Basal Protrusions Mediate Spatiotemporal Patterns of Spinal Neuron Differentiation

Graphical Abstract

Highlights

- Embryonic spinal neurons elaborate longitudinal basal processes prior to axogenesis
- Basal processes reach several cell diameters and last a few hours before retraction
- Basal processes express high levels of Delta protein
- Experiments and theory show processes mediate transient long-range lateral inhibition

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In Brief
Lateral inhibition mediated by Delta-Notch signaling is thought to operate at short range between neighboring cells. Hadjivasiliou et al. show that transient protrusions up to 100 μm long from differentiating neurons extend the range of lateral inhibition and regulate spatiotemporal patterns of neuronal differentiation along the spinal cord.
Basal Protrusions Mediate Spatiotemporal Patterns of Spinal Neuron Differentiation

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SUMMARY

During early spinal cord development, neurons of particular subtypes differentiate with a sparse periodic pattern while later neurons differentiate in the intervening space to eventually produce continuous columns of similar neurons. The mechanisms that regulate this spatiotemporal pattern are unknown. In vivo imaging in zebrafish reveals that differentiating spinal neurons transiently extend two long protrusions along the basal surface of the spinal cord before axon initiation. These protrusions express Delta protein, consistent with the hypothesis they influence Notch signaling at a distance of several cell diameters. Experimental reduction of Laminin expression leads to smaller protrusions and shorter distances between differentiating neurons. The experimental data and a theoretical model support the proposal that neuronal differentiation pattern is regulated by transient basal protrusions that deliver temporally controlled lateral inhibition mediated at a distance. This work uncovers a stereotyped protrusive activity of newborn neurons that organize long-distance spatiotemporal patterning of differentiation.

INTRODUCTION

During the early stages of vertebrate neurogenesis, neurons of particular subtypes initially differentiate along the spinal cord with a sparse periodic pattern but eventually produce more continuous columns of similar neurons (Figure 1A; Dale et al., 1987; Roberts et al., 1987; Higashijima et al., 2004a, 2004b; Kimura et al., 2006; Batista et al., 2008; England et al., 2011). The mechanisms that regulate this pattern of differentiation are unknown. Delta-Notch-mediated lateral inhibition is a regulator of vertebrate neurogenesis (Chitnis et al., 1995; Henrique et al., 1997; Appel et al., 2001; Okigawa et al., 2014), but this conventionally operates in a juxtacrine fashion between Delta-expressing cells and their immediate neighbors and cannot explain the spatial and temporal pattern of neuronal differentiation along the embryo spinal cord. Recent evidence, however, suggests the distance over which contact mediated signaling of various types can operate can be extended by cellular protrusions capable of spanning several cell diameters (reviewed in Buszczak et al., 2016; Pröls et al., 2016). For example, signaling through long cellular protrusions plays a role during limb patterning in the chick embryo (Sanders et al., 2013), in the development of zebrafish pigmentation stripes (Eom et al., 2015), and in neural plate patterning in the zebrafish (Stanganello et al., 2015). In fact, dynamic cellular protrusions from the basal surface of sensory organ precursor (SOP) cells have been proposed to mediate long-distance lateral inhibition to regulate the sparse distribution of mechanosensory bristles in the fly notum and wing disk (De Joussineau et al., 2003; Cohen et al., 2010; Hadjivasiliou et al., 2016; Hunter et al., 2016, 2019). Whether similar protrusive activity mediates long-distance spacing patterns in the vertebrate central nervous system (CNS) is not known, but long and short cellular protrusions expressing the Notch ligand Delta-like 1 have been described on intermediating progenitors in the embryonic mammalian cortex (Nelson et al., 2013). Furthermore, dynamic protrusive activity on the surface of recently born spinal neurons can be observed in slice cultures of chick embryo spinal cord (Das and Storey, 2014).

To determine whether cellular protrusions could also play a role in the patterning of spinal neuronal differentiation, we addressed these issues in the zebrafish embryo spinal cord. Live in vivo imaging revealed all spinal neurons transiently extend two long cellular protrusions along the basal surface of the spinal cord prior to axon initiation and apical detachment. We show these long basal protrusions express Delta protein at high level and Notch reporter activation is upregulated in cells in their vicinity. Furthermore, experimental reduction of the basal protrusion length results in reduced spacing between differentiating neurons. Our in vivo data are supported by a theoretical model, whose output is consistent with the hypothesis that neuronal differentiation is regulated by lateral inhibition mediated at a...
distance by transient basal protrusions. Our work thus reveals a
distinctive protrusive activity of differentiating neurons that
organizes long-distance spatiotemporal patterning of neuronal
differentiation in the embryo spinal cord.

RESULTS

Differentiating Spinal Neurons Transiently Elongate
Two Long Basal Protrusions along the A/P Axis before
Detaching from the Apical Surface
To study the early phases of neuronal differentiation in vivo, we
labeled small numbers of cells in the zebrafish embryo spinal
cord by mosaic expression of membrane-GFP and captured their
behavior with confocal time-lapse microscopy from 18 to
42 h post fertilization (hpf). Analysis of more than 100 cells
that differentiate into neurons reveals a stereotyped, transient
T-shaped transition from a cell that is attached to the apical
surface of the neuroepithelium to a basally positioned neuron
with the beginnings of a single axon extension. This transition
involves the elongation of two longitudinally directed cellular pro-
cesses that protrude along the basal surface of the neural tube,
one protruding anteriorly and the other posteriorly (Figure 1B,
time point 1h42 and 3h18; Figures 1B and 1C; Video S1).

Figure 1. Differentiating Spinal Neurons Transiently Elongate Two Long Basal Prom-
trusions along the A/P Axis before Detaching from the Apical Surface
(A) Diagram to show spinal neurons differentiate with an initial long-distance spacing pattern (t1). Later differentiating neurons of the same type subsequently fill in the gaps between the earlier differentiated cells (t2 and t3) to generate a near continuous column of neurons. Lateral view of spinal cord, dorsal to top.

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These basal protrusions can be asymmetric in length (17 out of

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28 cells) and each protrusion can reach up to 109 μm (mean ± SD, 42.6 ± 20.2 μm, n = 24 cells) with a combined length of up to 151.5 μm (mean ± SD, 86.8 ± 25.3 μm, n = 21 cells) (Figures 1D and S1). The basal protrusions are typically present on differentiating neurons for several hours (mean ± SD, 6.8 ± 2.2 h, n = 13 cells) and grow on average 6× longer than the basal extensions formed by the non-differentiating neural progenitors (mean ± SD, 14.3 ± 6.2 μm, n = 74 cells) (Figure 1D). After reaching their maximum length, basal protrusions begin to retract back to the cell body, and this is followed by the detachment and retraction of the apical process (19 out of 24 cells) (Figure 1B, from time point 3h18 to 5h42; Figure 1E). In a few cases (5 out of 24 cells) the apical detachment preceded the retraction of basal protrusions. Although apical and basal process retraction occurs at roughly the same time they do not appear to be strictly synchronized, suggesting they may be independent of one another. After these three processes have retracted, cells adopt a near spherical shape and the cell body becomes highly enriched in filopodial activity that diminishes prior to axon formation (23 out of 27 cells) (Figure 1B, time point 6h30 and 8h54; Figure 1B′; Video S1). The transient basal protrusions contain dynamic microtubules (Figure S2A) and often produce filopodia that are directed radially toward the apical surface (Figure S2B). Basal protrusions from nearby differentiating cells can overlap (Figures S2C–S2C′).

Differentiating spinal neurons thus stereotypically adopt a transient T shape prior to apical detachment and axon formation (summarized in Figure 1B′). These observations reveal a new in vivo cellular behavior that precedes axogenesis and distinguishes the neuronal precursors in the process of differentiation from surrounding neural progenitors.

**Stereotyped Axon Formation Follows Basal Protrusion Retraction**

Studies of neuronal differentiation in vitro have revealed that axons derive by selection and specialization of one neurite from several pre-existing neurites (Dotti et al., 1988; Craig and Banker, 1994; Barnes, and Polleux, 2009). To investigate whether the axons of spinal neurons in vivo might derive from the transient long basal protrusions, we monitored axon initiation. Neurons were located at many different dorsoventral (D/V) levels of spinal cord and thus likely represent many different subtypes of spinal projection neuron. Our 3D reconstruction analyses revealed that axonal outgrowth almost always follows the full retraction of basal protrusions (27 out of 31 cells) (Figures 2A and 2B; Videos S1 and S2), and in contrast to in vitro observations, axons never differentiated from an existing cellular protrusion. The majority of subtypes of spinal neurons have an axon that runs ventrally and circumferentially from the cell body before either crossing the ventral floor plate or turning anteriorly or posteriorly to join the ipsilateral longitudinal axon tracts (Bernhardt et al., 1990). Our observations show that this ventral circumferential axon trajectory is initiated stereotypically at the outset of axon growth, directly from the cell body and is spatially independent of and perpendicular to the preceding transient basal protrusions (Figures 1B, 2A, and 2B; Videos S1 and S2). In only one case have we seen a neuron generate what appears to be a forked axon with two ventrally directed branches. In this case, one of these branches was quickly retracted leaving the usual morphology of a single ventral axon.

Our analysis does not include the primary sensory Rohon-Beard neurons, which develop three axons (two central longitudinal axons and a peripheral axon) and are likely to use a different program of axogenesis (Andersen and Halloran, 2012). Our data also contain only one definitive motoneuron because their very ventral location impedes imaging. However, the single motoneuron has short basal protrusions and was the only neuron that did not have a ventral trajectory to its initial axon growth; instead, it directed its axon laterally from the cell body toward the nearby somite boundary before exiting the cord to innervate the muscles (Figure 2C, observations summarized in diagram in Figure 2D).

**Non-apical Progenitors in Spinal Cord Also Extend Basal Protrusions prior to Apical Detachment**

In addition to the apical progenitors that generate most of the neurons of zebrafish CNS, a scarce population of basal progenitors that divide in non-apical locations is also present (Alexandre et al., 2010; McIntosh et al., 2017). We call these progenitors non-apical progenitors (or NAPs) and previously demonstrated that the majority of spinal NAPs express Vsx1 and share molecular and regulatory mechanisms with neurons (McIntosh et al., 2017). This prompted us to investigate whether spinal NAPs might also share the morphological program of differentiation with neurons. We were able to monitor 7 NAPs by confocal time-lapse microscopy all of which undergo the stereotypical T-shape transition characteristic of differentiating neurons prior to their basal mitosis (Figure 3A; Video S3). The NAP exemplified in Figure 3A has a basal cell body that transiently extends a pair of long basal protrusions that are filopodia rich while still attached to the apical surface (Figure 3A; Video S3). The basal protrusions on NAPs are often asymmetric in length (6 out of 7 cases). On some cells, basal protrusions do not fully retract before NAP mitoses (4 out of 7 cells) (Figures 3A and 3B; Video S3). In these cases, the retraction of basal protrusions is completed after mitosis (green arrow in Figures 3A and 3B) but still prior to axon formation in the two daughter neurons (blue arrow in Figures 3A and 3B; Video S3).

These observations show spinal neurons and NAPs share common stereotypical morphological behaviors and further confirm that spinal Vsx1 NAPs and differentiating neurons share cellular and molecular characteristics as suggested previously (McIntosh et al., 2017).

**Differentiating Telencephalic Neurons Do Not Form Long Transient Basal Protrusions**

The elongation of basal protrusions seems to be a consistent feature of differentiating neurons and NAPs in the zebrafish spinal cord. To investigate whether the T-shape transition is common to differentiating neurons in other regions of the zebrafish CNS, we analyzed neuronal differentiation in the dorsal telencephalon from 20 to 40 hpf. Using this approach, we find that differentiating neurons in the telencephalon do not extend transient basal protrusions prior to apical detachment and axogenesis (n = 16 out of 16 cells) (Figure 3C; Video S4). In these cells, axon formation derives from the basal end of the new neuron’s...
radial process and usually immediately follows the detachment of the neuron from the apical surface (Figure 3C; Video S4). These observations demonstrate that the programs of axogenesis and apical release are regionally distinct, suggesting a region-specific role for the T-shape transition in spinal differentiation.

Neurons Rarely Differentiate Close Together in Time and Space
To quantify the spatiotemporal dynamics of spinal neuron differentiation, we used in vivo confocal microscopy to determine the spatiotemporal pattern of differentiation of Vsx1:GFP-expressing neurons in the zebrafish spinal cord. Vsx1:GFP neurons are...
born in pairs from the terminal division of vsx1-expressing NAPs (Kimura et al., 2008; McIntosh et al., 2017). GFP is detected in their progenitor immediately before terminal division and maintained in their daughters (Figure 4A). The appearance of adjacent GFP-expressing daughters thus offers a distinct and easily recognized time point to record as the start of differentiation of those neurons (Figure 4A). Using this criterion, we recorded the position and time of the start of differentiation of every pair of Vsx1 positive neurons in a 250- to 400-μm length of spinal cord at the level of somites 9–14 and between 19 and 27 hpf. We did this for both left and right sides in 17 embryos, thus recording 449 Vsx1 differentiation events in space and time within 34 equivalent stretches of spinal cord (Figure 4B; Data S1).

These data confirm that Vsx1 neurons differentiate in a long-distance spacing pattern with later born neurons differentiating in the gaps between already existing neurons (Figure 4C; Video S5). Time-lapse movies show no evidence that Vsx1 neurons or their progenitors migrate into this space; rather, these cells maintain stable positions. This pattern of sequential differentiation in the gaps continues for the next 6 h, at which time a near continuous line of Vsx1 neurons has been generated (Figures 4B and 4C; Video S5; Data S1).

To quantify this spatiotemporal pattern of differentiation, we looked at the timing of Vsx1 differentiation events that happened less than 20 μm apart. Neuroepithelial cells are typically 10.5 ± 4.1 μm (mean ± SD, n = 95 cells) wide at their basal pole, so this correlates to less than two cell diameters. Of the 449 Vsx1:GFP differentiation events, in only 7 cases (1.6%) were the differentiation events closer in time and space than 20 μm and 60 min apart (Figure 4D). The majority (68.3%) of events that occurred within 20 μm occurred between 2 and 3.5 h apart. Additionally, most consecutive Vsx1 differentiation events (i.e., those that occur closest in time) occur at a distance of 50–60 μm (Figures 4E and 4F).

These data suggest the presence of a mechanism that regulates the spatiotemporal differentiation of Vsx1 neurons in order to sequentially transform a long-distance spacing pattern into a continuous column of neurons.
Figure 4. Neurons Rarely Differentiate Close Together in Time and Space

(A) Vsx1:GFP expression in a single cell before, during, and after a NAP division. Following mitosis, GFP expression is maintained, and axogenesis can be followed in both daughter neurons.

(B) Spatiotemporal pattern of Vsx1:GFP neuronal precursor differentiation from 19 to 27 hpf. The location of Vsx1:GFP NAPs at the time of mitosis are represented as pairs of green circles and plotted in space (x axis) and time (y axis). The black lines descending through time from the pair of green circles represent the position held by the daughter cells after mitosis.

(C) Image sequence from a time lapse showing the differentiation of Vsx1:GFP neurons in one section of spinal cord through time. The left panel shows Vsx1:GFP neurons differentiating over time. In the right panel, cells have been color coded to denote sister pairs. All images are projections from small confocal z stacks. See also Video S5.

(D) Frequency distribution showing the difference in time between Vsx1:GFP mitoses that occur less than 20 μm apart.

(E) Diagram illustrating the method used to calculate the distance between successive Vsx1:GFP differentiation events from a time-lapse movie. $t_i$ indicates the time of differentiation and $d_i$ the distance between successive differentiation events.

(F) Histogram showing the distribution of the distance between successive Vsx1:GFP differentiation events in wild-type embryos.
Transient Basal Protrusions Express DeltaD and Notch Activity Is Upregulated in Their Vicinity

Our previous section analyzed Vsx1 neurons to show that neuronal differentiation in the embryonic zebrafish spinal cord occurs with an initial sparse pattern followed by sequential filling (Figure 4). Similar patterns of differentiation are also apparent in previous studies of other neuronal subtypes (Gribble et al., 2009; Hutchinson and Eisen, 2006; Hutchinson et al. 2007; Kimura et al., 2008; England et al., 2011). This data suggests a mechanism may exist to transiently inhibit neuronal differentiation over a distance of several cell diameters from each newly differentiating cell and that this mechanism is sequentially released to allow differentiation in the initially inhibited space. We hypothesize that the transient basal protrusions on newly differentiating neurons and NAPs could mediate lateral inhibition at a distance in time and space. Since Delta-Notch signaling has been suggested to mediate lateral inhibition at a distance to regulate sparse pattern formation in other systems (reviewed in Pröls et al., 2016), we tested whether the transient basal protrusions on differentiating neurons could potentially mediate transient Delta-Notch signaling in our system.

Using an antibody against the DeltaD protein and a DeltaD transgenic reporter line Tg(DeltaD:GAL4c;UAS:GFP) (Scheer et al., 2001), we were able to determine that the DeltaD transgene highlights cells with typical T-shape morphology and that DeltaD protein is specifically enriched in the basal protrusions and cell body of these cells (Figures 5A and 5A'). Furthermore, if the basal protrusions participate in long-range lateral inhibition we expect them to activate Notch signaling pathway in the surrounding cells contacted by the basal protrusions. Importantly, this should occur in cells out of range of contact from the neuronal cell body. To test whether this is the case, we randomly labeled differentiating neurons in the Notch reporter line Tg(Tp1: VenusPEST) (Ninov et al., 2012) and monitored the dynamics of Notch activation in nearby cells. We measured the relative mean intensity values of VenusPEST expression in a neuroepithelial region contacted by the labeled basal protrusion (but not the neuronal cell body) and compared it to a control region that had not been contacted by an identified protrusion (Figure 5B).

We assessed VenusPEST expression 2 h after basal protrusions reached their maximum length. We found the amount of VenusPEST expression is significantly increased in regions spatially related to the identified protrusions when compared to the control region (Figure 5C). These observations are therefore consistent with the hypothesis that basal protrusions activate Notch signaling in order to delay neuronal differentiation in cells at a distance from the differentiating neuronal body.

Since basal protrusions extend bidirectionally along the same D/V level as the differentiating cell body, these protrusions will be perfectly placed to preferentially interact with neural progenitors located at the same D/V level (i.e., progenitors likely to generate neurons of the same subtype) and promote the neuronal spacing pattern observed in the zebrafish spinal cord. This suggests that the relative positions of neurons of different subtypes could be independent of each other. To test this, we measured the relative positions between different neuronal subtypes (evx1, eng1b, and Vsx1:GFP; Figures S3A–S3H). This analysis revealed that positions of evx1 and eng1b neurons had no consistent alignment with Vsx1:GFP-expressing neurons (Figures S3F–S3H), suggesting that there is no pre-pattern for the relative position of different neuronal subtypes along the anteroposterior axis, and that regulation of differentiation of a particular neuronal subtype is independent of interactions with neurons of other subtypes.

Together, these results are consistent with the existence of a long-distance lateral inhibition mechanism that operates between differentiating neurons of the same subtype and their progenitors at the same D/V level. The expression of DeltaD in transient basal protrusions and the increase in Notch activation in cells spatially related to these basal protrusions suggests the basal protrusions could control both the spatial and temporal pattern of differentiation through long distance but transient Notch-Delta lateral inhibition.

Laminin Depletion Reduces Both Basal Protrusion Length and Spacing between Successively Differentiating Neurons

To further test whether basal protrusions could regulate the spatiotemporal pattern of Vsx1 neuron differentiation, we...
modified basal protrusion length and quantified the pattern of neuronal differentiation in vivo. Since the transient basal protrusions grow at the basal surface of the neuroepithelium, we predicted that extracellular matrix proteins in the basement membrane could be required for their growth. To test this, we monitored neuronal differentiation in lamc1 mutants that have no detectable Laminin at the basal surface of the neuroepithelium at the developmental stages we are studying. Neurons differentiating in lamc1 mutant spinal cords develop significantly shorter basal protrusions (mean ± SD, 12.3 ± 4.7 μm, n = 39) than neurons in wild-type embryos (mean ± SD, 42.6 ± 20.2 μm, n = 24 cells), unpaired one-tailed t test p-value < 0.0001 (Figures 6A and 6B; Video S6), consistent with a role for Laminin in basal protrusion extension.

To determine whether the reduced length of basal protrusions in Laminin-depleted embryos could affect the spatiotemporal pattern of neuron differentiation, we performed time-lapse microscopy and compared the pattern of differentiation of Vsx1:GFP neuron pairs in lamc1 mutants (Data S1) (n = 721 differentiation events in 50 stretches of spinal cord in 25 embryos) and wild type. We found that successive differentiation events occur closer together in lamc1 mutants than in wild type (Figure 6C, mean ± SD, 54.00 ± 1.52 μm in wild type and 45.3 ± 0.99 μm in lamc1, one-tailed t test p-value = 3.16 × 10⁻³), with the highest frequency of these events occurring 30–40 μm apart in the mutant compared to 50–60 μm apart in the wild type (Figures 4F and 6C).

Since wild-type basal protrusions extend 42.6 μm on average (and can potentially influence differentiation in this range), we then determined the proportion of sequential differentiation events that occurred within 42.6 μm of each other in the wild type and lamc1 background. This verified that differentiation events are twice as likely to occur within this range in the lamc1 mutant (0.19 ± 0.11) than in the wild-type embryos (0.080 ± 0.11) (Kolmogorov-Smirnov test, p-value = 0.000066) (Figure 6D). We further compared the wild-type and lamc1 differentiation data to randomly generated differentiation events and found that the proportion of sequential events that occurred within 42.6 μm in the wild type, but not the lamc1 data, is significantly different from random (Kolmogorov-Smirnov test, p-value = 0.000224 and p-value = 0.213) (Figure 6D).

To discard the possibility that a decrease in neuronal spacing in lamc1 mutants is due to an overall increase in neuronal differentiation we quantified the rate of neurogenesis. We determined the ratio of neurons to progenitors (N/P) at early stages of embryonic development and found no difference between wild type and mutant (Figures S4A and S4B). In addition, we analyzed the overall organization of the spinal cord in lamc1 mutants and showed that patterns of polarity proteins, the locations of progenitor divisions and the location of neuronal differentiation are normal (Figures S4A–S4C). These experiments suggest that gross neuroepithelial organization and rates of differentiation are normal in lamc1 mutant embryos at early stages of embryonic development.

Overall, these results are consistent with the hypothesis that basal protrusions transiently extend the range of influence of lateral inhibition and longer basal protrusions can regulate differentiation over a longer distance.

**Theoretical Predictions Support the Role of Basal Protrusions in Patterning Differentiation through Delta-Notch-Mediated Lateral Inhibition**

To determine whether the pattern of neuronal differentiation can be explained by Delta-Notch-mediated lateral inhibition delivered via transient basal protrusions, we developed a physical description of lateral inhibition coupled to the observed...
protrusions dynamics. The dynamics of Delta-Notch signalling have been modelled extensively (Binshtok and Sprinzak, 2018). Here we built on Cohen et al. (2010) and Collier et al. (1996) and describe the process of lateral inhibition by,

\[
\frac{dN}{dt} = R_N \frac{D_{in}}{a + D_{in}} - \mu N \quad \text{(Equation 1)}
\]

\[
\frac{dD}{dt} = R_D \frac{1}{1 + bN} - \rho N \quad \text{(Equation 2)}
\]

\[
D_{in} = \alpha \sum_{\text{soma}} D + \beta \sum_{\text{protrusions}} D. \quad \text{(Equation 3)}
\]

These equations describe the dynamic process of gene activation and inhibition between signaling proteins in contacting cells. \(N\) and \(D\) refer to the amount of active Notch and Delta within cells, and \(D_{in}\) is the total signal received by a cell from all cells in contact with it. We assume that cells only mediate signaling through their protrusions and set \( \alpha = 0 \) and \( \beta = 1 \). Nonzero values of \( \alpha \) are considered in the STAR Methods. We further assume that the probability of neuronal differentiation correlates with a cell’s level of Delta expression (Hunter et al., 2016) and that neuronal differentiation commences with basal protrusion extension. The temporal and spatial dynamics of basal protrusions follow the experimentally observed dynamics. See STAR Methods for further details of the theoretical setup.

We first performed simulations to predict the distribution of \(dx\) assuming differentiation events occur randomly along the spinal cord (Figure 7A). If differentiation events occur at random, the distance between successive events should also be random. With random differentiation, the predicted distribution of \(dx\) (mean ± SD, 40.90 ± 21.55 μm) differs significantly from both the wild-type experimental distribution (mean ± SD, 54.53 ± 18.92 μm; Kolmogorov-Smirnov test, \(p\)-value < 10⁻¹²) (compare Figure 7A to Figure 4F), and the lamc1 mutant distribution (mean ± SD, 46.32 ± 18.68 μm; Kolmogorov-Smirnov test, \(p\)-value = 9.6 x 10⁻⁷) (compare Figure 7A to Figure 6C), confirming that the spatiotemporal patterns of differentiation in vivo are unlikely to be randomly generated.

We then performed simulations assuming that the protrusion dynamics follow those of the wild-type fish. The predicted distribution between successive differentiation events \(dx\) differs significantly from both the wild-type experimental distribution (mean ± SD, 40.90 ± 21.55 μm) and the lamc1 mutant distribution (mean ± SD, 46.32 ± 18.68 μm; Kolmogorov-Smirnov test, \(p\)-value = 0.121). We repeated the analysis but now assuming that the length and dynamics of protrusions follow those of the Laminin-deficient lamc1 mutant. Now the predicted distribution is in agreement...
with the distribution of \( dx \) in the lamc1 mutant found in vivo (compare Figure 6C to Figure 7C; Kolmogorov-Smirnov test, \( p \) value = 0.181). Furthermore, the lamc1 mutant distributions are significantly different to simulations with wild-type length protrusions (compare Figure 6C to Figure 7B; Kolmogorov-Smirnov test \( p \)-value \(< 10^{-10} \)). These results together suggest that the spatiotemporal dynamics of differentiation in wild-type and the lamc1 mutant can both be explained by protrusion mediated lateral inhibition (Figure 7D). The differences in the distribution of \( dx \) between the wild-type and lamc1 mutant can be explained by differences in basal protrusions length.

To understand how changes in basal protrusion length and dynamics impact on the spatiotemporal pattern of differentiation, we performed simulations while continuously varying the protrusion length. We found that the average distance between sequential events (\( \Delta x \)) follows a linear relationship with the protrusion length (Figure 7E). However, a given change in the protrusion length, \( dl \), only confers a change in the mean spacing \( \Delta x \), which is about 22% of \( dl \) (Figure 7E). This can be understood as follows. The protrusions specify a transient region where neurogenesis is inhibited. Although this generates a minimal spacing between sequential events, the events do not have to occur right at the boundary, and this alters the mean of the distribution (as seen in the noise around the peaks in Figures 4F, 6C, 7B, and 7C). This effect becomes stronger as the protrusions become smaller, which explains why large changes in the protrusion length in the lamc1 mutant do not produce equally drastic shifts in the average value of \( dx \) (Figures 4F and 6C, see Quantification and Statistical Analysis in the STAR Methods and Figure S5). The relative impact of the protrusions on the spacing between sequential events in our region of interest declines for smaller and slower protrusions. These considerations together explain why large changes in the protrusion length in the lamc1 mutant do not produce equally drastic shifts in the average value of \( dx \) (Figure 6C; see Figure S5 for detailed mathematical derivation and explanation).

To explore how the position and timing of differentiation are related, we also computed the spatial and temporal relationship between differentiation events (see STAR Methods). These analyses both in vivo and using our theoretical model showed that there is a negative correlation between the distance between two cells and the time at which they differentiate so that cells that are closer in space tend to differentiate further apart in time (Figures S6 and S7). In vivo, the wild-type and lamc1-mutant data both followed this trend; however, the range over which this correlation was present in lamc1 mutants was reduced, consistent with the reduced basal arm length in lamc1 mutants (Figures S6C and S6F). These spatiotemporal correlations also appear in our theoretical model when long or short basal protrusions mediate lateral inhibition (but not when differentiation occurs randomly), further supporting the role of basal protrusions in patterning neuronal differentiation (Figure S7).

Finally, we have performed simulations that assess differentiation patterns when lateral inhibition takes place only at soma-to-soma contacts and a combination of soma and basal protrusion contacts or only via basal protrusion contacts (STAR Methods). We found that including soma-to-soma lateral inhibition prior to protrusion extension cannot recapitulate our in vivo observations (Figure S6). This suggests that soma-to-soma contacts play a minimal role in the mechanism that determines the pattern of differentiation between spinal neurons.

**DISCUSSION**

Using live imaging in zebrafish, we have uncovered a cellular behavior for vertebrate neurons that regulates the spatiotemporal dynamics of neuronal differentiation along the spinal cord. Differentiating neurons and NAPs transiently develop two long basal protrusions prior to apical detachment and axogenesis. These basal protrusions express Delta at high levels and activate Notch signaling at a distance from the cell body. The dynamics of basal protrusion extension and retraction are consistent with a role in delivering Delta-Notch-mediated lateral inhibition at a distance to regulate the position and time of spinal neuron differentiation. Additionally, previous work has shown that Delta expression is required for the sparse spatial pattern of zebrafish spinal neurons (Okigawa et al., 2014). We show that experimental manipulation of basal protrusions in vivo and in a mathematical model of cells with and without signaling basal protrusions also support the role of basal protrusions in mediating lateral inhibition at a distance to regulate both the position and the time of spinal neuron differentiation. Protrusion-mediated lateral inhibition has been proposed to control sparse differentiation patterns in the fly peripheral nervous system (De Joussineau et al., 2003; Cohen et al., 2010). Our work demonstrates that a similar cell-protrusion-mediated mechanism operates in the spinal cord of a vertebrate.

The extension and retraction of basal protrusions on spinal neurons is highly stereotyped and is the earliest morphological feature of neuronal differentiation once the nucleus of the newly born neuron has reached the basal surface of the neural tube. Therefore, influencing the differentiative behavior of surrounding cells is prioritized over other essential neuronal behaviors such as axon outgrowth. Basal protrusions are robust microtubule based processes and always appear in pairs—one directed strictly anteriorly along the spinal cord and one directed strictly posteriorly. In contrast to the random protrusive activity observed on vertebrate neurons differentiating in vitro (Dotti et al., 1988), protrusive activity on spinal neurons differentiating in vivo is highly directed and predictable. We hypothesize that this directed longitudinal growth of basal protrusions is an effective way to preferentially contact and influence the behavior of neural progenitors at the same D/V level in the spinal cord. Progenitors from the same D/V level will likely generate neurons of the same subtype, and this directed basal growth maximizes the chance of influencing differentiation of similar neuronal sub-types. We find that NAPs (Vsx1-expressing progenitors) also undergo this predictable basal protrusive activity prior to their terminal division close to the basal surface of the spinal cord. They will therefore also be able to influence differentiation of similar NAPs. Thus, this morphological transition is another similarity between neurons and NAPs during their paths to differentiation (McIntosh et al., 2017).

Our analyses suggest neuronal basal protrusions deliver Delta-mediated lateral inhibition at a distance, a similar role to that proposed for the basal protrusions of SOPs on the fly notum and wing disk (De Joussineau et al., 2003; Cohen et al., 2010; Hunter et al., 2019). Although basal protrusions on SOPs and
spinal neurons share some similarities, there are some major differences between the two systems. SOPs radiate thin actin-based filopodia in all directions along the basal surface of the epithelium, while zebrafish neurons develop two substantial microtubule-based protrusions that grow in predictable orientations. The basal protrusions of zebrafish neurons often also have filopodia on their surface, which may increase the interactions between differentiating cells and their near neighbors. Cell bodies of zebrafish neurons also have filopodia on their surface, although these are much shorter than the basal protrusions. Contrary to dynamic basal filopodia on SOPs, basal protrusions on spinal neurons remain relatively stable and extended for several hours. Importantly however, spinal neuron protrusions are transient, and their retraction releases cells from long-distance lateral inhibition and allows other neurons to differentiate in the previously inhibited space. This suggests that spinal basal protrusions regulate both the time and space of neuronal differentiation.

Protrusive activity that could influence surrounding cell behaviors has previously been suggested in the rodent cortex. There, basal intermediate progenitors (BIPs) in the rat and mouse subventricular zone have a large number of multidirectional membrane extensions that have alternatively been suggested to sense local factors prior to mitosis (Noctor et al., 2004) or to mediate Delta-Notch signaling between BIPs and apical radial glia cells, which maintains the proliferative progenitor population (Nelson et al., 2013). Although the protrusions on rodent progenitors do not appear to have a stereotypic orientation and their relation to the spatial and temporal progression of neurogenesis in the cortex has not been assessed, it remains possible that they serve similar functions to the basal protrusions of spinal neurons and progenitors. Our observations in the zebrafish telencephalon show that newborn neurons in this region behave quite differently to spinal neurons. Early telencephalic neurons do not elaborate long basal protrusions prior to axogenesis, and there is no obvious spatiotemporal pattern of differentiation in this region. Thus, programmes of cell morphogenesis and neuronal differentiation are region specific.

Many of the neurons in the spinal cord arise from asymmetrically fated divisions (Alexandre et al., 2010; Das and Storey, 2012; Saade et al., 2013; Kressmann et al., 2015) where daughter cell fate is also regulated by Delta-Notch interactions. In asymmetric divisions, Delta-Notch signaling is likely to be mediated exclusively between the sister cells of each division (Dong et al., 2012; Kressmann et al., 2015). Our modeling suggests that lateral inhibition between immediate neighbors cannot explain the long-distance spacing pattern of neuronal differentiation; nonetheless, this local mechanism that operates during progenitor divisions must be integrated with the long-distance mechanism delivered through basal protrusions. We have not investigated how these two processes might work together, but we favor the possibility that lateral inhibition through long basal protrusions delays neuron (and NAP) differentiation after their birth rather than regulating the time of their birth or particular fate. Our own unpublished data show that neurons born at the same time begin to express the neuronal transgene HuC:GFP within a very wide time window (4–12 h after their birth); thus, neurons can progress through their differentiation pathways at very different rates. Prospective neurons can initially maintain high levels of Notch activity, and reduction in Notch activation accelerates their differentiation (Baek et al., 2018), raising the possibility that the transient lateral inhibition mediated by basal protrusions controls the time of differentiation but does not change cell fate.

To test the potential for basal protrusions to mediate the spatial pattern of differentiation in vivo, we examined spinal neuron differentiation in Laminin-depleted spinal cords. We found that basal protrusion growth is significantly reduced in the absence of Laminin, and this correlates with a predicted reduction in the distance between differentiation events. Laminin depletion did not completely abolish basal protrusions from spinal neurons, and we show that the short protrusions that remain can explain the altered spatiotemporal dynamics of differentiation in the mutant. Although we cannot eliminate the possibility that Laminin depletion alters the spatial pattern of differentiation through mechanisms other than reduced basal protrusion length, this experimental approach is consistent with our major hypothesis. The overall architecture and cellular organization of the Laminin-depleted spinal cord is grossly normal, and we propose that a Laminin-rich extracellular matrix may be required for basal protrusion growth, perhaps in a similar way to Laminin’s proposed role in axonal growth at the basal surface of neuroepithelium (Randlett et al., 2011).

Theoretical modeling that captures the protrusion dynamics in our in vivo system supports the hypothesis that basal protrusions mediate the spatiotemporal pattern of differentiation. We show that the spacing between successively born neurons is linear with protrusion length. Furthermore, our theoretical model recapitulates the spatiotemporal patterns in vivo in both wild-type and Laminin-depleted measurements. Interestingly, the inclusion of lateral inhibition via soma-to-soma signaling in our model introduces discrepancies between model output and in vivo data (Figure S8), suggesting that soma-to-soma signaling may be particularly weak during these events in vivo.

The biological function of regulating neuronal differentiation in a spatiotemporal manner is unclear. However, we speculate that it may be advantageous for neuronal circuit formation if the initial connections are made between a minimal number of spatially distributed neurons. Later, differentiating neurons can then be added to a functioning circuit to consolidate or modify the circuit function. This could be particularly important in zebrafish and amphibian embryos, as they develop externally and need to quickly build a functional motor circuit for survival.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
- METHOD DETAILS
  - In Vivo Experimental Details
  - Theoretical and Computational Details
- QUANTIFICATION AND STATISTICAL ANALYSIS
  - Analysis of In Vivo Data
  - Computational Analysis of Simulated and In Vivo Data
SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.devcel.2019.05.035.

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The authors declare no competing interests.

AUTHOR CONTRIBUTIONS

P.A. and J.C. conceived the project, supervised the work, and wrote the manuscript. P.A. characterized the individual neuronal behaviors and prepared figures and movies. P.A. and R.E.M. characterized the neuronal behaviors and Vsx1 distribution in lamc1 mutants. R.M. characterized the Vsx1 distribution in wild-type embryos. P.A. and R.E.M. obtained the Delta and Tg(TP1:VenusPEST) experimental data. G.G., P.A., and Z.H. developed the model to quantitatively Tg(TP1:VenusPEST). Z.H. developed the mathematical model, wrote all computational code, performed simulations, did most of the statistical analysis, and contributed to writing the manuscript. P.A. and R.E.M. characterized the neuronal behaviors and Vsx1 experimental data. All authors commented on the manuscript.

DECLARATION OF INTERESTS

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| Mouse monoclonal anti-HuC/D (16A11) | Invitrogen | Cat#A-21271; lot1252835; RRID:AB_221448 |
| Rabbit polyclonal anti-aPKC \( \zeta \) (C-20) | Santa Cruz Biotechnology | Cat#SC-216; k0413 |
| Mouse monoclonal anti-DeltaD (zd2) | Cancer Research Technology | Cat#C7/2/14; lot255/06 |
| Chicken polyclonal anti-GFP | Abcam | Cat#ab13970; lotGR89472-7; RRID:AB_300798 |
| **Chemicals, Peptides, and Recombinant Proteins** |        |            |
| 1-phenyl-3-(2-thiazolyl)-2-thiourea | Sigma-Aldrich | Cat#P4015 |
| Sytox Green | ThermoFischer Scientific | Cat#S7020 |
| Fast Red substrate | Roche | Cat#1758888001 |
| MS-222 | Sigma-Aldrich | Cat#E10521 |
| **Critical Commercial Assays** |        |            |
| SP6 mMessenger mMachine kit | Ambion | Cat#AM1340 |
| **Experimental Models: Organisms/Strains** |        |            |
| Zebrafish – Ekkwill | N/A | N/A (wildtype strain) |
| Zebrafish – AB/Tuebingen (wildtype strain) | N/A | N/A (wildtype strain) |
| Zebrafish – Tüpfel long fin | N/A | N/A (wildtype strain) |
| Zebrafish – Tg(vsx1:GFP) | Kimura et al., 2008 ZFIN ID: ZDB-FISH-150901-23998 |
| Zebrafish – Tg(deltaD:Gal4;UAS:GFP) | Scheer et al., 2001 ZFIN ID: ZDB-FISH-150901-6106 |
| Zebrafish – Tg(TP1:VenusPEST) | Ninov et al., 2012 ZFIN ID: ZDB-FISH-150901-8023 |
| Zebrafish – sleepy; lamc1sa379 mutant | Kettleborough et al., 2013 ZFIN ID: ZDB-FISH-150901-23200 |
| Zebrafish – Tg(vsx1:GFP);lamc1sa379 mutant | This paper | N/A |
| **Recombinant DNA** |        |            |
| Plasmid: pCS2-mCherry-CAAX (referred to as m-RFP) | Laboratory of Chi-Bin Chien; Kwan et al., 2007 | N/A |
| Plasmid: pCS2-EGFP-CAAX (referred to as m-GFP) | Laboratory of Chi-Bin Chien; Kwan et al., 2007 | N/A |
| Plasmid: pCS2-mKate2-CAAX (referred to as m-mKate2) | This paper | N/A |
| Plasmid: pCS2-H2B-RFP (referred to as n-RFP) | Laboratory of Steffen Schlopp; Megason and Fraser, 2003 | N/A |
| Plasmid: pCS2-Eb3-GFP | Laboratory of William Harris; Norden et al., 2009 | N/A |
| **Software and Algorithms** |        |            |
| Volocity 3D Image Analysis Software | Perkin-Elmer | http://www.perkinelmer.com/pages/020/cellularimaging/products/volocity.xhtml; RRID:SCR_002668 |
| Fiji | Schindelin et al., 2012 | http://fiji.sc; RRID:SCR_002285 |
| Wolfram Mathematica | Wolfram | https://www.wolfram.com/mathematica; RRID:SCR_014448 |
| Prism 7 | GraphPad | https://www.graphpad.com/scientific-software/prism; RRID:SCR_005375 |

### CONTACT FOR REAGENT AND RESOURCE SHARING

Further requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Paula Alexandre (p.alexandre@ucl.ac.uk). Details on theory and computational models can be obtained from Zena Hadjivasiliou (Zena.Hadjivasiliou@unige.ch).
EXPERIMENTAL MODEL AND SUBJECT DETAILS

All animal procedures were performed according to the UK Animal (Scientific Procedures) Act 1986 and carried out under Home Office Project Licence number PPL P70880F4C, which was subject to local AWERB Committee review and Home Office approval. The following zebrafish lines were used: Ekkwill, AB/Tuebingen, Tuepfel long fin, Tg(vsx1:GFP) (Kimura et al., 2008), Tg(deltaD:Gal4;UAS:GFP) (Scheer et al., 2001), Tg(TP1:VenusPEST) (Ninov et al., 2012), and lamc1as379 mutant (sleepy; Kettleborough et al., 2013). Tg(vsx1:GFP) and lamc1as379 lines were crossed to establish a Tg(vsx1:GFP);lamc1as379 line. Adults were maintained under standard conditions as previously described (Westerfield, 2000), in a 14/10 hour light/dark cycle.

Embryos were obtained by natural spawning and raised in water or E2 medium at 28.5°C. If necessary, they were transferred to 0.003% 1-phenyl-3-(2-thiazolyl)-2-thiourea (Sigma-Aldrich) at 24 hpf to inhibit pigmentation.

Injections were performed at 16-64-cell stage. Embryos positive for mRNA expression, transgenic GFP expression and/or lamc1as379 phenotype were selected for imaging. Live imaging was performed at 18-42 hpf. In situ hybridisation was performed at 22 hpf and immunohistochemistry at 22-28 hpf. Sex is not yet determined at these stages in zebrafish so was not taken into account.

METHOD DETAILS

In Vivo Experimental Details

Immunohistochemistry

Whole-mount immunohistochemistry was performed on wild type and lamc1as379/- embryos to assess neurogenesis and epithelial cell polarity, and on Tg(deltaD:Gal4;UAS:GFP) embryos to assess Delta protein expression. Embryos were fixed for 2 hours at room temperature in 4% PFA at 22-28 hpf. Primary antibodies used were against HuC/D (mouse anti-HuC/D, Invitrogen, diluted 1:200), aPKC (rabbit anti-aPKC, Santa Cruz Biotechnology, diluted 1:500), DeltaD (mouse anti-DeltaD, Cancer Research Technology, diluted 1:50) and GFP (chicken anti-GFP, Abcam, diluted 1:1000). Embryos were incubated with primary antibody for 2 to 3 days at 4°C in PBS Triton 0.5%, 2% BSA, 10% goat serum (detailed protocol described in Wright et al., 2011). Embryos were incubated in secondary antibodies overnight at 4°C in the same blocking solution. Sytox Green (ThermoFischer Scientific, diluted 1:2,000) was added with secondary antibody to label nuclei and show mitotic divisions.

In Situ Hybridisation

Embryos fixed for 2 hours at room temperature in 4% PFA at 22 hpf were processed for whole-mount in situ hybridisation according to the protocol described in (Thisss and Thisse, 2000). For synthesis of antisense mRNA DIG-labelled probes DNA plasmids containing a cDNA fragment of eng1b (Batista et al., 2008), evx1 (Thaéron et al., 2000), and vsx1 (Passini et al., 1997) were linearised and the cDNA fragment was reverse transcribed using the RNA polymerases T3, T7, T3, respectively. Probes were detected in wild type embryos using Fast Red (Roche) substrate. Some embryos were incubated in Sytox Green (ThermoFischer Scientific, diluted 1:2,000) to label nuclei. To compare the relative distribution of neuronal subtypes, we performed in situ hybridisation for eng1b or evx1 in Tg(vsx1:GFP) transgenic embryos followed by the detection of GFP expression by immunohistochemistry (chicken anti-GFP, Abcam, diluted 1:1000).

mRNA Injection

Plasmids containing cDNAs coding for the following fusion proteins were linearised and the mRNA synthesised using SP6 mMessenger mMachine kit (Ambion): membrane tagged RFP (mCherry-CAAX; referred to as m-RFP)(Kwan et al., 2007), m-GFP (EGFP-CAAX)(Kwan et al., 2007), m-mKate2, (mKate-CAAX)(this paper), nuclear tagged RFP (H2B-RFP; referred to as n-RFP)(Megason and Fraser, 2003), and Eb3-GFP (Norden et al., 2009). mRNA was injected into a single cell of wild type, lamc1as379 mutant or Tg(TP1:VenusPEST) embryos at 16-64 cell stage to cause mosaic labelling and the embryos allowed to grow until imaging.

Confocal Imaging

Prior to imaging, live embryos were anaesthetised in MS-222 (Sigma-Aldrich). Fixed and live embryos were mounted in 1.5% low-melting point agarose (Sigma-Aldrich) in a petri dish with the dorsal spinal cord or dorsal telencephalon facing up. Fixed embryos were kept in PBS1x during imaging, while live embryos were kept at 28.5°C in E2 medium containing MS-222 (Sigma-Aldrich) and 0.003% 1-phenyl-3-(2-thiazolyl)-2-thiourea (Sigma-Aldrich).

Live imaging of individual cells was performed to observe neuronal differentiation and Notch activation. mRNA-injected wild type, lamc1as379/- or Tg(TP1:VenusPEST) embryos were imaged on a spinning-disk confocal microscope using an UltraVIEW VoX system (Perkin-Elmer) built on a Nikon Ti-E microscope, with a 40x water-immersion objective with numerical aperture (NA) of 1.0. Z-stacks were acquired at 0.5-1 μm. A series of z-stacks were obtained every 3 to 8 minutes for between 3 and 20 hours from 16 hpf.

Live imaging was performed on Tg(vsx1:GFP) and Tg(vsx1:GFP);lamc1as379/- embryos to assess spatiotemporal dynamics of neuronal differentiation on a SP5 confocal (Leica) microscope with a 20x water-immersion objective with an NA of 0.95. Z-stacks were acquired at 1 μm every 5 to 8 seconds for 8-10 hours.

Fixed whole-mount tissue from in situ hybridisation and immunohistochemistry was imaged on a SP5 confocal (Leica) microscope (described above) or on a LSM880 laser scanning confocal (Zeiss) microscope equipped with a 20x water-immersion objective with an NA of 0.95.

Image Processing and Analysis

Individual basal protrusions were measured from the cell body to the periphery of the basal protrusion. The maximum overall length reached by basal protrusions includes the cell body width. This analysis was performed in 3D at single and multiple timepoints using...
Volocity software (PerkinElmer). Images and movies shown in the manuscript result from a small projection of confocal z-stacks created using Fiji (Schindelin et al., 2012). Extra cells were occasionally removed from the field of view or pseudocoloured using Fiji to show examples of individual cells clearly.

To compare the intensity of Tg(TP1:VenusPEST) in the vicinity and away from the influence of the basal protrusions, we produced small z-projections, corrected drift and subtracted the background using Fiji. We used Fiji to measure the mean intensity values two hours after the basal protrusions reached their maximum length and analysed the area that had been in contact with the basal protrusions for at least 1h but was away from the neuronal cell body. For each case we calculated the ratio between the mean intensity under basal protrusions and control region (away from the basal protrusions).

Analysis of spatiotemporal dynamics of neuronal differentiation was performed at the level of somites 9 to 14 and between 19 and 27 hpf. The first appearance of adjacent GFP-expressing daughters following terminal division was considered to be the time of differentiation. Using Volocity, distances between temporally successive differentiation events were determined by measuring the distance (dx) between the last and the next neuronal pair born within a 80 μm (Figures 4F, 6C, and 7) and 42.6 μm (Figure 6D) space interval. The distance between neurons in fixed tissue was also measured using Volocity.

**Theoretical and Computational Details**

**Lateral Inhibition Driven Differentiation**

We used a mathematical model to simulate Notch-Delta mediated lateral inhibition. The model, as defined by Equations 1, 2, and 3 in the main text, describes the dynamics gene activation and inhibition via cell-cell signalling. $D_{in}$ in Equation 3 is the total amount of incoming Delta summed over soma-to-soma and basal protrusion mediated contacts. The parameters $\alpha$ and $\beta$ represent the relative amount of Delta at the soma-to-soma and in the basal protrusions respectively or the strength of the signal at the two locations. In the analysis presented in the main text we assumed that $\alpha = 0$ so that only basal protrusions mediate Notch signalling. We also relaxed this assumption (see Quantification and statistical analysis section “Signalling at soma-to-soma” and Figure S5) to investigate whether Notch signalling at soma-to-soma contacts could also be important. $R_N$ and $R_D$ are the baseline production rates for Notch and Delta molecules, $a$ and $k$ are parameters that determine how strongly incoming Delta induces Notch signalling, whereas $b$ and $h$ determine the strength of inhibition of Delta from Notch levels within the same cell. Finally, $\mu$ and $\rho$ are the degradation rates of Notch and Delta, respectively.

We applied the model to a 1D array of cells of variable size following the measured size distribution. We developed a theoretical description of lateral inhibition and cell differentiation in a one dimensional tissue (i.e. a row of cells). We construct the row of cells by sampling cell diameters from a normal distribution with mean 11.10 μm and s.d. 4.51 μm, the experimentally measured values in the neuroepithelium. This captures the diversity seen in the cell width of differentiating neurons, dividing cells and neuroepithelial cells. We used our setup to simulate differentiation events in the row of cells under different conditions as described below and in the main text. Signalling dynamics in individual cells could then be fully defined by the coupled system of differential equations 1, 2, and 3. Cells could make contact at the soma cell membranes and / or via basal cellular protrusions (Methods Image 1).
Methods Image 1: Signalling models considered in theoretical setup. Red indicates the presence and grey the absence of signalling. A: only basal protrusions can contribute to lateral signalling, B: basal protrusions and soma-to-soma contacts participate in lateral signalling, C: only soma-to-soma contacts contribute to lateral signalling.

**Protrusion Dynamics**

We modelled basal protrusion dynamics by allowing cells to extend protrusions if their Notch expression falls below a threshold (Hunter et al., 2016). Differentiating cells send but do not receive a signal (Sprinzak et al., 2010, 2011). Protrusions were extended at a constant rate and stopped growing when they reached length \( l_{\text{max}} \) where \( l_{\text{max}} \) was sampled from a normal distribution with mean 42.6 μm and s.d. 20.2 μm, following the in vivo measurements for maximum basal protrusion length. Once maximum length was reached the protrusions retracted at a rate 1.7 times faster than the extension rate (following in vivo dynamics). For the lamc1 mutants we modified the distribution of \( l_{\text{max}} \) to follow the mutant distribution with mean 12.3 μm and s.d. 4.7 μm and implemented extension and retraction rates that were 1.4 times slower than the wild-type and retraction rates 2.5 times slower than the wild-type, following the rates measured experimentally. A cell was assumed to have differentiated when both its right and left protrusion were fully retracted. Differentiated cells no longer participated in signalling and we ran simulations until all virtual cells underwent differentiation.

We assume that cells begin extending their protrusions with a probability that depends on the levels of their Notch expression so that differentiation becomes more likely as Notch levels fall below a threshold. We implement this following previous work (Hunter et al., 2016) by computing the probability of entering differentiation using a Hill function,

\[
P_{\text{diff}} = p \left( \frac{N}{N_{\text{th}}^q} \right) \quad \text{Equation 4}
\]

for each cell, where \( N \) is the Notch expression of that cell and the parameters \( N_{\text{th}} \) and \( q \) determine a Notch threshold and the window around this threshold that lead to differentiation. The prefactor \( p \) is the upper limit of the likelihood of differentiation per time step in the simulation. Differentiated cells no longer participate in lateral inhibition. In addition, protrusions are high in Delta but are assumed to carry a negligible number of free notch receptors (e.g. due to cis-inhibition) and so they only send but do not receive a signal (Sprinzak et al., 2010, 2011).

The values of all model parameters for all figures presented in the main and supplemental text are provided on Methods Table 1.

| Variable | Definition | Value |
| --- | --- | --- |
| \( N \) | Notch levels | 1.0 |
| \( D \) | Delta levels | 1.0 |
| \( D_{\text{in}} \) | Total incoming Delta | 100 |
| \( P_{\text{d}} \) | Arm extension and neuronal differentiation probability | 0.2 |
| \( R_N \) | Baseline Notch production rate | 0.91 |
| \( R_D \) | Baseline Delta production rate | 2 |
| \( a \) | Dissociation constant in Notch production | 0.1 |
| \( b \) | Hill coefficient in Notch production | 1.0 |
| \( h \) | Hill coefficient in inhibition of Delta by Notch | 3.0 |
| \( \mu \) | Notch degradation rate | 1.0 |
| \( \rho \) | Delta degradation rate | 1.0 |
| \( r \) | Maximal differentiation probability per unit time | 10^{-4} |
| \( N_{\text{th}} \) | Notch threshold for differentiation | 1 |

**Methods Table 1**: Definition of parameters in mathematical model. The table also indicates the values used for all figures that use simulated data in the main text and Supplemental Information.

**Randomly Differentiating Tissue**

We simulated a randomly differentiating tissue by initiating a row of cells as described above and then allowing cells to differentiate at random. In a row of \( n \) cells this corresponds to sampling from \( \{1, 2, \ldots, n\} \) without replacement and assuming that the \( i^{\text{th}} \) sampled number is equivalent to the \( i^{\text{th}} \) differentiation event. This allowed us to generate an ordered sequence of differentiation events and...
then compute the distance between cells (corresponding to the index Numbers 1 to $n$) that were sampled successively (Methods Image 2). In this way we were able to predict the expected distance between successive events in a randomly differentiating tissue (Figure 7A).

- In this example we have $n = 7$ cells
- Sample randomly without replacement from (1, 2, 3, 4, 5, 6, 7)
- An example output is: (1, 4, 7, 2, 5, 3, 6)
- Assume that cell 1 differentiated at time $t_1$, cell 4 at time $t_2$ and so on where $t_i < t_{i-1}$
- The distance between successive events can then be calculated as illustrated below

Methods Image 2: Algorithm for the generation of a randomly differentiating spinal cord. The positioning and size of cells were set according to experimental measurements.

**Numerical Details**

We initiate all simulations by randomly assigning each cell Notch and Delta levels sampled from $N(RN, 0.01RN)$ and $N(RD, 0.01RD)$ respectively where $N(\mu, \sigma)$ denotes the Normal distribution with mean $\mu$ and s.d. $\sigma$ for values of $RN$ and $RD$ given on Methods Table 1. Following this the Notch and Delta levels of each cell evolve according to Equations 1, 2, and 3 which we solved numerically using the Euler method (Euler step set to 0.01). Furthermore, a Gaussian noise term was applied to initiate protein concentrations and to the concentrations at each time step in the simulation.

At each step in the simulation each individual cell has a probability of initiating protrusion extension that is computed using Equation 4. Cells that begin extending protrusions spend $T_{ext}$ a.u. of time extending their protrusions and $T_{ext}/1.7$ a.u. of time retracting their protrusions, reflecting the relative amount of time cells were experimentally observed spending in the protrusion extension and retraction stages respectively. Once full protrusion retraction is achieved a cell is assumed to have differentiated to a neuron and no longer participates in the process of lateral signalling. The simulation parameter $T_{ext}$ was set to 0.05 units of time for all wild-type simulations and 0.05*(mean length of experimental lamc1 mutant/experimental wild-type basal protrusions)* $dT$ units of time in short protrusion simulations where $dT = 1.4$ reflecting that lamc1 protrusions extended 1.4 times slower than wild-type basal protrusions. The retraction time in mutant protrusions in the simulations was set to 1.1 times their extension time, again reflecting experimental measurements. We further discuss the role of $T_{ext}$ in the Quantification and Statistical Analysis section. A detailed outline of the algorithm we used throughout our analysis is shown on Methods Image 3. The numerical simulations produced a differentiation time for each individual cell together with its position. We used this information to compute ($dx$) as described in the main text (Figure 4C).
Establish a line of $n$ cells with apical radial sizes $A$, based on measured values ($\text{mean} = 11.10, \text{sd} = 4.51$)

for cells $i=1:n$

$A(i) = \text{Random Normal(mean, s.d.)} // \text{diameter of } i\text{th cell}$

$\text{pos}(1) = 0 \text{ // position of first cell}$

for cells $i=2:n$

$x(i) = x(i-1) + A(i-1) + A(i) \text{ // all other cells positioned relative to previous cell and according to their size}$

Assign each cell levels of Notch and Delta

for cells $i=1:n$

$N(i) = \text{Random Normal(1.0, 0.01)}$

$D(i) = \text{Random Normal(1.0, 0.01)}$

At each time step update Notch, Delta and arm distribution

For each time step

For all cells $i=1:n$

// Compute incoming delta from cell membrane contacts from immediate neighbours

$D_{\text{in, CM}}(i) = D(i+1) + D(i-1)$

// Compute incoming delta from basal protrusion contacts

For all other cells $j = 1:n$, $j \neq i$

If $(i-j) \& \& \text{DIFF}(j) \geq 0 \& \& (j\text{th cell's right arm's length}) > 0 \& \& (j\text{th cell's right arm's length}) \geq \text{pos}(i) - \text{pos}(j))$ OR

If $(i-j) \& \& \text{DIFF}(j) \geq 0 \& \& (j\text{th cell's left arm's length}) > 0 \& \& (j\text{th cell's left arm's length}) \geq \text{pos}(i) - \text{pos}(j))$

Then

$D_{\text{in, BP}}(i) = D_{\text{in, arms}}(i) + D(j)$

$D_{\text{in, total}}(i) = \alpha\ast D_{\text{in, CM}}(i) + \beta\ast D_{\text{in, BP}}(i)$

// Update Notch levels

$N_{\text{notch}}(i) = N_{\text{notch}}(i) + (-\mu\ast N_{\text{notch}}(i) + R_{\text{N,D}} \ast \text{in}(i) \ast \text{in}(j) / (a + D_{\text{in, total}}(i) \ast k)) \ast dt + \text{RandomNormal}(0, N_{\text{notch}}(i) \ast \text{rand error}) // \text{random error} = 0.01 \text{ and } dt = 0.05$

// Update Delta levels

$D_{\text{delta}}(i) = D_{\text{delta}}(i) + (-p\ast N_{\text{notch}}(i) + R_{\text{D}} \ast (1 + b\ast N_{\text{notch}}(i) \ast h)) \ast dt + \text{RandomNormal}(0, D_{\text{delta}}(i) \ast \text{rand error})$

// Update basal protrusions if time > 1.0 a.u.

If cell $i$ has no basal protrusions, initiate extension with probability equal to: $\text{value} = 1.0 - p\ast N_{\text{notch}}(i) \ast q / (N_{\text{notch}}(i) \ast q + N_{\text{th}}(i) \ast q)$

If basal protrusions extending, extend further by euler_step*T_ext:

$BP_{\text{left, L}}(i) = BP_{\text{left, L}}(i) + \text{euler_step} \ast T_{\text{ext}} \ast \text{Normal}(F_{\text{mean}}, F_{\text{SE}})$

$BP_{\text{right, L}}(i) = BP_{\text{right, L}}(i) + \text{euler_step} \ast T_{\text{ext}} \ast \text{Normal}(F_{\text{mean}}, F_{\text{SE}})$

If $F_{\text{mean}} = 42.3, F_{\text{SE}} = 3.09$

If a basal protrusion has exceeded Normal($F_{\text{mean}}, F_{\text{SE}}$) initiate retraction.

If basal protrusion retracting, retract further by euler_step*T_ext*1.7

$BP_{\text{left, L}}(i) = BP_{\text{left, L}}(i) - \text{euler_step} \ast T_{\text{ext}} \ast 1.7 \ast \text{Normal}(F_{\text{mean}}, F_{\text{mean}} \ast 0.1)$

$BP_{\text{right, L}}(i) = BP_{\text{right, L}}(i) - \text{euler_step} \ast T_{\text{ext}} \ast 1.7 \ast \text{Normal}(F_{\text{mean}}, F_{\text{mean}} \ast 0.1)$

When retraction complete, cell differentiates and exits signalling

Return differentiation time and position for each cell
Changes in dx with Protrusion Length (Figures 7 and S5)

\[ l_{\text{max}} \] extends a protrusion of length the same change in the average distance between sequential events (Figure 7E). A change in the main text we have shown that a specific change in the average maximum length reached by protrusions does not lead to the \( d_{\text{L}} \) extends a protrusion of length the same change in the average distance between sequential events (Figure 7E). A change in the main text we have shown that a specific change in the average maximum length reached by protrusions does not lead to the change in the value of \( dx \) with the protrusion length is only expected to lead to a change of 0.22 \( dL \) in the average value of \( dx \). This can be understood as follows. Consider a single differentiating cell which extends a protrusion of length \( l_{\text{max}} \) and inhibits any cell within a distance \( l_{\text{max}} \) from differentiating while the protrusion is present as shown in the diagram in Figure S5A.

In the limiting case where the protrusion extends instantaneously the following differentiation event will occur at a distance between \( d + l_{\text{max}} \) and \( L \) from the differentiating cell with equal probability, where \( L \) is the maximum distance away from our cell of interest (Figure S5A). It follows that the next differentiation event is expected to occur (on average) at a distance of \( d + L + l_{\text{max}} \) away from our cell of interest (the mean of a Uniform distribution 2 on the interval \( (d + l_{\text{max}}, L) \)). If we substitute \( L = 80 \mu \text{m} \) (the maximum dx value in our analysis) and \( d = 10 \mu \text{m} \) (the average cell diameter) we obtain, \( dx = 45 + 0.5l_{\text{max}} \). Assuming that the distribution of sequential differentiation events is stationary (i.e. time independent) it follows that,

\[ \bar{dx} = 45 + 0.5l_{\text{max}} \]

This means that in the limiting case where protrusions extend extremely fast a change in the average protrusion length equal to \( dL \) will lead to a change in \( dx \) equal to only 0.5 \( dL \).

Now consider a second case where the protrusions extend extremely slowly so they do not effectively inhibit neighbouring cells from differentiating. In this case, neighbouring cells will differentiate anywhere between \( d \) and \( L \) away from the differentiating cell and the expected value for \( dx \) become independent of the protrusions so that,
\[ \bar{x} = 45 \]

We expect a real tissue to lie in between these two limiting cases so that, \[ \bar{x} = 45 + \varphi_l^{l_{\text{max}}} \]

where \( \varphi \) is a constant between 0 and 0.5 and depends on the timescale of protrusion extension and lateral inhibition relative to the timescale of differentiation (shaded region in Figure S5B).

We can compute \( \varphi \) for our experimental data by substituting \( dx = 54.0 \mu \text{m} \) and \( l_{\text{max}} = 42.6 \mu \text{m} \) for the wild-type and \( dx = 45.3 \mu \text{m} \) and \( l_{\text{max}} = 12.3 \mu \text{m} \) for the lamc1 mutant. It follows that \( \varphi_{\text{WT}} = 0.22 \) and \( \varphi_{\text{lamc1}} = 0.024 \). The decrease in the slope in the mutant is consistent with a reduced speed in protrusion extension, as observed experimentally.

### Pairwise Differences in Space and Time (Figures S6 and S7)

In order to further investigate the coupling between the distance between any two differentiated cells and their time of differentiation we computed the distance in space, \( dx \) (not to be confused with \( dx \) which is the distance between sequential differentiation events), and differentiation time, \( dt \), between all pairs of cells in each experiment and different versions of the theoretical model setup. We then asked how the distributions of \( dx \) and \( dt \) depend on one another.

For experimental data, the distribution of the pairwise position differences \( dx \) for all pairs follows an approximately uniform distribution on the measured interval (Figure S6A). When we restrict this distribution to cells that differentiate within one hour of each other, however, we observe a change in the distribution: very few cells differentiate within less than 30 \( \mu \text{m} \) of one another and the distribution of \( dx \) becomes centred around 60 \( \mu \text{m} \) (Figure S6B). Furthermore, \( dx \) and \( dt \) were negatively correlated (Spearmann’s Rho = -0.26; Spearmann’s Rank test p-value = 2.2 \( \times \) 10\(^{-12} \)). We also plotted the mean \( dt \) for cells that differentiated within a specific space interval of one another (Figure S6C). The smaller the distance present up until an interval of 50-60 \( \mu \text{m} \) consistent with the average length of protrusions in the between two cells the larger the difference in the time of differentiation. This effect appears to be present up until an interval of 50-60 \( \mu \text{m} \) consistent with the average length of protrusions in the wild type.

We repeated the same analysis for the lamc1 mutant data. We once again found a negative correlation between \( dt \) and \( dx \) (Spearmann’s Rho = -0.16; Spearmann’s Rank test p-value = 4.6 \( \times \) 10\(^{-10} \) ). Furthermore, the distribution of \( dx \) shifts with very few cells differentiating right next to each other. However, the distribution of \( dx \) conditional on \( dt < 1 \text{hour} \) is shifted to the left in the lamc1 data when compared to the wild-type experimental data (Figure S6B versus S6E). This is consistent with shorter basal protrusions governing the spatiotemporal dynamics in the lamc1 mutant. We again plotted the mean \( dt \) for cells that differentiated within a specific space interval of one another (Figure S6F). The smaller the distance between two cells the larger the difference in their time of differentiation. Unlike the wild type data (Figure S6C), this effect is only present up until an interval of 20-30 \( \mu \text{m} \), consistent with a reduced range in lateral inhibition as reflected by the reduction in the protrusion length.

We then turned to pairwise differences for theoretical predictions. We asked whether these observations are consistent with a randomly differentiating tissue or a tissue where basal protrusions mediate lateral inhibition. In a randomly differentiating tissue (where basal protrusions extend but do not signal) we get no correlation between \( dt \) and \( dx \) (Spearmann’s Rho = 0.00185; Spearmann’s Rank test p-value = 0.502), and conditioning the distribution of \( dx \) on \( dt \) has no impact (Figures S7A–S7C).

On the other hand, simulations where differentiating cells extend signalling protrusions of wild-type length lead to negatively correlated \( dt \) and \( dx \) (Spearmann’s Rho = -0.10; Spearmann’s Rank test p-value < 10\(^{-10} \)) and the distribution of \( dx \) shifts to the right when we condition on \( dt \) much like the experimental data (Figures S6A and S6B versus S7D and S7E). Furthermore, when we plotted the mean \( dt \) for cells that differentiated within a specific space interval of one another we saw similar trends to those observed experimentally (Figure S7F versus Figure S6C).

The same simulations but with short signalling basal protrusions also led to negatively correlated \( dt \) and \( dx \) (Spearmann’s Rho = -0.12; Spearmann’s Rank test p-value < 10\(^{-20} \)), but with a weaker distribution shift for \( dx \) when conditioning on \( dt \) and a reduced range for lateral inhibition as in the experimental data (Figures S6D–S6F versus Figures S7G–S7L). Taken together, these results further support our hypothesis that the spatiotemporal dynamics of neuronal differentiation is contingent upon lateral inhibition mediated by the long and transient basal protrusions we see in vivo.

### Signalling at Soma-to-Soma (Figures 7 and S8)

Notch signalling typically occurs at soma-to-soma contacts between cells that are direct neighbours of one another (Lai, 2004). We therefore asked if soma-to-soma contacts could also play a role in our system. To investigate this, we run simulations that incorporate lateral inhibition at soma-to-soma contacts. This implies non-zero values for \( \alpha \) in Equation 3.

When signalling that occurs at all cell contacts is included (i.e. protrusion to soma and soma-to-soma) the predicted distribution differs significantly from that observed experimentally (Figures S8A and S8C; Kolmogorov-Smirnov test, p-value < 10\(^{-5} \)). The predicted and observed distributions are even more different when we assume that lateral inhibition is only mediated at soma-to-soma membrane (and not basal protrusions) contacts (Figures S8B and S8C; Kolmogorov Smirnov test, p-value < 10\(^{-10} \)). In fact, in this latter case the predicted mean value for \( dx \) is below that of a randomly differentiating tissue. This is because signalling taking place only at somal membrane contacts leads to differentiation events that occur in a typical checker-like pattern where cells that are one or two cell diameters apart tend to differentiate at a similar time (Collier et al., 1996; Hadjivasiliou et al., 2016). This can be seen by the peaks at \( dx = 30 \mu \text{m} \) in our histograms (Figures S8A and S8B). The absence of such a peak in our experimental data (Figure 6C in the main text) suggests that soma-to-soma contacts play a minimal if any role in the mechanism that determines the pattern of differentiation between spinal neurons.
Sensitivity Analysis of Simulated Data

In this section, we discuss the sensitivity of our simulated data and conclusions to variations in key parameters. We specifically explore the sensitivity of our conclusions to variations in parameters that determine the coupling between the basal protrusion growth dynamics and lateral inhibition. Parameters that determine feedback between Notch and Delta signalling have been explored in previous studies and we base our analysis on these published works (Collier et al., 1996; Cohen et al., 2010).

The simulation of neuronal differentiation in a randomly differentiating tissue is fully independent of any parameters. Our predictions were obtained using a random sampling algorithm on tissues of similar size and structure to the experiments (see Section 1.4). Therefore, our conclusion that the spatiotemporal dynamics observed experimentally are unlikely to come from a randomly differentiating tissue (p-value < 10^{-10} for wild-type data and < 10^{-6} for lamc1 data) is independent of any model parameters.

To explore the dependency between protrusion and differentiation dynamics, we varied three key parameters: the Hill exponent $q$, the speed of the protrusion growth determined by the duration of the protrusion extension period $T_{ext}$, and the upper limit for the probability of differentiation per simulation step, $p$. For each variation, we ran simulations as described in the Methods section and compared the simulated distribution of the distance between sequential differentiation events between the simulations and wild-type experimental data as in the main text using the Kolmogorov-Smirnov test.

Methods Image 4 shows the distributions for different parameters. We found that the comparison between simulation and experiment remains not significant so long as protrusion dynamics and the cellular decision to differentiate are tuned together.

Methods Image 4: Signalling simulations for various values of $p$, $q$ and $T_{ext}$ when lateral inhibition is mediated only through basal protrusions. Histograms of the distances between successive differentiation events ($dx$) in the simulations. The mean and s.d. of $dx$ are shown together with the p-value when the distribution was compared to the wild-type experimental data. Numbers in red indicate a significant deviation from the wild-type experiments. Simulations were repeated 100 times and simulation parameters other than the ones varied in this analysis are given in Table 1.

When the baseline probability of differentiation is very high (Methods Image 4C, F, I, L) or the extending basal protrusions are too fast (Methods Image 4A-C and A’-C’) the simulated distribution diverges from the experiment. Very high probability to enter differentiation per time (higher $p$) leads to a reduction in the average $dx$. On the other hand, fast basal protrusions together with a high Hill coefficient lead to more narrow $dx$ distributions with larger average $dx$. Furthermore, higher values for $q$ also lead to more narrow $dx$ distributions for the same values of $p$ and $T_{ext}$ (Methods Image 4A’-L’).

However, key features of the simulated distribution remain robust to these variations. In particular, the peak near $dx = 60\mu m$ and the skewed distribution away from small values of $dx$ are seen in all our simulations. Hence, this analysis suggests that the exact behaviour of the spatiotemporal dynamics depends on the coupling between the basal protrusion dynamics (actual speed of extension and retraction) and the initiation of cell differentiation as a response to levels of Notch signalling. The same should hold true in a real tissue: a weak dependency of differentiation on the basal protrusion dynamics and lateral inhibition would lead to weaker correlations in spatiotemporal dynamics of neuron differentiation.

We repeated the analysis now allowing lateral inhibition to take place both through basal protrusions and at membrane-membrane contacts. With this combination of signalling, nearly all parameter combinations we tested gave $dx$ distributions that deviate from the...
wild-type data (Methods Image 5). When we allowed very weak signalling at soma-to-soma contacts ($a = 0.01$) some of our simulations were not significantly different from wild-type simulations. Such small values of $a$ lead to membrane-to-membrane signalling is so weak it has a very minor impact on dynamics. These results suggest that soma-to-soma contacts may only contribute very weakly to lateral inhibition prior to protrusion extension.

Methods Image 5: Signalling simulations for various values of $p$, $a$ and $T_{ext}$ when lateral inhibition is mediated only through basal protrusions and soma-to-soma contacts. Histograms of the distances between successive differentiation events ($dx$) in the simulations. The mean and s.d. of $dx$ are shown together with the p-value when the distribution was compared to the wild-type data. Numbers in red indicate a significant deviation from the wild-type experimental data. Simulations were repeated 100 times and simulation parameters other than the ones varied in this analysis are given in Methods Table 1.

When cells only signal at their soma contacts the basal protrusion dynamics do not matter. To explore whether soma-to-soma only signalling could lead to wild-type-like distributions we run simulations varying $p$ and $a$. None of the simulated distributions were close to resembling the wild-type experimental data (Methods Image 6). As in the main text, we find a bias for $dx$ between 20 μm to 30 μm, the typical distance between cells that are two to three membranes apart. Weaker soma signalling (reducing $a$ to 0.01) led to distributions more similar to those of a randomly differentiating tissue. Therefore, our analysis suggests that the observed dynamics are unlikely to be due to lateral inhibition mediated at soma-to-soma contacts alone.

Methods Image 6: Soma-to-soma only signalling simulations for varying $p$ and $a$ when lateral inhibition takes place only at soma-to-soma contacts. Histograms of the distances between successive differentiation events ($dx$) in the simulations. The mean and s.d. of $dx$ are shown together with the p-value when the distribution was compared to the wild-type experimental data. Numbers in red indicate a significant deviation from the wild-type experiments. Simulations were repeated 100 times and simulation parameters other than the ones varied in this analysis are given in Methods Table 1.