Improving the understanding of cytoneme-mediated morphogen gradients by in silico modeling

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Abstract

Morphogen gradients are crucial for the development of organisms. The biochemical properties of many morphogens prevent their extracellular free diffusion, indicating the need of an active mechanism for transport. The involvement of filopodial structures (cytonemes) has been proposed for morphogen signaling. Here, we describe an in silico model based on the main general features of cytoneme-meditated gradient formation and its implementation into Cytomorph, an open software tool. We have tested the spatial and temporal adaptability of our model quantifying Hedgehog (Hh) gradient formation in two Drosophila tissues. Cytomorph is able to reproduce the gradient and explain the different scaling between the two epithelia. After experimental validation, we studied the predicted impact of a range of features such as length, size, density, dynamics and contact behavior of cytonemes on Hh morphogen distribution. Our results illustrate Cytomorph as an adaptive tool to test different morphogens gradients and to generate hypotheses that are difficult to study experimentally.

Key words: In silico modeling, Cytonemes, Drosophila Morphogenetic gradients, Hedgehog signaling, Cytomorph.
Introduction

During embryonic development, groups of cells are organized to give rise to tissues and organs. Precise spatio-temporal control of cell-to-cell communication is needed during proliferation, cellular three-dimensional organization and differentiation. Misregulation of these events is one of the most prevalent causes of diseases such as congenital malformations, neurological disorders and cancer (1). Several signaling molecules are defined as morphogens, messengers that are transported at a distance in a concentration-dependent manner to regulate the differential activation of target genes (2). The cellular mechanisms involved in the transport of the morphogens are still under debate (3).

Modeling has been a useful strategy to explore complex biological processes. Models to explain pattern generation during development have been mainly focused on the description of how, when and where a morphogenetic signal induces a specific cellular response within a particular tissue. Especially relevant were the early works of Turing (4) and Wolpert (5), who set the foundations of how a precise morphogen distribution could determine cell fate and patterning in a concentration-dependent manner. Subsequently, several works took into account the effect of the production and degradation of morphogens (6–9) and their transport was usually modeled by inferring a diffusion mechanism (10). Since the molecular properties of most morphogens impede them to diffuse freely in the extracellular environment, a different mechanism for their transport is required (11). A transport mechanism based on cytonemes (filopodia-like-structures) has been observed for most signaling pathways (12–15): Decapentaplegic (Dpp) (16,17), Wingless (Wnt/Wg) (18,19), EGF (20), FGF (20), Hedgehog (Hh) (21–23) and Notch (24–27). Cytonemes are actin-based membrane protrusions emanating from morphogen producing and/or receiving cells that deliver and/or collect morphogen by direct cell-cell membrane contacts (Fig. 1). Increasing experimental evidences by live imaging in
developing tissues highlight the implication of dynamic cytonemes in short- and long-distance cell communication (22,23,28–30).

A few mathematical models centered on different aspects of cytoneme-mediated signaling have been proposed (reviewed in (31)). They focus in characteristics such as vesicular transport along cytonemes (32,33), cytoneme contact mechanisms (34) or cytoneme guidance towards correct target receiving cells in a 1D system (35). To date there are also some models concerning the cytoneme-mediated establishment of morphogen gradient during pattern formation (36–38). Those models use static cytonemes and weight functions pondering the quantity of morphogen received. However, experimental evidence indicates that cytoneme dynamics can play an important role (22,23,28–30) and at present there are no data sustaining a pondered mechanism of signaling. Those models also assume a local source of morphogen, which is not true in most cases and theoretical studies emphasize the importance of using an extended source (39). Finally, previous models are not computationally implemented into a tool that can be used to test or load experimental data on cytoneme-mediated morphogen gradients.

In this work, we have developed a new dynamic model for cytoneme-mediated gradient formation in compartmentalized tissues during development, which validates this mechanism of cell signaling and has several advantages: 1) it was designed to be general enough to be applied to different morphogens or tissues, 2) it considers an extended morphogen source within a developing tissue, 3) the signaling is not based on weighted mechanisms, 4) it considers the dynamics of the cytonemes and 5) it was implemented into a computational tool (Cytomorph), which makes possible to develop in silico predictions for a variety of morphogens.

Finally, we tested and validated our model for the Hh signaling pathway in two different experimental paradigms of Drosophila: the wing imaginal discs and the
abdominal histoblast nests. We also made a preliminary analysis of the adaptability of Cytomorph making some in silico predictions of Dpp and Wg gradients in wing discs.

**Results**

**Theoretical framework: Mathematical model**

The morphogen gradient distribution is usually studied in biology as a spatial 1D function that can be generally determined as:

\[
\frac{\partial u(x,t)}{\partial t} = P(u, x, t) + T(u, x, t) - D(u, x, t) \quad \text{(Eq-1)}
\]

Where \(u\) is the concentration of a specific morphogen, \(P(u, x, t)\) is the production term, \(T(u, x, t)\) is the transport term and \(D(u, x, t)\) is the degradation term.

The effect of morphogen production and degradation on the gradient shape has been described in some modeling works (6–9). Here, we have focused on the transport mechanism; other terms were included using direct experimental data (see Supplementary Material). To model cytoneme-mediated morphogen transport, we focused in the main requirement of this type of signaling; the establishment of cell-to-cell membrane contacts for localized transmission. Thus, the core of our mathematical model is based on the determination of cytoneme contacts, specifically the contact distribution at receiving cells. Therefore, the transport can be determined as:

\[
T(u, x, t) = \alpha \cdot N(x_r, t) = \alpha \cdot \sum_{x_p=0}^{N_p} C(x_{p,r}, t) \quad \text{(Eq-2)}
\]

Where \(N(x_r, t)\) is the total number of contacts in receiving cells taking into consideration all the producing cells involved \((N_p)\). The \(C(x_{p,r}, t)\) is the contact function that defines the contact mechanisms for cytoneme signaling according to the position of producing \((x_p)\) and receiving \((x_r)\) cells.
Experimentally, three types of cell-to-cell contacts have been reported for cytoneme intercellular communication (Reviewed in (40,41)) (Fig. 1A-C):

Type 1: Cytonemes from receiving cells that contact signal-producing cell bodies (Fig. 1A).

Type 2: Cytonemes from signal-producing cells that contact receiving cell bodies (Fig. 1B).

Type 3: Cytonemes from both signal-producing and receiving cells that establish contacts (Fig. 1C).

Developing tissues are usually compartmentalized into two cell populations that divide the morphogenetic field in signal producing and receiving areas. Therefore, it is common to describe gradients using a spatial 1D frame of reference (Fig. 1D) in terms of the discrete cell positions \((x_{p,r} \in \mathbb{N}_0)\) in these areas.

To generate a model for the cytoneme mediated signaling observed experimentally, three different contact functions \(C_i(x_{p,r}, t)\) are defined in terms of spatial conditions as follows:

Types 1 and 2: In order to establish contacts, the distance between cells must be smaller than, or equal to, the length of the cytonemes.

Type 3: In order to establish contacts, the distance between a producing and its receiving cell must be smaller than, or equal to, the sum of the lengths of the cytonemes extending from these cells.

Which mathematically can be represented, in our frame of reference, as:

- Type 1: 
  \[C_1(x_{p,r}, t) = \begin{cases} 
  0 & \text{if } x_p \geq \lambda_r(t) - x_r \\
  \psi(\mu, x_r) & \text{if } x_p < \lambda_r(t) - x_r 
  \end{cases} \quad \text{(Eq-3.1)}\]

- Type 2: 
  \[C_2(x_{p,r}, t) = \begin{cases} 
  0 & \text{if } x_p \geq \lambda_p(t) - x_r \\
  \psi(\mu, x_r) & \text{if } x_p < \lambda_p(t) - x_r 
  \end{cases} \quad \text{(Eq-3.2)}\]
Type 3:

\[ C_3(x_{p,r}, t) = \begin{cases} 
0 & \text{if } x_p < \lambda_r(t) - x_r \\
\psi(\mu, x_r) & \text{if } \lambda_r(t) - x_r \leq x_p < \lambda_r(t) + \lambda_p(t) - x_r \\
0 & \text{if } x_p \geq \lambda_r(t) + \lambda_p(t) - x_r 
\end{cases} \quad (\text{Eq-3.3}) \]

These equations describe the contacts between a receiving cell at position \( x_r \) and a producing cell at position \( x_p \), depending on the temporal dynamics of cytoneme length \( \lambda_r(t) \) [\( \lambda_p(t) \)] and the probability of contact \( \psi(\mu, x) \).

\( \lambda_r(t) \) and \( \lambda_p(t) \) describe the dynamics of elongation and retraction of cytonemes emanating from either receiving cell position \( x_r \) or producing cell position \( x_p \). The dynamics observed in experimental data (30) showed that cytonemes not only can elongate and retract (named as Triangular behavior by (30) ) (Fig. 1E), but they can also have intermediate stationary phases, during which the cytonemes maintain their maximum elongation (named as Trapezoidal behavior by (30) ). (Fig. 1F). Therefore, in our model we mathematically defined \( \lambda_r(t) \) [\( \lambda_p(t) \)] considering the experimental results of the more basic cytoneme behavior (Triangular dynamics) but also including the possibility of a stationary phase (Trapezoidal dynamics) (see Supplementary Materials).

The function \( \psi(\mu, x) \) determines if there are contacts between cytonemes satisfying the minimum distance condition in a specific place \( x \) with a probability of \( \mu \). This function takes the value of 1 in the case of establishing contact (see Supplementary Materials). Since the cellular mechanisms to create a contact are starting to be elucidated (42), but not fully understood, here we used a probabilistic approximation to the real mechanism of signal transfer.
Computational framework: Model implementation

The general design of our model allows its application to different cytoneme-mediated morphogen gradients. Thus, to take full advantage of this approach, we created a Matlab-language-based software called Cytomorph, in which it is possible to simulate different experimental data and test different hypotheses in silico.

Cytomorph was designed to introduce different inputs (experimental data and variables under study), compute in different modules the dynamics of cytonemes and their contacts and, finally, the morphogen distribution using previous equations of the model (Fig. 2 shows a general workflow of the software). For practical purposes, in what follows we will focus on the inputs and outputs of the model. A detailed schematic view of the steps of the Cytomorph machinery can be found in Supplementary Figure S.1.

Cytomorph inputs:

Cytomorph was designed with inputs divided into two sets (Fig. 2A). The first set is loaded to Cytomorph via a spreadsheet (Fig. 2A-1) encompassing all experimental distributions of cytoneme lengths and temporal distributions of elongation, retraction and stationary phases during cytoneme dynamics (See Supplementary Table S.1). The second set is loaded via a graphical user interface (GUI) (Fig. 2A-2, see also Supplementary Figure S.2). This group comprises: 1) average experimental values (e.g. cell size in a tissue and velocities of elongation and retraction), 2) parameters difficult to measure with the current experimental techniques (e.g. the contact probability and the temporal contact dynamics) and 3) features or parameters without experimental data but which are required for morphogen simulations (e.g. the number of cells needed for gradient formation and the role of cytoneme dynamics). A detailed illustration of features and parameters of the second group are described in Supplementary Figure S.2.
Cytomorph modules:

For an intuitive use of Cytomorph we designed a GUI to run simulations (Fig. 2A,B) and Cytomorph was subdivided in different scripts and modules of the next three types: 1) A group of scripts for the GUI. 2) Modules to numerically simulate the cytoneme dynamics and to compute contacts and their spatial distribution over time. 3) A module to plot the simulated contacts together with different gradient properties (see Supplementary Materials).

Cytomorph outputs:

To study cytoneme features and assess their role in the Hh gradient formation, Cytomorph was implemented to analyze different characteristics:

- Contact distribution: The contacts per cell along simulations (Fig. 2C-1); violin plots are shown (Fig. 2C-2) to visualize the contact distribution along simulations.

- Signal variability: To study the predicted in silico variability we computed the distribution of coefficients of variation per case (Fig. 2C-3).

- Temporal evolution: To observe the number of contacts in each receiving cell per time lapse (Fig. 2C-4) and the total evolution of contact distribution and gradient shape over simulated signaling time (Fig. 2C-5).

- Gradient distribution: Assuming that each contact transmits the same amount of morphogen ($\alpha = \text{const}$ in Eq-2), the distribution of the morphogen $u(x, t)$ can be estimated through the $N(x, t)$ calculated in the model (Fig. 3C-6).

A detailed description of how the outputs were computed and calculated can be found in Material and Methods and Supplementary Materials sections.
**Experimental framework: Model validation**

To validate Cytomorph we used the formation of the Hedgehog (Hh) gradient in two different *Drosophila* tissues: the imaginal wing disc and the abdominal histoblast nest. The latter has been used to study cytoneme dynamics as it easily allows *in vivo* imaging. Both tissues have the same cell distribution, in which the Hh producing region (Posterior (P) compartment) signals over the receiving region (Anterior (A) compartment).

We first characterized and quantified the biological magnitudes needed as inputs for Cytomorph simulations. For length characterization, we overexpressed Ihog, a transmembrane protein and co-receptor of the Hh pathway present in all epithelial cells, since its overexpression stabilizes cytonemes without affecting their length (30). This effect on cytoneme dynamics makes Ihog overexpression a good tool for cytoneme visualization at the basal side of fixed tissues, such as the wing imaginal disc (Fig. 3A). Abdominal histoblast nests keep the same cell and cytoneme distributions in A and P compartments (Fig.3B and movie 1) as well as the same apico/basal polarity (Fig. 3C). For the wild type dynamics of cytonemes, we used markers that do not affect cytoneme dynamics (Life-actin-RFP and mCD8-GFP) that we simultaneously overexpressed using the binary systems UAS-Gal4 and QUAS-QF, allowing the *in vivo* visualization of both receiving (from A cells) and producing (from P cells) cytonemes (Fig. 3C and movie 2).

Looking at the quantified length of cytonemes in the wing disc we observed a statistically significant difference, receiving A cytonemes being shorter than producing P cytonemes (Fig. 3D). In addition, comparing both receiving and producing wing disc cytonemes with those of the abdominal histoblast nests (30), we observed that the former are significantly longer than the latter (Fig. 3D). We also quantified the cell size ($\phi$) in both tissues and found a difference in cell size: $\phi = 3.05\pm0.65 \, \mu m$ in wing discs and $\phi =
4.37±0.89 μm in abdominal histoblast nests (see Materials and Methods for the measurement protocol and the statistical study of this average values).

To quantify the Hh experimental gradient (and its experimental signal variability) as a validation for the model-simulated gradient profile, we analyzed the signal intensity of the endogenous Hh using a GFP fluorescent reporter (Hh:GFP BAC) in the two selected Drosophila tissues. Then, to compare the Hh gradient responses between different samples (see Materials and Methods and Supplementary Figure S.3 for details), we also used a genetic tool (EnhancerPtcRed) that allows simultaneous visualization of Hh and the transcriptional response of its receptor Patched (Ptc) (Fig. 4A). After statistical analysis, we found that, despite the tissue similarities, the Hh gradients are not identical (Fig. 4B), with the gradient decaying faster in abdominal histoblast nests than in the wing imaginal discs. It is important to mention that in our cytoneme model the shape of the gradient is a consequence of the contact distribution (Supplementary Figure S.4) and this distribution is due to cytoneme dynamics and cytoneme distribution along the tissue, which are elements experimentally settled. In addition, we observed that the range of the Hh gradient in both tissues can be determined by the sum of the maximum cytoneme lengths, emphasizing again the importance of cytonemes as mechanism for gradient formation.

The scaling of Hh gradients between different tissues provides an opportunity to study the adaptability of our cytoneme model. Studying the parameter space after loading the experimental data (length, temporal dynamics of cytonemes and cell size $\phi$), the in silico simulations have shown that our model is able to predict the shape of Hh gradients in the two tissues (blue fitted curve in Fig. 4 C,D). In this way, we demonstrate the ability of the model to adapt to different biological conditions and correctly forecast the signaling gradient. The parameter space has been selected as a reference case for further simulations.
(Supplementary Table S.2). More information can be deduced when analyzing the parameter space used to fit the gradient. For example, model simulation for the Hh gradient in the wing disc agrees with the cytoneme contact type 3, which fits with the experimental data. Thus, modeling emphasizes the importance of the direct cytoneme-cytosome (cyt-cyt) interaction for the correct development of the wing disc Hh gradient, so far assumed but not demonstrated. As for the gradient in the abdominal histoblast nests, a lower probability for cyt-cyt interaction fits better with the experimental gradient, indicating that, in contrast with the wing disc tissue, in abdominal histoblasts, cyt-cyt interaction is not as critical as cytoneme to cell body contact.

Since many theoretical models still assume free diffusion as the mechanism for morphogen transport (10), we next compared our cytoneme model predictions with those of the classical diffusion-degradation model (see Supplementary Materials), and then both with the experimental gradient measurements. Interestingly, in the case of the wing disc, both model predictions fall statistically within the experimental variability (Fig. 4E), with the cytoneme model slightly closer to the experimental mean. However, this is not the case for the abdominal histoblast nests where the diffusion model does not adapt and is not able to predict the gradient as accurately as the cytoneme model does (Fig. 4F, black). Indeed, the diffusion model requires the assumption of a three times smaller diffusion constant to fit simulations with experimental data in abdominal histoblast nests (Fig. 4F, red).

**In silico framework: Model predictions**

The model can be used to study the effect of different parameters on gradient formation and to predict their biological implications. As examples, we have selected two parameters from which we could obtain experimental data (cytoneme length/cell size ratio and number of producing cells) and another parameter lacking quantified experimental...
data (number of cytonemes per cell). We then used Cytomorph simulations to predict their effects and values.

- **The ratio between cytoneme length and cell size** is a default unit used in the software to intuitively visualize the extent of cytonemes. *In silico* simulations showed that this ratio seems to be responsible of controlling the shape and length of the morphogen gradient (Fig. 5A). Although this ratio also affects signal variability, this is not statistically significant in most cases (Fig. 5A’). Therefore, after simulation we can conclude that both, length of cytonemes and cell size, are key to understand how cytoneme signaling defines the shape and extension of the gradient (Fig. 5A’’).

- **The number of signaling source cells** has been previously suggested to be crucial for plausible gradient formation modeling (39) and experimental data also indicated this importance for Hh signaling (43). Therefore, we decided to analyze the effect of this parameter in our model. We have observed that, starting count from the A/P compartment border, the first 5-7 rows of producing cells are key in shaping the Hh gradient (Fig. 5B), while the next producing cell rows (rows 8-10) refine the gradient shape lowering the variability. Finally, increasing the number of Hh producing cells to more than 10 cell rows does not affect the Hh gradient. This can be observed in both the amount of morphogen and the signal variability (Fig. 5B and B’). Therefore, the number of producing cell rows is indeed important in the scaling and determination of the gradient shape in the wing imaginal disc (Fig. 5B’’). This dependence on the size of the producing region can be extrapolated to other tissues, as suggests our simulations in the abdominal histoblast nests (Supplementary Figure S.5). In contrast, analysis of the number of receiving cell rows did not show any effect on Hh distribution (see Supplementary Figure S.6).
The density of cytonemes (the number of cytonemes per cell) is key for the amount of morphogen distributed (Fig. 5C) and significantly affects signal variability (Fig. 5C’), but it is not determinant for the gradient shape (Fig. 5C’’). Counter-intuitively, experimental variability can be estimated in silico (error bars versus green shaded area in Fig. 5A) for a low number of cytonemes per cell, which is what can be inferred from wild type (Fig. 3C). These results are of biological importance because they indicate that the shape of the gradient is mainly determined by the behavior of cytonemes and not by their number.

Hypotheses based on cytoneme behaviors

Cytomorph is an adaptable tool devised to answer different questions and test hypotheses on cytoneme mediated signaling. Since most of our working hypotheses are related to contact dynamics, we will next focus on how the three types of contacts (Fig. 1A-C) might affect gradient features. Types 1 and 2 can be considered mathematically the same, while type 3 should have an additional contribution, as both presenting and receiving cells emit contacting cytonemes. Our model predicts a significant effect in the gradient when considering the type 3 cytoneme contacts compared with those of types 1 or 2, as we found that the amount of morphogen (Fig. 6A) and the length of the signal (Fig. 6A’) were doubled. Type 3 seems then to be the most probable situation for Hh gradient formation (Fig. 6A), although types 1 and 2 can still be functional forms for other signaling situations.

Further analysis of contact dynamics properties using Cytomorph has also allowed the study of the contact effect for reception, which is currently not well understood due to the difficulty of approaching it experimentally; it was defined in the model by the
contact probabilistic function $\psi(\mu, x)$. In particular, we implemented in different Cytomorph modules three working hypothesis of the contact function $\psi(\mu, x)$:

1. A contact dynamic in which the probability of contact only depends on the condition that cytonemes are close enough; then the probability to contact is $\psi = \psi(\mu)$.

2. A contact dynamic in which different contacts can be established along the overlapping cytoneme membranes. This multiple-contacts approach was designed by the special contact function $\psi = \Psi(\mu)$.

3. A contact dynamic in which, in addition to the previous distance condition, the cell position is important and can be treated as a variable $\psi = \psi(\mu, x)$.

Comparing the first two hypothesized contacts, the *in silico* simulations showed that the overlapping multiple contacts function can significantly change the number of contacts and subsequently the amount of morphogen transferred (Fig. 6B), also resulting in significant changes over signal variability (Fig. 6B’) and gradient shape (Fig. 6B’’). Similarly, comparison between cases 1 and 3 showed statistically significant changes in the number of contacts (amount of transmitted morphogen) and signal variability when including cell position as a variable (Fig. 6C and C’). Furthermore, after the analysis of different scaling across receiving cells we could also infer that gradient distribution was affected (Fig. 6C’’), with case 3 showing a faster and more linear decay as a consequence of its dependence on cell position (see Supplementary Materials).

Since our results suggest that the contact probability function only depends on the variable $\mu$, we carried on an *in silico* study to test the impact of this variable over gradient features. Interestingly, model simulations for different values of $\mu$ only significantly contribute to the amount of morphogen transferred (Fig. 6D), but they do not disturb neither the variability nor the shape of the morphogen gradient (Fig. 6D’ and D’’).
Cytoneme dynamics in Hh gradient evolution

To this point we have validated Cytomorph in steady state conditions (Fig. 5C and D), and from now on we will test its capability to study temporal aspects during gradient formation. For this purpose, we performed fluorescence recovery after photobleaching (FRAP) experiments in the abdominal histoblast nests. In this tissue, the gradient is established previous to histoblast migration and it allows the dynamic characterization of the Hh signaling gradient by in vivo recording (movie 3).

Previous to photobleaching a reference Z-stack was taken and the signal was then bleached to 80-90% of the initial maximum value (Fig. 7A); recovery was then recorded in a Z-stack every 45 seconds. To automatize the acquisition of the gradient profile, a FIJI macro was written. The results showed that the Hh gradient recovered up to 92% of the initial value in less than 50 minutes (Fig. 7B), while the receptor graded response (EnhancerPtcRed) presented a 64% recovery in the same time. (Fig. 7C). This difference in the percentage of recovery was expected, as there is a delay in the reporter response, which requires both transcription and translation. To validate the temporal evolution of the model, we then simulated the Hh gradient in histoblast nests using Cytomorph (Fig. 7D) and compared the predicted with the experimental curves. As we can observe in Fig. 7E, the temporal prediction and the experimental signal recovery are the same, corroborating the capacity of Cytomorph to predict the temporal evolution of the gradient.

Cytomorph could also be used to solve pending questions regarding cytoneme dynamics. We were particularly intrigued by how each type of the observed dynamics (30) might affect the shape of the gradient. To analyze whether each cytoneme behavior represented or not an advantage for the distribution of the gradient and its variability, we modified the fraction of the cytoneme population having each behavior and simulated how these changes could affect the gradient (Fig. 7 F and F’). It was interesting to find
out that the impact was stronger on the tail of the gradient when considering that cytonemes simply elongate and retract (triangular behavior), while the impact was greater near the source of morphogen (Fig. 7F’’) when considering that all cytonemes have a stationary phase between elongation and retraction (trapezoidal behavior).

Although the experimental data showed the importance of the dynamics of cytonemes for the formation of gradients (22,23,28–30), we also included in the model the case in which all cytonemes were static. In fact, our simulations showed that dynamic and static cytonemes generate different gradient shapes (Supplementary Figure S.7). This is noteworthy since many theoretical models do not consider this temporal aspect and our results indicate that using dynamic cytonemes the predicted gradient better fits the experimental data (Supplementary Figure S.7). It should be remembered that the temporal dynamics of the contacts is not yet well defined and, furthermore, it is experimentally difficult to unravel how the dynamic contacts coordinate with growth during gradient formation. Cytomorph can be used to study plausible biological scenarios for Hh signaling, even in the absence of experimental data. In fact, simulations of contacts and growth (Supplementary Figure S.7) allow to study the dynamic of signaling and reflect the complexity in the coordination of different cytoneme features in the formation of a gradient.

**Robustness of cytoneme signaling**

Although failures can occur during the development of organisms, cell signaling has been shown to have a robust control mechanism (44–46) in which a combination of parameters can compensate for a possible developmental failure. In this context, Cytomorph can also serve to identify compensation mechanisms; as an example, we have found an interesting interaction between the number of producing cells and the density of cytonemes per cell: a reduction in the number of producing cells can be compensated by
an increase in the number of cytonemes per cell (Supplementary Figure S.8). Both predicted gradients fall within the experimental variability, creating a “functional” gradient that should be able to activate the same target genes.

**Cytomorph as a predictive tool for other morphogen gradients**

We have proposed Cytomorph as a computational tool that could be used to simulate the formation of cytoneme-mediated gradients for different morphogens. To test this hypothesis, we made a simple approximation using Cytomorph to simulate Dpp and Wg signaling pathways in the wing imaginal disc.

Dpp signaling has been described to form a gradient mediated by cytonemes in *Drosophila* wing discs (16,47) and it gradient has been highly study (48–53). In this system, Dpp is produced by the first cell rows of the A compartment (~7 cell rows) and forms two morphogenetic gradients at both sides of the wing pouch as Dpp spreads across A and P compartments.

Using the described cytoneme characteristics in the imaginal wing disc, we simulated Dpp gradient in P compartment cells and compared it with the experimentally quantified Dpp gradient taken from (51). Our results correctly predict the shape of the Dpp gradient (Supplementary Figure S.9) and allow to extrapolate some predictions for Dpp cytoneme behavior. Specifically, Cytomorph simulations suggest that for Dpp signaling cytonemes elongate from both producing and receiving cells, as in the case for Hh signaling, reproducing the described Dpp gradient extension and shape (51). However, our simulations predict that instead of cytonemes establishing contacts while growing, as in the Hh pathway, they are more likely to establish contacts after elongation (Supplementary Figure S.9).
Although the Wg gradient has not yet been shown to be mediated by cytonemes in the wing disc, there are evidences of the involvement of cytonemes in Wg signaling in other *Drosophila* tissues (18) and in vertebrate system (19). In wing disc, Wg signaling satisfies the cell distribution assumed by Cytomorph: cells at dorso/ventral (D/V) compartment border of the wing pouch are the source of Wg, which spreads and signals to both D and V compartments. Using the cytoneme length in the imaginal wing disc described here, we estimate that the expected length for the Wg gradient according to eq-3 is 42 μm (~14 cells). This prediction for Wg gradient length is in agreement with the experimental data for the extension of the Wg signaling gradient (described by the extracellular detection of the Wg protein), which is up to 40 μm in imaginal wing disc (54).

**Discussion**

In this work, we present a general *in silico* model for morphogen gradient formation considering that the morphogen dispersion is mediated by cytonemes. In particular, we demonstrate that this model validates a mechanism of cytoneme-mediated Hh gradient that can be extrapolated to other morphogens such as Wg and Dpp. We have implemented our model in an open computational software (Cytomorph), which allows the introduction of experimental data to study the role of different biological parameters. With this approach we try to overcome the previous lack of connection between theoretical models and experimental data in cytoneme mediated cell signaling. To improve our understanding of how specific cytoneme features impact the gradient properties, Cytomorph is able to plot results in graphs showing the final shape of the morphogen distribution, the number of contacts, the signal variability, the time course and the
gradient scaling. To facilitate the use of this tool, we also designed a GUI that allows straightforward control of software commands.

Model validation

To experimentally validate the model and its adaptability to real gradient predictions, we studied the Hh gradient formation in two different Drosophila tissues: wing imaginal discs and abdominal histoblast nests. Using different genetic tools, we experimentally quantified several parameters in both tissues, such as the length of cytonemes, the cell size and the Hh gradient distribution. Cytomorph was able to predict the Hh scaling and correctly simulate the signal gradients in both tissues, emphasizing the involvement of cytonemes for a correct signaling. Although the quantified gradient scaling in these two tissues had not been previously characterized, we expected them to be different because, despite their similarities, both systems have different behaviors: while the wing disc is an expanding but static epithelium, the abdominal histoblasts divide and migrate simultaneously reducing, for instance, the probability of cyt-cyt contacts, as our model suggests.

Cytoneme model versus diffusion model

The diffusion model is still the mathematical model most commonly used in biophysics, although the biochemical properties of most morphogens argue against their transport via Brownian motion. Comparing our cytoneme model with the classic diffusion-degradation model, we found that our model thoroughly predicts the shape of the Hh gradient in two different tissues, the wing imaginal discs and the abdominal histoblast nests. The diffusion model, however, required a readjustment of the diffusion constant to predict the Hh gradient in the abdominal histoblast nests. It is not clear why the diffusion constant has to be different for the same protein in similar epithelial tissues. Nevertheless, it is
important to point out that the measurement of the diffusion constant is an effective parameter that summarizes a collective behavior and does not give information regarding the transport mechanism involve. Actually, it has been described that the diffusion coefficients can significantly vary depending on the morphogen, the tissue and the experimental approach (55).

**In silico study of cytoneme features**

After the experimental validation of Cytomorph, we studied *in silico* different aspects of cytoneme-mediated signaling as a way to understand the role of cytoneme features and to be able to generate hypotheses regarding this signaling mechanism. We initially tested the cytoneme length/cell size ratio, a parameter for which we already had experimental data, and our simulations suggested that it is a crucial parameter for the Hh gradient scaling but not for the variability of the signal. This model prediction was experimentally supported in abdominal histoblast nests and in imaginal wing discs.

Theoretical analysis (39) emphasizes the importance of considering an extended source to predict realistic gradients, however previous models do not take this element into account. Therefore, we used Cytomorph to clarify the effect of an extended source in shaping the gradient. The resulting simulations gave a detailed description of how the gradient is affected by changing the number of cell rows involved in the production of morphogen in a tissue.

Cytomorph also provides the possibility to analyze the effect on gradients of parameters for which there are no experimental data, such as the number of cytonemes per cell. Interestingly, our results suggest that this particular parameter is key for both the variability of the signal and the amount of transmitted morphogen but not for the
distribution or the scaling of the gradient. Moreover, simulations also allowed to estimate
the probably low number of cytonemes per cell.

In parallel with cytonemes parameters we have also studied other features of the
cytoneme-mediated signaling. By observing the effect of different types of signaling
contacts, our model predicts that the type 3 will be different from types 1 and 2, since the
amount of transmitted morphogen and the length of the gradient increase due to cyt-cyt
contacts. In agreement with experimental observations, our in silico results have shown
that type 3 cytoneme interaction is the most likely situation for Hh gradient formation in
the wing imaginal disc.

Previous approaches to cytoneme signaling used weight functions, with a
dependence on cell position, to ponder the quantity of received morphogen. To ascertain
if this dependence is required, we tested three different hypotheses for the contact
function $\psi(\mu, x)$. In contrast with previous approaches, our model suggests that this
contact probability does not require a pondered mechanism based on cell position, since
the simplest case $\psi = \psi(\mu)$ fits the gradient distribution better than $\psi = \psi(\mu, x)$.
Besides, the existence of multiple contacts between cytonemes ($\psi = \Psi(\mu)$) fits better the
experimental data for Hh signaling. Since Cytomorph predicts a low number of
cytonemes per cell, what is important in determining the shape of the gradient is the
distribution of contacts across the receiving cells. This conclusion is in agreement with
the results of other cytoneme-mediated morphogen distributions (56).

Finally, we studied the effect of the probability of contact $\mu$ in the gradient
properties. Our results showed that this parameter has a significant impact on the amount
of morphogen transmitted but not on the signal variability or the gradient shape.
Cytoneme dynamics in Hh gradient formation

One of the main advantages of our model was the inclusion of the temporal dynamics in the equations, which has been experimentally found to play a significant role in the correct activation of the target genes during development. To validate the temporal dynamics of our model we performed experiments to study the temporal recovery of the Hh signal after photobleaching (FRAP technique), and then compared this data with our model simulations. Comparison of the predicted and the experimental gradient curves proved that our dynamic model is able to simulate physiological temporal features.

We then used Cytomorph to study the role of the two types of cytoneme behavior (Triangular and Trapezoidal) observed experimentally. Our simulations allowed us to generate the hypothesis that the two cytoneme dynamic might have a distinctive impact on specific regions of the Hh gradient, suggesting the importance of these cytoneme behaviors on the precise spatial control of the gradient shape. Besides the generation of this new hypothesis, our simulations also show that a population half Triangular and half Trapezoidal cytoneme dynamics fits better the experimental data, what happens to be the proportion found experimentally (30).

Our simulations point out that static and dynamic cytonemes give rise to quite different Hh gradient. Consequently, this characteristic of cytoneme signaling should be included in theoretical models to study cytoneme signaling. Moreover, cytoneme temporal dynamics could provide robustness to the progressive establishment of signaling gradients, an advantage for both growing (wing imaginal discs) and migrating while growing systems (abdominal histoblast nests). Static cytonemes are less likely to adapt to tissue changes, increasing the probability of failure, while dynamic cytonemes can allow constant regulation of the gradient shape throughout development. Nevertheless, other
static tubular structures, such as tubulin-based channels, could be significant for other morphogens or biological models \((57)\).

The *in silico* model also emphasizes the role of different features in the gradient properties and, more importantly, predicts that the shape of the gradient is a consequence of the contact distribution; in turn, this distribution is due to cytoneme dynamics and cytonemal distribution along the tissue. In fact, the model suggests that during the correct establishment of graded distribution, cytoneme dynamics are more critical than the amount of available morphogen from producing cells. This hypothesis has been recently corroborated experimentally for several morphogens. The analysis of the dispersion of Hh, Wg, and Dpp in the *Drosophila* wing imaginal disc indicates that their delivery to target cells is regulated since an increment in their gene doses does not alter the extent or shape of their gradients \((58)\). For Hh signaling, the receiving cells take up less than the 5% of Hh produced; under conditions of Hh production up to 200% of the normal amount, neither the protein uptake nor the extent of the gradient changes. These findings show that the amount and destination of delivered morphogens are regulated, in agreement with a cytoneme model but not with a diffusion model.

**Other uses of Cytomorph**

Based on uncomplicated mathematical premises, our model improves the understanding of cytoneme signaling mechanisms. Although our interest along this work has been focused on identifying individual roles for different cytoneme parameters that could affect the morphogen gradient formation, Cytomorph was also able to detect interactions or compensation mechanisms between cytoneme features able to counteract malfunctions, emphasizing the robustness of the cytoneme model as a signaling mechanism.
Finally, we propose that Cytomorph can be adapted to different biological systems and morphogens. As a prove, we used Cytomorph to simulate Dpp and Wg signaling. Although this has only been an initial attempt to verify its use for other morphogens, we have shown that the simulated Dpp gradient reproduces the experimental gradient and that the length of Wg gradient can accurately be estimated with our model equations. However, each morphogen and tissue has its own characteristics. For instance, in both Wg and Dpp gradients some of the signal-producing cells are also part of the receiving cells which is not the case in the Hh pathway. Therefore, for a precise analysis of the shape of these gradients, it would simply be necessary to implement the software with adequate computational modules. This adaptability is achieved by the modular architecture used to design the Cytomorph software, in which new modules can be added to adjust the particularities of each system or to test emerging biological hypotheses, as soon as new findings or experimental conditions become available.

Material and Methods

Experimental methods

_Drosophila_ lines

_Drosophila melanogaster_ stocks were maintained according to protocols described in Ashburner manual (59). Crosses were maintained at 18°C until the time of gene expression induction. The description of mutations, insertions and transgenes is available at Fly Base (http://flybase.org).

The following drivers were used to induce ectopic expression using the Gal4/UAS (60) and QUAS-QF (61) systems: tubGal80ts (Bloomington Drosophila Stock Center, BDSC),

Overexpression stocks: The pUAS-transgene strains used were: UAS.ihog-YFP (64) and UAS.LifeActinRFP (BDSC 58362). The QUAS-transgene strains used were: QUAS.mCD8-GFP (BDSC 30002).

Other stocks: EnhancerPtcRed (Kyoto stock center, DGRC 109138) and Hh:GFP BAC (65).

**Experimental data acquisition and quantification in wing imaginal discs**

Laser scanning confocal microscopes (LSM700 and LSM800 Zeiss) were used for confocal fluorescence imaging of imaginal discs. Fluorescence signal of Hh:GFP BAC protein and EnhancerPtcRed reporter were obtained using 40 x magnification and taking Z-stacks with a step size of 0.7-1 μm. Fiji software (ImageJ software, National Institutes of Health) was used for image processing and analysis.

- **Filopodia extension**

  Cytonemesis were labeled overexpressing UAS-Ihog-YFP in either Hh-Gal4 (P compartment) or Ptc-Gal4 (A compartment) domains for 24-48h before dissection. The length extension of cytonemesis was manually measured using the Straight tool from FIJI software. The statistical analysis and software simulations were done using a total of 984 cytonemesis, 729 in the P compartment and 255 in the A compartment.

- **Cell diameters for gradient normalization**

  Since the software computes the data in cell diameters, to compare the experimental data with model simulations, it is important to know the characteristic cell diameter in μm of each specific tissue studied. For the normalization of the gradient length, we manually measured approximately 100 cells along the X-axis in each wing imaginal disc (n=19).
- **Hh gradient imaging in wing discs.**

Hh protein gradient and the graded response of *ptc* enhancer reporter were measured experimentally using Plot profile tool of FIJI taken an average Z-stack projection to get all the morphogen distribution along apicobasal sections of the wing disc epithelium. The fluorescence profiles of the corresponding channel for the *Hh:GFP BAC* and *EnhancerPtcRed* signals were measured in a 90x35 μm² region of the A compartment with the start positioned at ≈25 μm from the A/P border inside the P compartment.

- **Mathematical protocol for the Hh gradient data.**

The Hh protein gradient and *ptc* enhancer reporter gradient response in wing discs and their experimental variabilities were estimated using 19 different wing disc samples, as follows: the background was estimated measuring the mean signal level over a 20 μm region in areas in which each reporter genetic tool is not active; for the *ptc* reporter signal, the region corresponds to the entire P compartment, while for Hh protein the region corresponds to the A compartment cells located away from the A/P compartment border. After subtracting the background, the intensity was normalized with the mean of the maximum intensities (3 values for the region of maximum *ptc* enhancer reporter expression and the whole P compartment signal for Hh protein levels). Finally, to compare the resulting data, we translated the measured profiles to the same reference origin; for the beginning of the Hh gradient we used the A/P compartment border. This origin was mathematically estimated using the well-defined sharp increase in *ptc* expression at the A/P compartment border. The graphic steps of the process are depicted in the Supplementary Figure S.3.
Data acquisition and quantification for in vivo imaging in abdominal histoblast nests

Pupal abdominal histoblasts imaging was performed in a chamber to seat and orient the pupae to look under the microscopy as described in (29). The dorsal abdominal segment A2 was filmed using 40x magnification; Z-stacks of around 30 μm of thickness with a step size of 1.1-1.3 μm were taken using a LSM800 confocal microscope. The overnight movies of the Hh-GFP gradient and the ptc-RFP enhancer graded response during the histoblast migration (movie 3) were done recording Z-stacks every 2 min. For optimal recording of dynamic cytonemes, in vivo experiments using at the same time Gal4 and Q systems were taken in different conditions (Z-tack of 18.5 μm with a step size of 0.5 μm every-one minute). Also movie 2 (see some sequences in Fig.3) was computationally treated using a deconvolution method (Huygens software) for cleaning the fluorescence signal.

- Cell diameters for gradient normalization.

The cell diameters for Hh and ptc profile normalizations were measured for 14-32 different histoblasts per pupae along the X-axis (n=9). Since we did not find statistical differences in cell diameters between A (ϕ_{anterior} = 4.285 ± 0.886 μm with n=228) and P compartment cells (ϕ_{posterior} = 4.453 ± 0.887 μm with n=245). We used the average value (ϕ = 4.37 ± 0.89 μm, n=473) for abdominal histoblast simulations.

- Imaging Hh gradient in abdominal histoblast nests.

The Hh protein gradient and ptc enhancer reporter signals were measured using the Plot profile tool of FIJI in an average Z-stack projection, as we have done for the wing imaginal disc samples. In each channel, profiles were measured for the same region of 200x130 pixels (51.51x33.48 μm²) located in the A compartment close to the A/P compartment border.
- Mathematical protocol for Hh Gradient data.

The experimental variability of Hh signal and its gradient in abdominal histoblast nests were estimated using 14 different regions (in the A compartment close to de A/P border) extracted from 9 pupae. In each sample, the background signal was subtracted using the corresponding minimum value and then the intensity was normalized with the maximum value in each case. Finally, the profiles were translated to the same position using the maximum as a reference.

- Statistical analysis of filopodia extensions

To study the parametric behavior of the data, we first performed a Shapiro-Wilk normality test. After testing the non-parametric condition of the experimental distributions, we studied the significance using the Wilcoxon rank Sum test of homogeneity of variances (Implemented in Matlab2015a).

- Data analysis of cytoneme dynamics

The experimental data of filopodia dynamics have been taken from previous studies (30). Here, we have statistically studied in R language the differences between the times of the elongation, retraction and stationary phases of Triangular and Trapezoidal behaviors using a Kolmogorov-Smirnov statistical analysis (Supplementary Table S.3), as we observed statistically significant differences between Triangular and Trapezoidal times we develop the model to consider both behaviors.

Analysis of the Hh gradient formation by FRAP experiments

Fluorescence recovery after photobleaching (FRAP) is a method to study temporal evolution of fluorescent signals. An initial z-tack covering from apical to basal sections of the tissue was performed using Zeiss-LSM800 confocal microscopy to record the pre-bleaching conditions of the sample. To avoid damaging the tissue, the photobleaching
was done over a ROI of (48.7x61.3 μm²) in a region located at the A/P compartment border (we used ptc expression as a reference for the A/P compartment border (Fig.7A)). Photobleaching of the abdominal histoblast nests was done by series of short expositions of 488 nm laser at 100% intensity until the signal at that z-plane reached less than 10% of the initial value. Since the Hh signal is present through apicobasal length of the tissue, we repeated the photobleaching conditions 7-10 times at different sections covering the total apico-basal tissue length. To obtain the Hh signal recovery over time we recorded the same region used in the pre-bleach z-stack conditions every 45 seconds immediately after photobleaching.

The acquired image samples of the photo-bleached ROI area were then treated with the imaging protocol described above (quantification of the wild type Hh gradient in abdominal histoblast nets). Since the resulting file is not a simple image but a temporal sequence of images, we automatized the process creating a macro script in Fiji that measures the Hh and Ptc profiles over time in the region where the photobleaching was performed. Since the experimental conditions can generate undesired photobleaching, a control of the Hh signal intensity in the P compartment was also measured each time. To study the FRAP recovery we used the previously described mathematical protocol that translates all signals to the same origin, but normalization was done using the P compartment control values as follow:

\[
\frac{\text{Intensity where FRAP was done} - \text{background signal}}{\text{control intensity} - \text{background signal}}
\]

This equation is used in FRAP experiments to mathematically remove the possible undesired photobleaching in the recovery measurements.

Finally, to visualize the recovery evolution of the pre-bleaching gradient in percentages, the resulting values were normalized to the pre-bleaching maximum value.
All the samples studied (n=6) showed the same recovery tendency for Hh and ptc expression profiles than the representative case shown in Figure 7 B and C.

**Theoretical and computational methods**

**Software code:**

Cytomorph was generated implementing the cytoneme model in Matlab language (MatlabR2015a). Since Cytoneme-mediated signaling has been reported for many different morphogens and for different animal systems, our goal was that Cytomorph could be used as a computational tool to help other scientist. We then decided to develop Cytomorph as an open source software under a 3-clause BSD FOSS license. The software code and a manual for users are available in the software repository:

[https://github.com/AdrianA-T/cytomorph](https://github.com/AdrianA-T/cytomorph)

The available version of Cytomorph has been divided into different modules that can be updated to incorporate new discoveries in the formation of gradients, these modules can also be remodeled to simulate specific requirements of the system under study.

**Units used in the model:**

The frame of reference selected for the model is summarized in Fig.1D. The distance expressed in terms of cell diameters was selected for two main reasons: first, it is an intuitive unit commonly used in biology, that helps to visualize the data; and second, for practical reasons, since the mathematical equation of the model and the software code implementation are simplified using this distance unit. Therefore, the distance estimated through the experimental data (initially in μm) was normalized to cell diameters dividing by the average cell size, as described in cell size measurement protocols. The temporal unit used in this model was the second, so, the time calculated in the model from the *in*
dynamic cytonemes were expressed in seconds. The rest of variables in the model are either dimensionless or expressed in terms of cell diameters and seconds.

**Requirements to compare experimental data of the gradient with the theoretical contact function.**

The conditions to compare *in silico* simulations with experimental data have been estimated in the supplemental material and can be summarized as:

- **Mathematically:** The degradation rate of the morphogen should be taken into account.
- **Experimentally:** Confocal images must have been taken according to the linear gamma function and within the limits of the acquisition range.

**In silico simulations**

- **Numerical simulations:**

  Each *in silico* prediction was computed 2000 times per simulation and case, the different values obtained over those 2000 times were used to generate the predicted gradient for those conditions. The standard deviation of the data obtained of those 2000 times was used as the expected signal variability.

- **Parameters and data for simulations:**

  Supplementary Table S.2 details the parameters used in each simulation. The updated experimental data in those simulations were obtained from the already described measurements of the wing disc cytoneme length and the average cell diameter ($\phi = 4.37 \pm 0.89$ $\mu$m for abdominal histoblasts and $\phi = 3.05 \pm 0.65$ $\mu$m for wing disc cells). The experimental cytoneme dynamics were obtained from previous studies (30). Finally, the
average Hh gradient profiles obtained in the validated simulations was used in both systems. The degradation rate of Hh was obtained from a previous study (66).

- **Statistical analysis.**

The Wilcoxon rank sum test was performed per pairs and the resulting p-values were graphically coded in matrixes with a green color that is graded depending on the significance. The code used was: black for no significance (= n.s) and dark green to light green respectively for the p-values: * = p-value < 0.05, ** = p-value < 0.01, *** = p-value < 0.001, **** = p-value < 0.0001.

As mentioned, *in silico* predictions were computed over 2000 times per simulation in each case; the different values obtained over those 2000 samples were used for the statistical analysis using Wilcoxon rank sum test (Implemented in Matlab2015a).

For the Coefficient of variation, we divided the 2000 simulations in 100 subgroups of 200 samples each. Then the coefficient of variation distribution per case was performed over those 100 subgroups. Finally, we used those 100 values per case for the posterior statistical analysis of the signal variability using the coefficient of variation.

- **Simulations of the diffusion-degradation model**

Simulations for the diffusion-degradation model (next equation) were performed using in Matlab2015a language using *pdepe* function for 1-D parabolic and elliptic PDEs (See Supplementary Figure S.10).

\[
\begin{align*}
\text{PDE: } & \quad \frac{\partial u}{\partial t} = D\Delta u - \delta \cdot u \\
\text{BC: } & \quad u(0, t) = \langle u^N_{\text{exp}} \rangle_{x=0}, \quad 0 < t < \infty \\
& \quad \frac{\partial u(L, t)}{\partial x} = 0, \quad 0 < t < \infty \\
\text{IC: } & \quad u(x, 0) = 0, \quad 0 < x \leq L
\end{align*}
\]
The boundary and initial conditions are required to solve the equation. We decided to use the condition of the morphogen flux equal to zero at the tissue end \( \frac{\partial u(L,t)}{\partial x} = 0 \). This derivative boundary condition is commonly used and in biological terms means that the morphogen cannot escape from the tissue. The initial conditions the morphogen gradient is zero \( u(x,0) = 0 \) since there is no previous diffusion of the morphogen. Finally, to have more precise experimental conditions for the diffusion simulations, the boundary condition at the origin has been selected as the experimental average of the maximum values for the normalized Hh gradient \( u(0,t) = \langle u_{exp}^N \rangle_{x=0} \).

The same experimental data were used for both: cytoneme and diffusion models. A table detailing the parameters can be found in supplementary material (Supplementary Table S.4).

**Cytomorph simulations for other morphogens**

*Decapentaplegic (Dpp) gradient*

The Dpp signaling pathway is functional in the same wing disc cells as Hh signaling. Therefore, we used the length distribution of cytonemes protruding from the A and P compartment cells quantified in this work using non-morphogen-specific markers (membrane markers) (Fig 3.D). Since some of the receiving cells are also producing Dpp in the A compartment of the wing disc, we only simulated the gradient in the Dpp receiving cells of the P compartment.

Although we have not focused on the temporal aspects of the Dpp gradient formation, for the proper simulation of the Dpp gradient we considered the experimental degradation rate of Dpp (53). It has been reported that the experimental half-time of the Dpp protein was shorter than that of the Hh protein \( \text{Dpp} = 45 \text{ min} \) (53) vs \( \text{Hh} = 166 \text{ min} \) (66), therefore, we simulated the Dpp gradient during a short period of development.
exact values of the parameters simulated are available in Supplementary Table S.2. To be able to compare our simulations with the experimental Dpp gradient, we used the WebPlotDigitizer tool to convert images in numerical data extracting the published Dpp experimental information from (51).

Wingless (Wg) gradient extension

The \textit{in silico} simulation of the exact shape of the Wg gradient in the wing disc would require to update Cytomorph with new modules because some of the receiving cells are also producing Wg. We calculated the expected length of the Wg gradient using the spatial conditions of Eq-3, in which the range of the gradient is limited by the sum of the lengths of cytonemes emanating from producing and receiving cells: \(x_p < \lambda_r(t)+\lambda_p(t) - x_r\). Therefore, the total length of the gradient is limited by maximum length of cytonemes \(\lambda_{\text{max},r}+\lambda_{\text{max},p}\) protruding from the border of producing/non-producing cell. Assuming again that the receiving and producing lengths of Wg cytonemes are the same in the wing disc as those of others morphogens (\(\lambda_{\text{mean},r} = 11.8 \, \mu m\) with a standard deviation of \(4.1 \, \mu m\) and \(\lambda_{\text{mean},p} = 14.5 \, \mu m\) a standard deviation of \(5.5 \, \mu m\)), the expected length of the Wg gradient will be \(\lambda_{\text{max},r}+\lambda_{\text{max},p} \approx 15.9 + 20 = 35.9 \, \mu m\) from the last Wg producing cell, which corresponds to \(\sim 42 \, \mu m\) from the D/V border.

References and Notes


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Figure 1. Schemes of cytoneme-mediated cell signaling based on experimental evidence: A) Type 1: Receiving cells emit cytonemes to collect the morphogen from producing cells. B) Type 2: Producing cells emit cytonemes to deliver the morphogen to receiving cells. C) Type 3: Both producing and receiving cells emit cytonemes to deliver and collect the morphogen respectively. D) Frame of reference used to develop the mathematical equations. E) Schematic representation of the cytoneme triangular dynamics. F) Schematic representation of the cytoneme trapezoidal dynamics.
### A. Experimental data

<table>
<thead>
<tr>
<th>Gradient distribution</th>
<th>Cytoneme length</th>
<th>Time elongation</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_{1,2} / y_{1,2}$</td>
<td>$L_1$</td>
<td>$t_{e1}$</td>
</tr>
<tr>
<td>...</td>
<td>...</td>
<td>...</td>
</tr>
<tr>
<td>$x_n / y_n$</td>
<td>$L_n$</td>
<td>$t_{e_n}$</td>
</tr>
</tbody>
</table>

### B. GUI parameters

- Cytomorph: Software to simulate morphogen gradients through cytonemes

#### Static parameters
- Number of receiving cells: 15
- Number of producing cells: 15
- Average cell size (microns): 3
- Number of cytonemes per cell: 4
- Probability of contacts: 0.1
- Contact lifetime (s): 60
- Degradation Rate (1/s): 0.00007

#### Simulation parameters
- Number of simulations per case: 2000
- Developmental time simulated (s): 3000
- Show experimental gradient

### C. Outputs

#### Variable role (Scan case) Comparison between cases

1. Simulation parameters
2. Contact distribution
3. Coefficient of variation
4. # of contacts per time iteration
5. Contact distribution (over time)
6. Normalized gradient
Figure 2. General outline of the Cytomorph workflow: A) Inputs of the Cytomorph, divided into two groups: A.1) Loading the data through an Excel table. A.2) Loading it through the main GUI. B) Cytomorph secondary GUI windows: B.1) Window in which different parameter combinations (cases) can be loaded to compare with the reference case. B.2) Window in which a scan of variable values can be selected to study their effect. B.3) Window in which graphical properties can be selected. C) Graphic outputs of Cytomorph simulations: C.1 and C.2) Contacts per cell along simulations. C.3) Signal variability measured by coefficient of variation. C.4) Contacts per cell and iteration. C.5) Temporal evolution of the contact distribution. C.6) Final gradient and expected variability (error bars).
Figure 3. Experimental cytoneme characterization in *Drosophila* tissues. A) Wing imaginal disc cytonemes protruding from A (top left) and from P (top right) compartment cells marked with Ihog-RFP. Bottom panels show 3D reconstructions of a confocal Z-stack taken at the basal side of the tissue showing cytonemes protruding and from A (bottom left) and P (bottom right) compartment cells. B) A confocal Z-stack taken from the apical to basal side of the abdominal histoblast epithelium with the A compartment marked with life-actin-RFP (red) and the P compartment marked with CD8GFP (green). C) *In vivo* temporal sequence of abdominal histoblast cytonemes taken at one-minute intervals. Top image sequences show both A and P compartment labelled cytonemes (A in red, P in green), middle image sequences show a single channel of A compartment cytonemes, and bottom image sequences show the single channel of P compartment cytonemes. D) Statistical violin plots of cytoneme length distribution in the A (blue) and the P (green) compartments in wing disc (left) and abdominal histoblast nest (right). Scale bars: 15μm
Figure 4. Experimental and simulated Hh gradients in Drosophila tissues. A) Confocal sections of Drosophila epithelia labeled with Hh:GFP BAC and EnhancerPtcRed. Top: abdominal histoblast nest. Bottom: imaginal wing disc. B) Quantified data of the Hh gradient in both epithelia: wing disc (green) and abdominal histoblast nest (blue). C) Comparison between the wing disc experimental gradient (green) and the predicted gradient estimated by our cytoneme model (blue). D) Comparison between the abdominal histoblast nest experimental gradient (green) and the predicted gradient estimated by cytoneme model (blue). E) Comparison between the wing disc experimental gradient (green) and the predicted gradients applying different models: cytoneme model (blue) and diffusion-degradation model (black). F) Comparison between the abdominal histoblast experimental gradient (green) and the predicted gradients applying different models: cytoneme model (blue) and diffusion-degradation model with different diffusion coefficient (red 3 times smaller than black). Scale bars: 30μm.
Figure 5. *In silico* study of different cytoneme variables and their predicted impact on gradient features. Reference simulation in red, simulations after modifying a specific parameter in blue (graded light to dark depends on the value) and experimental data in green. X) Left. Morphogen distribution for different cases, normalized to the maximum value of the reference case, along receiving cells including the expected variability per cell row (error bars). Right. Study of the number of contacts in the first row of receiving cells $x_0$, normalized to the average value of the reference case: top, violin plots of 2000 simulations per case; bottom, green-color-coded matrix of p-values for the violin distributions. X’) Coefficient of variation per case in the first row of receiving cells $x_0$ (left). Green-color-coded matrix of p-values for violin distributions (right). X’’) Distribution of contacts normalized to their maximum value to compare changes in gradient shape along receiving cells (left). Coefficient of the normalized distributions to study the scaling along receiving cells (right). A) Simulations for different cell size/cytoneme length ratios ($\phi=2.5$ to 3.5 each 0.2 μm (blue), $\phi=3$ μm (red)). B) Simulations for different number of producing cells rows involved in the signaling (Np=1 to 14 (blue), Np=15 (red)). C) Simulations for different number of cytonemes per cell (n$cyt$=1 to 3 (blue), n$cyt$=4 (red)).
Morphogen distribution

\[ N_i(x_r) = \max \left( N_{ref}(x_i) \right) \]

\[ N_{s,i}(x_0) = \frac{\text{mean}(N_{ref}(x_0))}{s} \]

Signal Variability

\[ C.V = \frac{\text{std}(N_{s,i}(x_0))}{\text{mean}(N_{s,i}(x_0))} \]

Scaling

\[ N'_i(x_r) = \frac{N_i(x_r)}{\max(N_i(x_r))} \]

\[ N'_i(x_r) / (N'_{ref}(x_i)) \]
Figure 6 *In silico* study of different cytoneme presumptions and their predicted impact on gradient features. Reference case in red, simulations after modifying a feature in blue and experimental data in green. X) Left. Morphogen distribution along receiving cells for different cases, normalized to the maximum value of the reference case, showing the expected variability per cell row (error bars). Right. Study of the number of contacts in the first row of receiving cells $x_0$, normalized to the average value of the reference case. Top, violin plots of 2000 simulations per case. Bottom, green-color-coded matrix of p-values for the violin distributions. X’) Coefficient of variation per case in the first row of receiving cells $x_0$ (left). Green-color-coded matrix of p-values for violin distributions (right). X”’) Distribution of contacts normalized to their maximum value to compare changes in gradient shape along receiving cells (left). Coefficient of the normalized distributions to study the scaling along receiving cells (right). A) Simulations for different cytoneme signaling type (type 3 in red and type 1-2 in blue). B) Simulations for different contact functions (type $\psi(\mu)$ in red and type $\psi(\mu, x)$ in blue). C) Simulations of the hypothetical case of multiple contacts between cytonemes along the overlapping surface (single contact in blue, multiple contacts in red). D) Simulations for different probability of contact ($\mu = 40\%$ to $80\%$ each $20\%$ in blue, $\mu = 100\%$ in red).
**Morphogen distribution**

<table>
<thead>
<tr>
<th>Receving cells</th>
<th>50% Triangles</th>
<th>10% Triangles</th>
<th>90% Triangles</th>
</tr>
</thead>
<tbody>
<tr>
<td>15 µm</td>
<td></td>
<td></td>
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</tbody>
</table>

**Signal Variability**

\[ C.V = \frac{\text{std} \left( N_{s,i} (x_0) \right)}{\text{mean} \left( N_{s,i} (x_0) \right)} \]

**Scaling**

\[ N'(x_r) = \frac{N_i (x_r)}{\text{max} \left( N_i (x_r) \right)} \]

\[ N'(x_r) = \frac{N_i (x_r)}{N'_\text{ref} (x_r)} \]

**In silico vs exp Hh recovery**

**Statistical p-values matrix**

<table>
<thead>
<tr>
<th>C-R</th>
<th>C-2</th>
<th>C-3</th>
</tr>
</thead>
<tbody>
<tr>
<td>n.s</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

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Figure 7. FRAP experiments to study the temporal gradient formation. A) Representative image of FRAP experiments in abdominal histoblast nests in which the signal is eliminated after photobleaching over a specific ROI. B) Hh (Hh:GFP BAC) gradient profile shortly before bleaching in black and Hh signal recovery over time coded in a hot colormap; each step corresponds to 45 seconds. C) Ptc (EnhancerPtcRed) expression profile shortly before bleaching in black and ptc signal recovery over time coded in a hot colormap, each step is 45 seconds. D) In silico signal evolution predicted for abdominal histoblast nests. E) A graphical comparison every 3 minutes between in silico simulations and experimental data. F) Simulations for cytonemes contacting while growing with a different proportion of triangular/trapezoidal cytoneme dynamics (10% triangles in light blue, 50% triangles in red, 90% triangles in dark blue). Scale bars: 15 μm.