development 139: 2321–9.

11. Formosa-Jordan P, Baetes M, Ares S, Frade JM (2013) Lateral inhibition and neurogenesis: novel aspects in motion. Int J Dev Biol 57: 341–50.

12. Heidler P, Simpson P (1991) The choice of cell fate in the epidermis of drosophila. Cell 64: 1083–1092.

13. Collier JR, Monk NA, Maini PK, Lewis JH (1996) Pattern formation by lateral inhibition with feedback: a mathematical model of delta-notch intercellular signalling. J Theor Biol 183: 429–46.

14. Jacobsen TL, Brenman K, Ares AM, Muskavitch MA (1998) Cis-interactions between delta and notch modulate neurogenic signalling in drosophila. Development 125: 4531–40.

15. Cordle J, Johnson S, Tay JZY, Roversi P, Wilkin MB, et al. (2008) A conserved face of the jagged/serrate dsl domain is involved in notch trans-activation and cis-inhibition. Nat Struct Mol Biol 15: 849–857.

16. Fiuza UM, Klein T, Arias AM, Hayward P (2010) Mechanisms of ligand-mediated inhibition in notch signaling activity in drosophila. Dev Dyn 239: 798–805.

17. Becam I, Fiuza UM, Arias AM, Milà M (2010) A role of receptor notch in ligand-cis-inhibition in drosophila. Curr Biol 20: 554–60.

18. Fleming RJ, Hor i, Sen A, Filoramo GV, Langer JM, et al. (2013) An extracellular region of serrate is essential for ligand-induced cis-inhibition of notch signaling. Development 140: 2039–49.

19. Heidler P, Simpson P (1993) Altered epidermal growth factor-like sequences provide evidence for a role of notch as a receptor in cell fate decisions. Development 117: 1113–1113.

20. Sakamoto K, Ohsawa T, Takagi M, Takeda S, Katsube K (2002) Intracellular cell-autonomous association of notch and its ligands: A novel mechanism of notch signal modification. Dev Biol 241: 313–326.

21. Laci E, Nicholls JT, Ge W, Miyamoto A, Yao C, et al. (2005) The divergent dsl ligand dll does not activate notch signaling but cell autonomously attenuates signaling induced by other dsl ligands. J Cell Biol 170: 983–92.

22. Gittenberg M, Pitsouli C, Garvey C, Delidakis C, Bray S (2006) Role of conserved intracellular motifs in serrate signalling, cis-inhibition, and endocytosis. EMBO J 25: 4697–4706.

23. Matuda M, Chitnis AB (2008) Interaction with notch determines endocytosis of specific delta ligands in zebrafish neural tissue. Development 135: 197–206.

24. del Alamo D, Schwesiguth F (2009) Notch signalling: Receptor-cis-activation to achieve directionality. Curr Biol 19: R833–R834.

25. Sprinzak D, Lakhanpal A, Lebon L, Santat LA, Fontes ME, et al. (2010) Cis-interactions between notch and delta generate mutually exclusive signalling induced by other dsl ligands. J Cell Sci 170: 983–92.

26. Gitter Y, Miles R, Kanapin A, Elkon RB, Patel AP, et al. (2011) The nuclear notch signaling pathway: a conserved cis-inhibition mechanism. Nat Immunol 12: 13–14.

27. Miller AC, Lyons EL, Herman TG (2009) cis-inhibition of notch by endogenous delta biases the outcome of lateral inhibition. Curr Biol 19: 1378–83.

28. del Alamo D, Rosaulin H, Schwesiguth F (2011) Mechanism and significance of cis-inhibition in notch signalling. Curr Biol 21: R80–7.

**Figure S14** Cell labeling scheme. Arrays of \(N \times M\) perfect hexagonal cells with the subindex labeling schemes used that number each cell along the array. In (A) one subindex is used, while in (B) we use two subindices. The two main spatial directions of the cellular array are depicted.

**Figure S15** Decomposition of the elements determining the linear stability of the homogeneous state. Parameter values as in Fig. 3D (\(\varepsilon = 0.4, h = 1000\) and \(h = 4\)). (A) Strength of trans-activation \(\alpha\) across the \(r_t-r_0\) parameter space. (B) Strength of ligand repression \(\beta\). (C) Strength of cis-inhibition (for \(\varepsilon < 0\)) and of cis-activation (for \(\varepsilon > 0\)). Trans-interactions strengths (\(r_t\)) are crucial for determining the cis-role at intermediate cis-signaling efficiencies. Cis-inhibition promotes spontaneous patternning at high cis-interactions strengths (Eq. 14 in Methods). In each panel, color codes are detailed on the color bar.

**Figure S16** Simulation results agree with the spontaneous pattern formation regions predicted from LSA. Phase diagram in the \(r_t-r_0\) parameter space for \(\varepsilon = 0.4\) as in Fig. 3D, with the blue triangles indicating the boundaries of the spontaneous pattern formation regions computed from simulation results (Text S1). Other parameter values as in Fig. S3C.

**Text S1** Supplementary text contains more detailed aspects on the models and on the analytical and computational tools being used.

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Conceived and designed the experiments: PFJ MI. Performed the experiments: PFJ MI. Analyzed the data: PFJ MI. Contributed reagents/materials/analysis tools: PFJ MI. Wrote the paper: PFJ MI.
57. Held W, Mariuzza RA (2011) Cis-trans interactions of cell surface receptors:

52. Mukherjee T, Kim WS, Mandal L, Banerjee U (2011) Interaction between

51. Yamada K, Fuwa TJ, Ayukawa T, Tanaka T, Nakamura A, et al. (2011) Roles

50. Cho B, Fischer JA (2011) Ral gtpase promotes asymmetric notch activation in

49. Fortini M (2009) Notch signaling: the core pathway and its posttranslational

47. Sanders PGT, Mun˜oz-Descalzo S, Balayo T, Wirtz-Peitz F, Hayward P, et al.

46. Wilkin M, Tongngok P, Gensch N, Clemence S, Motoki M, et al. (2008)

43. Guy CS, Vignali KM, Temirov J, Bettini ML, Overacre AE, et al. (2013)

42. Hsieh EH, Lo DD (2012) Jagged1 and notch1 help edit m cell patterning in

40. Fü¨rthauer M, Gonza ´lez-Gaita´n M (2009) Endocytic regulation of notch

39. Coumailleau F, Fü¨rthauer M, Knoblich JA, González-Gaitáñ M (2009) Directional
delta and notch trafficking in sar endosomes during asymmetric
cell division. Nature 450: 1051–1053.

36. Wang R, Liu K, Chen L, Aihara K (2011) Neural fate decisions mediated by

35. Sprinzak D, Lakhanpal A, Lebon L, Garcia-Ojalvo J, Elowitz MB (2011) Mutual

dynamics for pattern formation. J Stat Mech 03: 019.

15: 221–231.

dynamics of regulatory and signaling pathways in the cell. Curr Opin Cell Biol

13: 1565.

52. Mukherjee T, Kim WS, Mandal L, Banerjee U (2011) Interaction between

51. Yamada K, Fuwa TJ, Ayukawa T, Tanaka T, Nakamura A, et al. (2011) Roles

50. Cho B, Fischer JA (2011) Ral gtpase promotes asymmetric notch activation in

49. Fortini M (2009) Notch signaling: the core pathway and its posttranslational

47. Sanders PGT, Mun˜oz-Descalzo S, Balayo T, Wirtz-Peitz F, Hayward P, et al.

46. Wilkin M, Tongngok P, Gensch N, Clemence S, Motoki M, et al. (2008)

43. Guy CS, Vignali KM, Temirov J, Bettini ML, Overacre AE, et al. (2013)

42. Hsieh EH, Lo DD (2012) Jagged1 and notch1 help edit m cell patterning in

40. Fü¨rthauer M, Gonza ´lez-Gaita´n M (2009) Endocytic regulation of notch

39. Coumailleau F, Fü¨rthauer M, Knoblich JA, González-Gaitáñ M (2009) Directional
delta and notch trafficking in sar endosomes during asymmetric
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36. Wang R, Liu K, Chen L, Aihara K (2011) Neural fate decisions mediated by

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dynamics for pattern formation. J Stat Mech 03: 019.

15: 221–231.

dynamics of regulatory and signaling pathways in the cell. Curr Opin Cell Biol

13: 1565.

52. Mukherjee T, Kim WS, Mandal L, Banerjee U (2011) Interaction between

51. Yamada K, Fuwa TJ, Ayukawa T, Tanaka T, Nakamura A, et al. (2011) Roles

50. Cho B, Fischer JA (2011) Ral gtpase promotes asymmetric notch activation in

49. Fortini M (2009) Notch signaling: the core pathway and its posttranslational

47. Sanders PGT, Mun˜oz-Descalzo S, Balayo T, Wirtz-Peitz F, Hayward P, et al.

46. Wilkin M, Tongngok P, Gensch N, Clemence S, Motoki M, et al. (2008)

43. Guy CS, Vignali KM, Temirov J, Bettini ML, Overacre AE, et al. (2013)

42. Hsieh EH, Lo DD (2012) Jagged1 and notch1 help edit m cell patterning in

40. Fü¨rthauer M, Gonza ´lez-Gaita´n M (2009) Endocytic regulation of notch

39. Coumailleau F, Fü¨rthauer M, Knoblich JA, González-Gaitáñ M (2009) Directional
delta and notch trafficking in sar endosomes during asymmetric
cell division. Nature 450: 1051–1053.

36. Wang R, Liu K, Chen L, Aihara K (2011) Neural fate decisions mediated by

35. Sprinzak D, Lakhanpal A, Lebon L, Garcia-Ojalvo J, Elowitz MB (2011) Mutual

dynamics for pattern formation. J Stat Mech 03: 019.

15: 221–231.

dynamics of regulatory and signaling pathways in the cell. Curr Opin Cell Biol

13: 1565.