Morphogen-based simulation model of ray growth and joint patterning during fin development and regeneration

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SUMMARY
The fact that some organisms are able to regenerate organs of the correct shape and size following amputation is particularly fascinating, but the mechanism by which this occurs remains poorly understood. The zebrafish (Danio rerio) caudal fin has emerged as a model system for the study of bone development and regeneration. The fin comprises 16 to 18 bony rays, each containing multiple joints along its proximodistal axis that give rise to segments. Experimental observations on fin ray growth, regeneration and joint formation have been described, but no unified theory has yet been put forward to explain how growth and joint patterns are controlled. We present a model for the control of fin ray growth during development and regeneration, integrated with a model for joint pattern formation, which is in agreement with published, as well as new, experimental data. We propose that fin ray growth and joint patterning are coordinated through the interaction of three morphogens. When the model is extended to incorporate multiple rays across the fin, it also accounts for how the caudal fin acquires its shape during development, and regains its correct size and shape following amputation.

KEY WORDS: Pattern formation, Zebrafish, Bone development

INTRODUCTION
Understanding how shape is controlled during development is one of the greatest challenges in developmental biology. Not only do organisms develop into particular shapes according to their species, but some species are able to regenerate organs of the correct size and shape following amputation (Yin and Poss, 2008). How this is achieved largely remains a mystery, but the control of growth, and therefore of shape, has been shown to involve morphogens (Bénazet and Zeller, 2009; Lee et al., 2005; Schwank and Basler, 2010). Morphogens are mobile substances that can elicit the expression of different sets of genes at different concentrations, hence generating spatial patterns of gene expression that can instruct tissue differentiation or alter tissue growth (Schwank and Basler, 2010). Therefore, spatial patterns of gene expression lead to the correct spatial organization of the many cell and tissue types of an organism. As morphogens can affect growth, changes in shape and growth may, in turn, affect morphogen concentration and distribution, thus altering patterning processes.

Simulation modeling provides a way to test hypothetical mechanisms of growth and patterning that would otherwise be difficult to comprehend owing to the highly dynamic nature of development in time and space (Baker et al., 2008; Kondo and Miura, 2010; Meinhardt, 2009). In this paper, we use simulation modeling to explore how growth and patterning might be coordinated through the action of morphogens during the development and regeneration of bony fin rays in the zebrafish (Danio rerio), an emerging model system for the study of bone morphogenesis (Akimenko et al., 2003; Iovine, 2007; Poss et al., 2003).

The zebrafish, like most teleost fish studied to date (Wagner and Misof, 1992), grows throughout its life and can regenerate many tissues, including its fin rays (Poss et al., 2003). The zebrafish caudal fin comprises 16 to 18 long dermal bony rays, each of which is composed of two hemirays that face each other (Fig. 1). Bone growth along the proximodistal axis of each ray occurs distally through discrete pulses of cell proliferation (Jain et al., 2007), and growth slows down as the fish ages (Iovine and Johnson, 2000). The caudal fin is bi-lobed (Fig. 1), indicating the lobe rays must grow faster than the rays between the lobes (cleft rays), at least during part of fin development. As a fin develops, each fin ray extends distally through the addition of bone matrix, and distal joint formation gives rise to segments (Haas, 1962) (Fig. 1). Mechanisms underlying fin growth and segment formation have been studied using morphological and genetic analyses (Iovine, 2007; Mari-Befa and Murciano, 2010).

Upon shortening of the fin through partial amputation, a distal blastema forms, which then allows regeneration to proceed (Akimenko et al., 2003). After amputation, lobe rays regenerate faster than cleft rays (Lee et al., 2005), allowing the fin to regain its pre-amputation size and shape, and then to carry on growing normally. Molecular mechanisms at play during development and regeneration are thought to be similar (Iovine, 2007), although growth during regeneration is faster than ontogenetic growth (Iovine, 2007). The level of amputation has an effect on regeneration rates, as regeneration initially proceeds faster if the fin is cut in proximal positions than if it is cut in more distal positions (Lee et al., 2005). However, a fin cut in a proximal region and a fin cut in a distal region attain the pre-cut fin size in similar amounts of time (Iovine, 2007). At the molecular level, several factors, such as fibroblast growth factors (Fgfs), sonic hedgehog a (Shha) and the bone morphogenetic protein 2b (Bmp2b) have been implicated in distal ray growth (Lee et al., 2005; Quint et al., 2002). In particular, Fgfs have been shown to control rates of cell proliferation in the fin, during both ontogeny (Wills et al., 2008) and regeneration (Lee et al., 2005), and domains of active Fgf signaling and cell proliferation are larger after proximal...
Based on the above qualitative information on growth and joint patterns in caudal fins, we propose a simple plausible model for the control of fin ray growth and regeneration, as well as for joint patterning. The model involves three morphogens and accounts for qualitative data from the literature regarding growth rates during both ontogeny and regeneration, and variations in growth and segment length within and between rays. The model extended to multiple rays also accounts for the regeneration of a properly shaped fin following amputation.

**MATERIALS AND METHODS**

**Modeling software and statistical analysis**

Programs for the ray models, as well as programs for model output analysis, were developed using Matlab version R2010b for Windows (The Mathworks, Natick, MA), and are available at http://mysite.science.uottawa.ca/arolland/. Analysis of variance was carried out using SPSS Statistics 19 (IBM corporation, Somers, NY).

**Model assumptions and simplifications**

Mechanisms of bony fin ray development involve ray growth and regeneration, joint formation and ray bifurcation. Rays can grow and form segments even in the absence of neighboring tissue (Mari-Beffa et al., 1999), whereas grafting experiments have shown that inter-ray tissue is important in the control of ray bifurcation (Murciano et al., 2002). Here, we only consider the mechanisms of ray growth in length and joint pattern formation; hence, we do not need to consider inter-ray tissue. In addition, although rays are composed of two facing hemirays, single hemirays can grow and form segments normally in absence of their facing hemiray (Murciano et al., 2007). Therefore, we decided to consider rays as a whole without making a distinction between the two hemirays. In the present model, we only focus on mechanisms controlling growth along the proximodistal axis of the rays, and ignore growth along the dorsoventral axis of the fin (Fig. 1). Bifurcation formation and growth along the dorsoventral axis of the fin will be examined in separate studies.

In the model implementation, a fin ray is discretized into compartments. Each compartment is treated as if it were a cell, and given an arbitrary length \( l \) equal to one (unit length). Compartments are numbered from one at the base of the ray to \( n \) at the distal end. However, owing to the large number of cells involved in whole rays, each compartment in reality represents a group of cells.

The control of growth in fins has been shown to involve gradients (Lee et al., 2005). In molecular terms, such gradients could arise from signal transport from cell to cell, and/or from a cell lineage-based mechanism (Warlick et al., 2009). The fact that the sof mutant, which is defective in the gap-junction protein connexin 43 (Iovine et al., 2005), has shorter fins and shorter ray segments than the wild type (Iovine and Johnson, 2000) indicates that signal transport is important in the control of fin ray growth and joint patterning. The most parsimonious model to simulate transport involves the use of diffusion equations. As pointed out by Kondo and Miura (Kondo and Miura, 2010), even if in complex systems the specific mode of transport at the molecular level might not involve free diffusion, diffusion equations often correctly recapitulate the behavior of the system, which justifies the use of such equations as a simple way to model complex patterning phenomena. For more information on simulation models of pattern formation and growth involving diffusion equations, we refer the reader to the classical models of sea shell patterns (Meinhardt, 2009).

Our model does not explicitly consider the role of the nervous system or the vasculature; however, it has been shown that the nervous system is necessary for a blastema to form following fin amputation in Fundulus (Géraudie and Singer, 1985). In the model, this is indirectly taken into account by assuming that a blastema automatically forms following amputation. Regarding the vasculature, recent data show that the fin ray pattern formation precedes vascularization in the fin (Huang et al., 2009), indicating that vascularization is not required for initial patterning of the rays. Altered vasculature can affect ray spacing and fin growth (Huang et al., 2009), but as the current model does not address ray spacing mechanisms, it does not involve the vasculature.
Model 1: Model of ray growth during fin development and regeneration

Based on the literature, a model of fin ray growth has to account for the following experimental evidence: ray growth along the proximodistal axis of the fin occurs distally (Jain et al., 2007); fins display indeterminate growth, which slows over time (Iovine and Johnson, 2000); upon amputation, regeneration growth occurs and is faster than ontogenetic growth until the fin reaches its pre-cut size (Iovine, 2007); regeneration proceeds faster if the fin is amputated in proximal position (Lee et al., 2005); a fin cut in a proximal region and a fin cut in a more distal region will attain the pre-cut size in a similar amount of time (Iovine, 2007).

Substances affecting growth

As ray growth along the proximodistal axis of the fin occurs distally (Jain et al., 2007), we postulate the presence of a substance, arbitrarily named \( G \), which acts distally to promote growth during both development and regeneration. We assume an inherent particularity about the distal compartment of a ray, so that \( G \) can only be produced in that compartment.

We model \( G \) transport using diffusion equations. \( G \) has a low diffusion constant, and/or high decay rate, which limits growth to the most distal part of the fin.

As growth was experimentally shown to slow down over time (Iovine and Johnson, 2000), it follows that \( G \) levels at the distal end of the ray must decrease as a ray develops and enlarges. At the same time, based on experimental data on regeneration from proximal and distal amputations (Lee et al., 2005), \( G \) levels in the distal part of a ray after amputation must be higher if the amputation is in a proximal part of the fin than if the amputation is more distal. In order to account for such variations in \( G \), another substance is required in the model, which is arbitrarily named \( X \), and is present in decreasing concentrations from ray base to ray tip. \( X \) activates \( G \) production in the distal compartment of the ray. In order to account for the fact that regeneration proceeds faster than ontogeny (Haas, 1962; Iovine, 2007), \( X \) levels have to increase with time (as the fish increases in size and age).

The simplest way to fit the above requirements for \( X \) is to assume that \( X \) is produced in the proximal compartment (at the base of the fin), at a rate that increases over time, and that \( X \) is transported between compartments. Using diffusion equations and setting a high diffusion constant for \( X \) and/or a low decay rate then allows it to reach the distal compartment and activate \( G \) production.

Note that the range of action of morphogens depends not only on the diffusion constant, but also on the decay rates. Hence, we can manipulate concentration profiles of \( G \) and \( X \) by altering those two parameters (e.g. \( X \) and \( G \) could both be transported at the same rate but \( G \) might have a much higher decay rate).

The model operates along one dimension. In the model, each compartment has two neighbors, except for the first and the last compartment. The fluxes of \( G \) and \( X \) coming into compartment \( i \) from compartment \( j \) per unit time are described using Fick’s law:

\[
\phi_{ij} = D_X \frac{G_j - G_i}{l}, \\
\phi_{ji} = D_X \frac{X_j - X_i}{l},
\]

where \( \phi_{ij} \) represents the flux of \( G \) from compartment \( j \) to compartment \( i \), \( D_X \) is the diffusion constant of \( G \), and \( G_i \) and \( G_j \) are concentrations of \( G \) in compartments \( j \) and \( i \), respectively. Similarly, \( \phi_{ji} \) represents the flux of \( X \) from compartment \( j \) to compartment \( i \), \( D_X \) is the diffusion constant of \( X \), and \( X_i \) and \( X_j \) are concentrations of \( X \) in compartments \( j \) and \( i \), respectively. \( l \) is the length between the centers of compartments \( j \) and \( i \), and \( l \) is constant.

The equations describing the variations in the amounts of \( G \) and \( X \) in a given compartment \( i \) during time interval \( dt \) are as follows:

\[
\frac{dG_i}{dt} = -\delta_G G_i + \sum_{j=1}^{i} \phi_{ij} \quad \text{if } i < n, \tag{3a}
\]

\[
\frac{dG_i}{dt} = \alpha_G X_i - \delta_G G_i + \phi_{ji} \quad \text{if } i = n, \tag{3b}
\]

with

\[
\frac{d\alpha_G}{dt} = \beta \quad \text{if } \alpha_G < \alpha_{G_{\text{max}}}, \tag{4b'}
\]

\[
\frac{dX_i}{dt} = 0 \quad \text{if } \alpha_G = \alpha_{G_{\text{max}}}, \tag{4b''}
\]

and

\[
\alpha_{G_{(i+n)}} = \alpha_{G_{\text{max}}}, \tag{4b'''}
\]

where \( G_i \) and \( X_i \) are, respectively, the concentrations of \( G \) and \( X \) in compartment \( i \), \( \alpha_G \) and \( \alpha_X \) are production coefficients for \( G \) and \( X \), \( \delta_G \) and \( \delta_X \) are rates of decay for \( G \) and \( X \). \( \phi_{ij} \) and \( \phi_{ji} \) represent the net fluxes of \( G \) and \( X \), respectively, from neighbor compartment \( j \) to compartment \( i \) (see Eqs 1 and 2; note that the net flux can be negative, representing outflow from \( i \) to \( j \)). \( n \) is the last compartment in the model, and \( k \) is the number of neighbors for compartment \( i \). The production rate of \( G \) in compartment \( n \), \( \alpha_X X_n \), depends on the amount of \( X \) present. The rate of \( X \) in compartment 1, \( \alpha_X \), increases linearly with time at rate \( \beta \) (the initial production rate is \( \alpha_X(0) \)), until reaching a saturation value \( \alpha_{X_{\text{max}}} \) (in practice, our chosen value for \( \alpha_{X_{\text{max}}} \) is not reached over the course of the simulations).

Implementation of growth

Growth is implemented by considering that each compartment can divide to give rise to two ‘daughter compartments’ of the same size as the mother compartment. In each compartment, \( G \) promotes growth/division by increasing the level of a division substance named \( C \). The initial value for \( C \) in each compartment is \( C_0 \). The level of \( C \) in a compartment \( i \) is subsequently given by:

\[
\frac{dC_i}{dt} = \alpha_C C_0 G_i, \tag{5}
\]

where \( C_i \) is the level of division in compartment \( i \), and \( G_i \) is the concentration of \( G \) in that compartment. \( \alpha_C \) represents how close a compartment is to dividing; therefore, it indirectly represents compartment growth. \( \alpha_C G_i \) is the rate of production of \( C_i \), which depends on both \( C_i \) and \( G_i \). \( \alpha_C \) is a constant. Unlike \( G \) and \( X \), \( C \) is not a transported substance. For the purpose of this model, \( C \) growth is considered irreversible, therefore \( C \) cannot decrease.

Once \( C \) doubles its initial level (i.e. reaches \( 2C_0 \)), the compartment divides into two compartments. A compartment that is about to divide (mother compartment) will give rise to two daughter compartments, each of the same size as the original mother compartment. This is equivalent to the mother compartment doubling in volume before dividing into two. Therefore, concentrations of \( X \), \( G \) (and \( S \) in model 2 below), and levels of \( C \) in each daughter compartment are divided by two compared with concentrations of the mother compartment. The formation and maintenance of the gradients in \( X \) and \( G \) (and \( S \) in model 2) is therefore controlled by transport between compartments, as well as by growth and compartment division.

Amputation

Amputation is implemented by simply removing compartments beyond a user-defined number of compartments at a user-chosen time during the simulation. We then assume that the new distal-most compartment has the specific property that \( G \) can be produced there. So, we effectively assume the formation of a blastema, as is observed experimentally (Akimenko et al., 2003).

Model 2: Model of ray joint pattern formation during fin development and regeneration

The model of joint pattern formation is built upon the ray growth model. Based on the literature, a model of ray joint formation has to account for the fact that bone segments are added distally (Haas, 1962), and that
segments tend to be smaller towards the distal end of the fin (Haas, 1962; Iovine and Johnson, 2000; Sims et al., 2009). This indicates that new segments formed along a ray are progressively smaller (Jain et al., 2007) as the fin develops.

The phenotypes of lof and evx1 mutants suggest that joint patterning and fin growth may be regulated independently (Iovine and Johnson, 2000; Schulte et al., 2011). Therefore, it is reasonable to assume the presence of a morphogen specifically involved in joint formation. On the other hand, joints in the nof mutant are closer together and fins are shorter than in the wild type, showing that growth and joint formation are closely linked processes (Sims et al., 2009). Hence, the mechanism of joint formation can be linked to the mechanism controlling growth. We tested several hypothetical mechanisms for joint patterning that involve interactions between X, G and a morphogen that either promotes or inhibits joint formation. Only the most parsimonious model is presented here (examples of outputs for alternative models are briefly presented in supplementary material Fig. S1).

We postulate the existence of a joint-inhibiting substance S. Segment joints are formed in compartments where S and G are each below a set threshold. In each compartment i, at each time step of the simulation, if Si < \sigma and Gi < \gamma, a joint is formed. For simplicity, we keep the position of the joint within the compartment abstract and represent it to be in the center of the compartment in which it formed.

Joint formation is irreversible (once a joint has formed, it cannot disappear). The presence of a joint triggers production, in response to the presence of X. This inhibits joint formation in the vicinity of the newly formed joint (because S concentrations will be above the threshold \sigma).

The equations describing the variations in the amounts of S in a given compartment i during time interval dt are as follows:

\[
\frac{dS_i}{dt} = \alpha_X X_i - \delta_X S_i + \sum_{j=1}^{k} \phi_{i,j} \quad \text{if } i \text{ contains a joint},
\]

\[
\frac{dS_i}{dt} = -\delta_X S_i + \sum_{j=1}^{k} \phi_{i,j} \quad \text{if } i \text{ does not contain a joint},
\]

with

\[
\phi_{i,j} = \frac{D_X S_j - S_i}{l^2}.
\]

where S_i and X_i are the concentrations of S and X in compartment i. \alpha_X is a production coefficient for S, \delta_X is the rate of decay for S. \phi_{i,j} represents the flux of S from compartments j to compartment i. D_X is the diffusion constant of S. l is the length between the centers of compartments j and i. k is the number of neighbors of compartment i.

**Experimental procedures for model validation**

Two groups (A and B) of 10 wild-type zebrafish aged 167 days and of fin length S to 7 mm were amputated at 2 mm (proximal cut, group A) or 4 mm (distal cut, group B) from the fin base and imaged twice a week for 80 days. Fish were kept in water tanks maintained by a linked system with water temperature kept at 28.5°C. During each imaging session, fish were processed one at a time. On the day of amputation (day 0), each fish was processed by taking it out of its water tank, anesthetizing it in a 91.8585 μmol tricaine methanosulfate (TMS) solution, imaging its caudal fin before amputation, amputating the fin, imaging the fin again following amputation and finally returning the fish to its water tank where it woke up. Images

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**Table 1. Default parameter values and initial conditions**

<table>
<thead>
<tr>
<th>Parameters related to X</th>
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<tr>
<td>\delta_G</td>
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<table>
<thead>
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<td>\gamma</td>
<td>0.1</td>
</tr>
<tr>
<td>\sigma</td>
<td>0.1</td>
</tr>
</tbody>
</table>

| Time step | 0.01 |

Initial conditions: 1 compartment, with X=0, G=0, S=0

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were taken using a Canon powershot A640 digital camera mounted on a Zeiss V8 Discovery microscope. After day 0, fish were imaged twice a week for 80 days following the same anesthetizing and imaging procedure as on day 0. Nine fish survived the experiment in group A, and eight fish survived in group B.

Digital images were used to measure fin length above the point of amputation, before amputation and over the course of regeneration, as well as to compare the length of segments cut during amputation before they were amputated and after they regenerated. Measures were taken using custom software written in Matlab (A.-G.R.-L., V. Lefebvre and M.-A.A., unpublished). For each fish, fin length was measured along the third long ray from the ventral side of the fin. For measurements to be accurate, length was measured from a ray joint below the amputation point up until the edge of the fin. The length measured from the image taken just after amputation (length from joint to amputation level) was then subtracted from all length measurements at other time points (including before amputation) to give the length above amputation level. For fish from group A, we also used the image taken just after amputation to identify cut segments, which allowed us to measure cut segment length before amputation and after regeneration using pictures taken before amputation and at day 14 post amputation (all cut segments had regenerated by then).

RESULTS
Single ray model of development and regeneration
We simulated fin ray growth and regeneration through the interaction between three morphogens $X$, $G$ and $S$. Only $X$ and $G$ are involved in the control of ray growth, but all three morphogens are involved directly or indirectly in joint patterning. $X$ is produced at the base of the fin and its production increases over time. As the most parsimonious model involves diffusion equations, $X$ has a long transport range, allowing it to reach the distal compartment (distal tip) of the ray (Fig. 2), where it activates $G$. $G$ is responsible for cell proliferation, can only be produced at the distal end of the ray in response to $X$, and has a short transport range. This limits ray outgrowth to the distal part of the fin, as observed experimentally.
(Jain et al., 2007), and $G$ concentrations decrease from the distal end of the ray (Fig. 2). When $G$ concentration in a compartment is low enough, a joint is formed in that compartment, which triggers the production of joint-inhibiting substance $S$. $S$ production at a joint ensures that no further joints are formed in the immediate vicinity of that joint. Hence, $G$ and $S$ together control joint spacing. $S$ is produced at joint locations in response to $X$. As $X$ concentrations decrease from a ray base to its tip, the further a joint is from the ray base, the less $S$ will be produced at that joint, and the closer the next joint will be (Fig. 2). Hence, $X$ indirectly controls joint spacing, and the interactions between $G$, $S$ and $X$ create a pattern of $S$ expression with quasi-periodic peaks of decreasing intensity (Fig. 2). Default parameters for the model as well as initial conditions are given in Table 1.

In order to compare the dynamics of ray growth and segment formation during development and regeneration, we simulated ray development for 50,000 steps (Fig. 3A-D), and at step 50,001 we performed an amputation, leaving either 15 ray compartments intact (proximal amputation, Fig. 3E) or 30 ray compartments intact (distal amputation, Fig. 3F) out of 54 compartments present at step 50,000. In each case, the ray was left to regenerate until it reached the pre-amputation length (Fig. 3F,H).

We found that as growth proceeds, $X$ concentration in the distal compartment decreases (Fig. 4B, before amputation), and therefore $G$ concentration in the distal tip also decreases (Fig. 4B). This causes growth to slow down over time (Fig. 4C), as observed experimentally (Iovine and Johnson, 2000). As $X$ concentrations decrease from the proximal to the distal end of the ray (Fig. 2), immediately after amputation time the $X$ concentration in the new distal compartment is higher than the $X$ concentration in the distal compartment before amputation (Fig. 4A). Moreover, as $X$ production increases over time, the distal compartment at a given ray length will be exposed to higher $X$ concentrations during regeneration than during development (supplementary material Fig. S2). High levels of $X$ in the new distal compartment following amputation give rise to high levels of $G$ (Fig. 4B). Therefore, regeneration growth is faster than ontogenetic growth, which is in agreement with data from the literature (Iovine, 2007).

As levels of $X$ are highest at the ray base and decrease along the proximodistal axis (Fig. 2B), the blastema of a ray cut in a proximal position is exposed to higher $X$ levels than a blastema in a more distal position (Fig. 4A). Hence, a ray cut proximally initially produces more $G$ (Fig. 4B) than a ray cut distally, and regenerates faster (Fig. 4C,D), as shown by the higher slope of regenerate length as a function of time (Fig. 4D). These results obtained from the model fit with experimental data (Lee et al., 2005). As reported in the literature, rays cut at different levels regain the pre-cut length in similar (but not equal) amounts of time (Fig. 4C,D) (Iovine, 2007), as rays cut proximally re-grow faster but have a larger length to regenerate than rays cut distally. Experimental data on fins amputated in proximal and distal positions (supplementary material Fig. S3) confirm that fins amputated proximally regain their pre-amputation length in a time similar to (but slightly longer than) fins amputated distally, as in the model (Fig. 4C,D). However, we noted that regeneration times varied between samples of each group, and that a few fish from each amputation type did regenerate but did not regain their pre-amputation length over the 80 day period of the experiment.

Segment lengths in intact and regenerated fins decrease from the base of the ray to its tip (Fig. 4E), as seen in real fins (Iovine and Johnson, 2000; Sims et al., 2009), and segment numbers increase over time (Fig. 4F). An analysis of variance of the mean segment lengths in non amputated fins, regenerated fins cut proximally and regenerated fins cut distally showed no statistical differences between the three (df=2, F=0.519, $P=0.598$, Fig. 4G). Interestingly, new joints at the onset of regeneration are initially further apart than prior to amputation. In particular, segments cut during amputation regenerate to be longer than their original length (Fig. 3D,F,H). To test whether this also happens in real fins, we collected data on the length of segments cut by amputation before amputation and after regeneration (supplementary material Fig. S3). A Wilcoxon signed rank test confirms that in real fins average cut segment length per fish is shorter before amputation than after regeneration ($n=9$, $T=45$, $P$ (one tailed)=0.004).

**Effect of parameter variations**

In order to explore the effect of parameter variations on the model, $X$, $G$ and $S$ production and transport were altered, effectively generating virtual mutants. Parameters used for each virtual mutant are shown in Table 2. Decreasing $X$ production (virtual mutant $mxp1$) leads to a shorter ray with fewer and shorter segments, which is the same effect as the $sof$ mutation (Sims et al., 2009);

<table>
<thead>
<tr>
<th>Table 2. Parameters used for each virtual mutant compared to the default parameters</th>
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<tbody>
<tr>
<td>Mutants</td>
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<tr>
<td></td>
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<tr>
<td>$mst1$</td>
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<tr>
<td>$mst2$</td>
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</table>

Naming of each virtual mutant reflects which morphogen is altered ($X$, $G$ or $S$), whether production ($p$) or transport ($t$) is altered, and whether the relevant parameter is decreased (1) or increased (2).
conversely, increasing $X$ production ($mXp2$) leads to a longer ray with more numerous and longer segments (Fig. 5, supplementary material Fig. S4). Altering $X$ transport ($mxt1$ and $mxt2$) has similar effects on fin length and segment numbers, but average segment lengths are similar to those of the default model. In particular, increased $X$ transport ($mxt2$) leads to a longer fin with increased segment numbers of normal length, an effect reported for the lof mutation (Iovine and Johnson, 2000). Decreasing $G$ production ($mgp1$), which is equivalent to decreasing the sensitivity of $G$ to $X$, leads to a shorter ray with fewer segments, but segments are on average longer than with the default parameters. Conversely, increasing $G$ production ($mgp2$) produces a longer ray with more numerous and shorter segments (Fig. 5, supplementary material Fig. S4). Increasing or decreasing $G$ transport ($mgt1$ and $mgt2$) has minor effects for the range of parameter values explored. As $S$ is not involved in ray growth, varying $S$ production ($mspj1$ and $mspj2$) has no effect on fin length (Fig. 5A, supplementary material Fig. S4). However, decreasing $S$ production ($mspj1$) or transport ($mxt1$) produces a ray with more and shorter segments (i.e. joints are closer to each other) (Fig. 5B,C, supplementary material Fig. S4). As parameter changes for $mspj$ and $mst$ had similar effects to the $sof$ and lof mutations, but the quantitative effects on the phenotypes were small, we also altered parameters more drastically, and confirmed that we could generate strong phenotypes as seen in the $sof$ and lof mutants (Fig. 5B, supplementary material Fig. S5). Detailed quantitative analyses of mutant phenotypes will be needed to calibrate and test the model further.

**Whole-fin model**

Lower levels of $X$ production lead to shorter rays and segments, and fewer segments (Fig. 5), which matches the $sof$ phenotype (Sims et al., 2009), as well as the differences in ray length, segment numbers (Goldsmith et al., 2006) and segment lengths (Jain et al., 2007) between lobe rays and cleft rays. Therefore, we can easily extend the model to represent multiple rays of the fin by postulating a graded bimodal distribution of $X$ production rates at the base of the fin along the dorsoventral axis. The resulting fin (Fig. 6A-D) naturally displays a bi-lobed shape, and smaller and fewer segments in cleft rays compared with lobe rays, which is consistent with the literature (Goldsmith et al., 2006; Jain et al., 2007; Sims et al., 2009). After amputation (Fig. 6E), the fin automatically regains its bi-lobed shape and similar joint patterns to the pre-amputated fin (Fig. 6F-I). The regenerated fin (Fig. 6I) is therefore similar to the pre-amputation fin (Fig. 6D), although new joints at the onset of regeneration are initially slightly further apart than prior to amputation, as observed in the single ray model (Fig. 3D,F,H) and experimentally (supplementary material Fig. S3).

As $G$ promotes the production of a division factor that accumulates until reaching twice its initial value, growth in the model occurs in pulses. Those pulses are less frequent when $G$ levels are lower, hence pulses are less frequent in cleft rays compared with lobe rays (supplementary material Movie 1), and less frequent over time as rays increase in length (supplementary material Movie 1). Those results are in agreement with experimental observations (Goldsmith et al., 2003; Jain et al., 2007).

**DISCUSSION**

The regulation of growth and shape through the action of morphogenetic gradients is poorly understood (Schwank and Basler, 2010). We propose an integrated simulation model of fin ray growth and joint patterning during development and regeneration, which can account not only for the regulation of fin ray length and joint patterns, but also for the regulation of fin shape. The model postulates and therefore predicts the existence of a proximally derived morphogen $X$ that is transported and activates the distal production of a morphogen $G$ involved in cell proliferation. $X$ production increases over time, which accounts for faster growth during regeneration than during ontogeny. A third morphogen $S$ is involved in joint patterning through inhibition of joint formation. Low levels of $G$ and $S$ trigger joint formation, and joint formation activates the production of $S$. $S$ production at a joint prevents the formation of other joints in the immediate vicinity. Hence, $S$ directly affects joints spacing without affecting growth, and $G$ directly affects growth and the distance from the fin tip at which joints can form, depending on $X$. $X$ affects joints spacing indirectly as high $X$ levels at a joint will lead to higher $S$ production at that joint, hence a larger inhibition zone and larger space between joints. Strikingly, in the whole-fin model, the variation in
joint spacing between rays qualitatively matches experimental data (Goldsmith et al., 2006; Jain et al., 2007; Sims et al., 2009), and the fin naturally regains its bi-lobed shape and proper joint patterns following amputation. Two predictions of the model were validated experimentally: first, fins amputated distally and proximally do regenerate in similar but not equal amounts of time (data and model both indicated that fins amputated in distal positions regained their pre-cut length first); second, the first segments formed after amputation are longer than the segments formed at the same position during development.

A possible candidate for morphogen $G$ would be a fibroblast growth factor (Fgf). Fgf is a distal morphogen that affects fin growth, and Fgf signaling upon amputation closely mirrors $G$ signaling in the model (Lee et al., 2005). Loss of Fgf signaling in fins has been shown to cause distal tissue loss, indicating a role for Fgf in controlling fin homeostasis (Wills et al., 2008). It will therefore be interesting in the future to extend the model to incorporate a role for $G$ in compartment homeostasis. It is unclear which molecule $X$ might be, but Fgfs have been proposed to respond to some positional information graded along the proximodistal axis of the fin (Lee et al., 2005), which is consistent with the existence of a proximally derived morphogen that enhances Fgf activity. In the absence of experimental data, a model based on diffusion equations is the most parsimonious to model signal propagation, even if the molecular mechanism itself is not of a diffusive nature (Kondo and Miura, 2010). Indeed, morphogen gradients established by diffusion so far have been reported to operate at a short distance compared with the length of fin rays (Lander et al., 2009; Wartlick et al., 2009). $X$ production rate in the model increases over time; therefore, it increases with the age or size of the fish. Such an increase in morphogen production was recently shown in Drosophila, as the production rate of the morphogen bicoid scales with the embryo volume (Cheung et al., 2011). Finally, a bi-modal distribution of

![Fig. 6. Whole-fin model of ray growth and joint patterning during development and regeneration. Model is identical to the single ray model, except that $X$ production at the base of each ray follows a bimodal distribution along the dorsoventral axis of the fin. From the most dorsal ray to the most ventral ray, $\alpha_0 = 5, 6, 7, 6, 5, 4, 3, 2, 1, 1, 2, 3, 4, 5, 6, 7, 6, 5$, respectively, and for each ray $\beta = 0.001 \alpha_0$. (A-D) Development, (E) amputation, (F-I) regeneration. Fin in I has reached the pre-amputation length (the distal compartment of the two longest rays has just divided). See also supplementary material Movie 1.](image-url)
a signaling substance across fin rays is plausible, as there has to be a bi-modal distribution of ray growth rates along the dorsoventral axis of the fin, during at least part of development, for the fin to be bi-lobed. There is currently no direct experimental evidence for a morphogen corresponding to S. However, defects in connexin 43 (a gap-junction protein) yield joints that are closer to each other (Sims et al., 2009). This supports the involvement of a transported substance in joint spacing, and the idea that such a substance would have an inhibitory effect on joint formation.

Segmentation patterning to date has been mostly studied in the context of somite formation in the mouse embryo. Somites arise from the anterior to the posterior end of the vertebrate embryo at constant spacing during development, and later give rise to vertebrae and associated muscles (Dubrulle and Pourquié, 2004). As in somites, joints in the fin rays form in a sequential manner, with joints being added distally as the fin grows. However, contrary to somites, the ray joints form increasingly closer to each other. Somite pattern formation models involve a gradient of FgR8 with higher levels at the posterior end of the embryo, and a somite is thought to form when Fgf signaling reaches a sufficiently low level (Baker et al., 2008), which has been termed ‘determination front’ (Dubrulle et al., 2001). This is similar to the gradient of G and its involvement in joint formation. However, models of somite formation also involve an oscillatory clock to determine the timing of somite formation (Murray et al., 2011). In our model, interaction between G and S at the distal end of the ray repeatedly induces joint formation and expression of S at new joint positions. The time intervals between the apparition of joints and the spacing between joints are not constant, owing to the spatiotemporal dynamics of X concentrations. Hence, the relationship between time and space is highly integrated in the model. It will be interesting in the future to investigate potential parallels between somite formation models and ray joint-patterning models.

It has recently been proposed that interactions between a distal signal, involving Fgf, and a proximal signal pattern bone structures in the mouse limb bud (Mariani et al., 2008; Tabin and Wolpert, 2007) and the chick limb bud (Cooper et al., 2011; Rosello-Diez et al., 2011). In the case of the chick limb bud, the proximal signal was shown to be diffusible. Interestingly, as in the fin ray, the length of the skeletal elements decreases from the base to the tip of the tetrapod limb. It is therefore possible that a related model may underlie the control of endochondral bone length in tetrapods and dermal bone length in teleosts.

Advances in the molecular genetics of fin development and regeneration point to the involvement of multiple genes and signaling molecules, including not only Fgfs but also Wnts and activins (Yin and Poss, 2008). Here, we propose a simple mechanism whereby interactions between three morphogens lead to the conversion of spatial gradients into the formation of repeated segments of varying lengths as the fin grows in pulses of cell proliferation. The mechanism is very robust as fins regain their proper shapes and joint patterns following amputation. The model may be extended in the future to include further mechanisms such as ray bifurcation or the control of fin homeostasis (Wills et al., 2008). Modeling can be a useful tool to explore how postulated mechanisms give rise to complex developmental patterns. The parameter values of the model are currently arbitrary, owing to a lack of published quantitative data. Obtaining detailed quantitative data on fin ray growth and joint patterning during development and regeneration in wild-type and mutant fins will make it possible to test the model further, refine it and calibrate it. This will provide a link between molecular and computational analyses, opening the way for integrating simulation modeling and molecular data in future studies.

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Competing interests statement
The authors declare no competing financial interests.

Author contributions
A.-G.R.-L and M.-A.A. designed the research, A.-G.R.-L. wrote the simulation model, analyzed experimental data and wrote the paper, and M.P. and V.T. collected experimental data.

Supplementary material
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References


Model of fin ray morphogenesis


Fish sample number

Segment length before amputation (in mm)

Segment length after regeneration (in mm)

Amputation level: Distal
Proximal

Ray length above cut level (in mm)

Days after amputation