

# Waves and patterning in developmental biology: vertebrate segmentation and feather bud formation as case studies

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**ABSTRACT** In this article we will discuss the integration of developmental patterning mechanisms with waves of competency that control the ability of a homogeneous field of cells to react to pattern forming cues and generate spatially heterogeneous patterns. We base our discussion around two well known patterning events that take place in the early embryo: somitogenesis and feather bud formation. We outline mathematical models to describe each patterning mechanism, present the results of numerical simulations and discuss the validity of each model in relation to our example patterning processes.

**KEY WORDS:** *developmental waves, patterning mechanisms, mathematical modelling, somitogenesis*

## Introduction

Complex patterns arise as organisms grow, develop and interact with their environment. Be it a spatially heterogeneous distribution of cell types, density or gene expression, there are a plethora of examples of pattern formation in the early embryo, from *Drosophila* segmentation (Gilbert, 2006; Sanson, 2001) to digit formation and limb chondrogenesis (Tickle, 2000; Tickle, 2006), subdivision of the vertebrate antero-posterior (AP) axis into somites (Gossler and Hrabé de Angelis, 1998; Pourquié, 2001b) and formation of the avian integument (Yu *et al.*, 2002; Yu *et al.*, 2004). In spite of the dramatic advances made in genetic and molecular biology, we are still looking for answers to fundamental questions: in particular, what are the underlying mechanisms that give rise to patterning in development?

Developmental biologists have shown that graded distributions of morphogens play important roles in pattern specification and generation (Green and Smith, 1991; Murray *et al.*, 1988; Tabata and Takei, 2004; Wolpert *et al.*, 2006). However, there are still many unanswered questions as to: how these patterns are specified; how morphological changes take place; how patterns are regulated; and how the underlying mechanisms display robustness to intrinsic and environmental perturbations.

Wolpert proposed a mechanism for providing positional information via a morphogen gradient with his French Flag model

(Wolpert, 1969). In the model, each cell in a line of cells has the potential to be either blue, white or red. The line of cells is exposed to a concentration profile of a morphogen and each cell interprets the information from the concentration profile by varying its response to different concentration thresholds of morphogen: cells become blue, white or red according to their interpretation of the information. In general, it is accepted that Wolpert's definition implies at least two thresholds of morphogen concentration, or the division of the field into at least three different states.

In the majority of applications of this model, it is supposed that the gradient is fixed or the morphogen concentration evolves towards a stationary gradient, so cells differentiate according to their position within the field and the functional role of the gradient is to provide spatial information. However, there are patterning events observed in biological systems in which the gradient is not stationary: rather, it travels across the tissue in a wave-like fashion, conferring, as it passes, some degree of determination (Dubrulle *et al.*, 2001; Lin *et al.*, 2006). Typically this kind of gradient acts in an *all-or-nothing* manner with a single threshold: above threshold concentration, cells remain in an immature state and have no pattern forming ability; below threshold concentra-

*Abbreviations used in this paper:* AP, antero-posterior; ECM, extracellular matrix; PDE, partial differential equation; PSM, presomitic mesoderm.

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tion they are competent to form patterning structures. Thus, in this case, the wave may control both spatial and temporal aspects of development.

It is this latter case that we will discuss here, both in relation to pattern specification and morphological events. How may a wave of competence combine with a local patterning mechanism to control the spatio-temporal patterns visualised in the embryo? Our studies are motivated by two paradigms from developmental biology: (i) somitogenesis (segmentation of the AP axis of vertebrate embryos) and (ii) formation of the avian integument. We begin by outlining the main events involved in each process, in the context of waves and patterning. Next we introduce several well-known mathematical models for pattern formation, discuss their application to our biological examples and the role of waves in controlling pattern forming ability. We conclude with a short discussion: comparing and contrasting each model and its validity in the light of current experimental knowledge. Finally, we remark upon the new challenges that interdisciplinary modelling in this area has brought about.

### Somitogenesis

Somites are tightly bound groups of cells that lie along the AP axis of vertebrate embryos. They are transient structures and further differentiation of the somites gives rise to the vertebrae, ribs and other associated features of the trunk (Gossler and Hrabé de Angelis, 1998). Somitogenesis is tightly regulated in both space and time (Pourquié, 2003a), with each somite forming from a seemingly uniform field of cells via a mechanism that involves the interaction of a moving gradient of morphogen and a segmentation clock.

Somites form from the pre-somitic mesoderm (PSM), thick bands of tissue that lie along the AP axis. At regular intervals (every 90 minutes in the chick), groups of cells in the anterior PSM undergo changes in their adhesive and migratory behaviour and

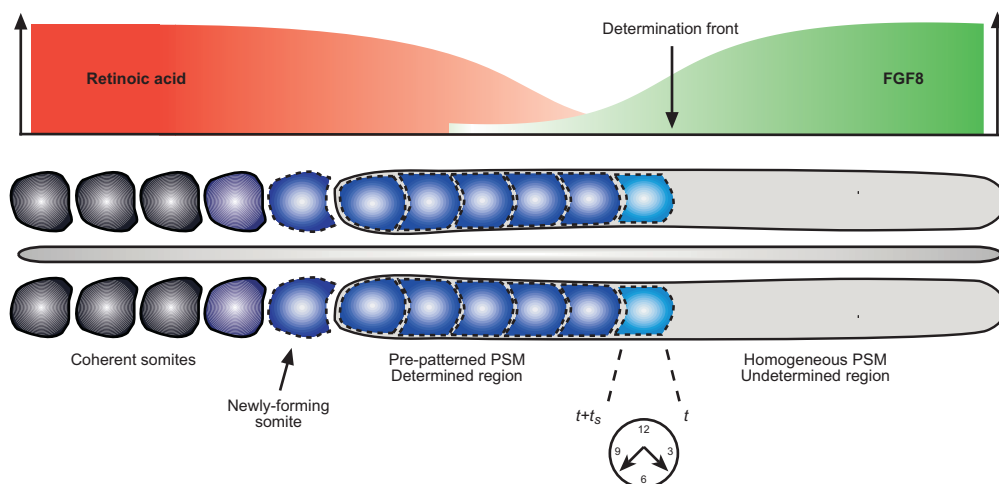
condense together to form an epithelial block of cells known as a somite. In this way, somites form in a very strict AP sequence (Gossler and Hrabé de Angelis, 1998; Stickney and Devoto, 2000; Stockdale *et al.*, 2000) and the budding of cells from the anterior end of the PSM compensates for the addition of cells at the posterior end of the PSM as the body axis lengthens. Each band of the PSM stays approximately constant in length throughout the process of segmentation and a wave of cell determination appears to sweep along the AP axis leaving somites in its wake (Baker *et al.*, 2008; Schnell and Maini, 2000; Schnell *et al.*, 2002).

Several genes are expressed dynamically in the PSM with cycling times equal to the time taken to form one somite (Déqueant *et al.*, 2006). For example, gene expression bands of *c-hairy1* (Palmeirim *et al.*, 1997) and *l-fng* (McGrew and Pourquié, 1998) sweep along the PSM in a posterior to anterior direction coming to rest in the newly forming somites. Expression is considered to arise as a result of a segmentation clock acting within PSM cells (Pourquié, 2001a).

Another gene with dynamic expression in the PSM is *fgf8*. A gradient of FGF signalling exists along the AP axis, with elevated levels in the posterior PSM gradually decreasing with movement in an anterior direction (Dubrulle *et al.*, 2001; Dubrulle and Pourquié, 2002). This gradient is mirrored by an opposing gradient of retinoic acid (see Díez del Corral *et al.*, 2002, 2003; Díez del Corral and Storey, 2004 and Fig. 1 for more details). As the axis elongates posteriorly and somites form anteriorly, the wavefront of FGF8 moves in a posterior direction along with the tail, so that the FGF8 gradient stays in a constant position relative to the PSM throughout somite formation. Hence cells are initially part of the region where FGF signalling prevails but as the gradient regresses they become part of the region where FGF signalling is virtually absent.

Dubrulle and co-workers (Dubrulle *et al.*, 2001; Dubrulle and Pourquié, 2002) have shown that the different levels of FGF8 in the posterior PSM coincide with regions of varying segmental determination and gene expression. In the posterior section cells are undetermined and plastic with respect to their future differentiation whilst in the anterior section cells are irreversibly committed to a particular developmental pathway (Dubrulle *et al.*, 2001). The border which separates these regions is known as the *determination front*. It has been shown that down-regulation of FGF signalling is necessary for cells to be specified as somitic, and go on to form part of a somite. Fig. 1 is an illustration of the vertebrate body plan during somitogenesis and the different regions of the PSM are clearly marked.

The first indications of somite patterning are observed as certain genes begin to exhibit striped expression patterns immediately anterior to the determination front (McGrew and



**Fig. 1. Schematic illustration of the vertebrate body plan during somitogenesis.** The top part of the diagram shows the opposing wavefronts of FGF8 and retinoic acid with the determination front (threshold level of FGF signalling) marked. The middle section of the diagram depicts the AP axis with several pairs of somites at the anterior end, followed by a sequence of five potential somite pairs, marked by a genetic pre-pattern, and the PSM. The bottom part of the diagram illustrates the interaction with the segmentation clock. The light blue blocks mark the position of the next somite pair to be specified: the posterior boundary is fixed by the level of the determination front at the time at which cells at the anterior boundary become able to signal.

Pourquié, 1998; Pourquié, 2001b, Pourquié, 2003b, Sawada *et al.*, 2000). In mouse and zebrafish, members of the *Mesp* family are periodically activated, with expression initially occurring on a domain larger than a somite and subsequently narrowing to occupy the future posterior somite half (Saga *et al.*, 2001; Sawada *et al.*, 2000).

However, the first overt signs of somite formation arise as cells begin to condense – undergo changes in their adhesive and migratory properties – and continue later as they differentiate (Gossler and Hrabé de Angelis, 1998; Keynes and Stern, 1988). The aggregation process occurs in the anterior portion of the PSM and is triggered by the interactions of cell adhesion molecules (CAMs) on the surface membrane of somitic cells. Somite formation in avian and mouse embryos is preceded by compaction of the anterior region of PSM and a simultaneous increase in cadherin, N-CAM and N-cadherin expression (Duband *et al.*, 1987; Kimura *et al.*, 1995). Integrins also play an essential role in somite formation (Drake *et al.*, 1992; Drake and Little, 1991). As a consequence it has been assumed that differential expression of adhesion molecules underlies the morphogenetic changes that take place during somite boundary formation (Murakami *et al.*, 2006; Newman, 1993).

Experimental data show that the extracellular matrix (ECM) filaments have a reproducible morphogenic destiny that is characterised by directed transport (Czirok *et al.*, 2004). Fibrillin 2 particles initially deposited in the PSM are translocated and eventually polymerise into an intricate scaffold of cables parallel to the AP axis. The cables coalesce near the midline before the appearance of the next-formed somite. This experimental evidence suggests that a direct translation from pre-pattern to coherent somites occurs via the changes in adhesive and migratory properties described previously.

There have been several models for somite formation and these have previously been reviewed by the authors (Baker *et al.*, 2003, 2006a, 2008). In all but a single case (Schnell *et al.*, 2002), the models specify only a pre-pattern for somitogenesis. The most recent models, both empirical and mathematical, are built upon a revised version of the *clock and wavefront* model which was first proposed by Cooke and Zeeman in 1976 (Cooke and Zeeman, 1976). The original model postulates the existence of a longitudinal positional information gradient along the AP axis of vertebrate embryos, which interacts with a smooth cellular oscillator (the clock) to set the time in each cell at which it will undergo a *catastrophe*. By catastrophe, they mean a rapid change of state, which could possibly be the change in locomotory and adhesive

behaviour of cells as they form somites.

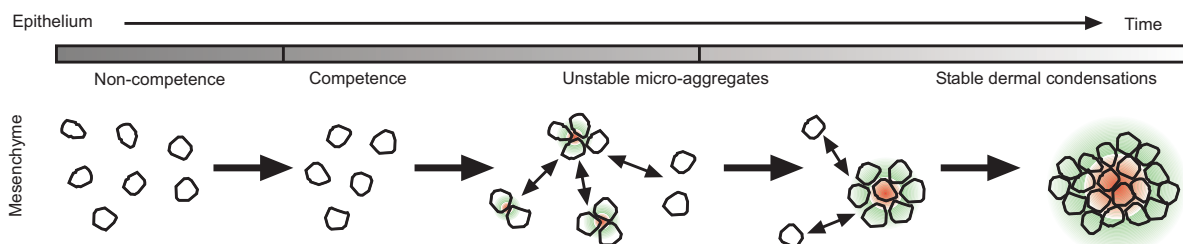
With the discovery of a segmentation clock (Palmeirim *et al.*, 1997; Pourquié, 2001a) and the wavefront of FGF8 travelling along the AP axis (Dubrulle *et al.*, 2001; Dubrulle and Pourquié, 2004), Pourquié and co-workers proposed a revised clock and wavefront model; involving the interaction of clock and wavefront to gate cells into somites. For a cell at a particular point along the axis, they assume that competence to segment is achieved only when the level of FGF8 falls below a certain threshold: the position of which defines the determination front (Dubrulle and Pourquié, 2002). In (Baker *et al.*, 2006a, 2006b) we developed a mathematical formulation of this model and we refer readers to Appendix A for more details.

### Feather germ formation

Feather buds, the precursors of the feathers, become visible shortly after fertilisation of the egg, at about 6-9 days. Each bud consists of a thickening of the top layer of skin with an aggregation of mesenchymal cells from the second layer of skin lying beneath it. The epithelial cells making up the top layer of skin are unable to move, whereas the underlying mesenchymal cells are motile and can move around in the ECM (Wolpert, 1998). Feather buds form via the interaction of these layers; first, the epithelium over the tract (the feather forming region) becomes competent and then, via mechanisms mediated by cell adhesion and regulated by reaction-diffusion and competition, the cells migrate to form individual feather primordia (Jiang *et al.*, 1999; Yu *et al.*, 2002). See Fig. 2 for more details of the feather bud formation process.

The nature of the patterns formed is controlled by the presence of various activators and inhibitors which promote and suppress bud formation, and by the initial density of mesenchymal cells. Addition of promoters of bud formation, such as FGFs and Follistatin, increases bud density, whilst addition of inhibitors such as BMPs have the opposite effect (Jung *et al.*, 1998; Widelitz *et al.*, 1996). The final pattern of buds and interbuds can be overlain by a pattern of activators and inhibitors of bud and interbud formation (see Fig. 2).

*In vivo*, feather buds first appear as dense regions on the skin surface; they develop in well defined lines and with a strict temporal ordering. The primary bud of each tract forms in the lumbar region at the level of the hindlimbs and patterning spreads bi-directionally along the midline axis. A wave of patterning then spreads out symmetrically and bilaterally from the midline (Jiang *et al.*, 1999; Lin *et al.*, 2006). Cells which lie initially in more lateral regions of the tract are unable to form feather buds until a wave



**Fig. 2. Schematic illustration of the processes involved in feather bud formation.** Initially the field is incompetent and cells cannot form buds. As a wave of competency passes, cells become able to form microaggregates and begin to secrete bud-promoting and -inhibiting factors. Promoting factors (red) recruit more cells into each aggregate whilst inhibiting factors (green) stop aggregates from becoming too large. Eventually, some aggregates are stabilised and go on to become dermal condensations.

of competency has passed (see Fig. 2): although expression of  $\beta$ -catenin has been shown to be a reliable marker for competency, it is not clear what drives this morphogenetic wave (Lin *et al.*, 2006).

Experimental techniques *in vitro* can be used to reset cells: embryonic chicken skin is dissected and the two skin layers are separated. With the mesenchymal cells dissociated from one another, the two layers are recombined. In this case, feather buds form simultaneously (Jiang *et al.*, 2004), showing that the morphogenetic wave is not essential for patterning. The effects of perturbation of bud density can also be seen *in vitro* using a fixed size piece of epithelium and varying the initial density of mesenchymal cells. Below a certain threshold, there is a complete absence of buds, but as the initial density is increased, regularly-sized buds form, until maximal packing is achieved with a hexagonal arrangement (Jiang *et al.*, 1999). Cell adhesion molecules have also been shown to play a role in pattern formation and their expression becomes progressively restricted to the feather buds as development proceeds (Jiang *et al.*, 2004).

These results suggest that there are two mechanisms at work *in vivo*: a positional information gradient, which renders cells competent to form primordia, overlying a local patterning mechanism which is dependent on a number of activators, inhibitors, cell adhesion molecules and also upon cell density. However, it is not yet known whether the local patterning mechanism can be divided into two separate stages of pattern specification and cell differentiation/rearrangement, or whether the processes are intimately coupled.

### Mathematical modelling in developmental biology

Mathematical modelling is now playing an increasingly important role in developmental biology: somitogenesis and avian appendage formation are two important paradigms for modelling in this field. Although still at an early stage, the experimental community is beginning to recognise that mathematical models allow one to put ideas and hypotheses into a concrete theoretical framework. Biological interactions and processes are often non-linear, and this is where intuitive, verbal reasoning may let us down: mathematical techniques allow us to analyse the effects of many interacting biological processes. Moreover, they allow us to direct research, discover/ask and even answer, pertinent questions and to design statistically sound and accurate experiments.

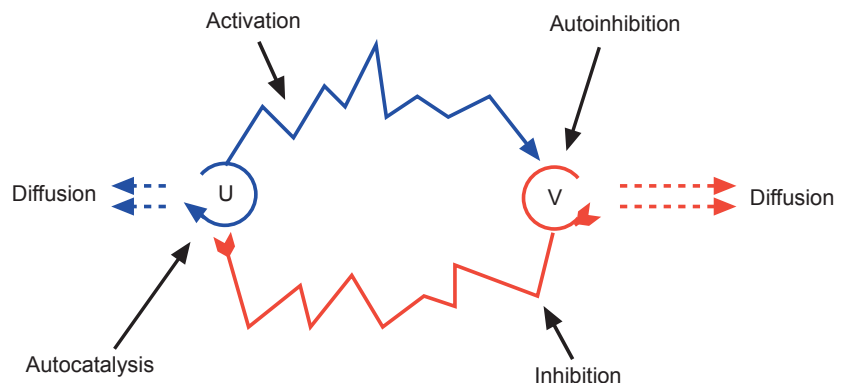
The impact of experimental perturbations, such as changes in the production and/or removal rates of a certain molecular constituent, can be mimicked in a mathematical model by changing, for example, certain parameter values. This is an important step in model validation: after one has verified that a model can display the results seen in wild type embryos, it is important to see whether it can display the behaviour observed when different model components are perturbed. If this is the case, then one can have greater confidence that the model captures correctly the biological phenomena being investigated: if not, some mechanism must be missing or incorrect, and current understanding is incomplete. This often leads to a greater insight into the

system.

In both somitogenesis and avian appendage formation, many of the molecular players have been characterised and laboratory techniques now allow extremely sophisticated experiments to be carried out. Both are excellent candidates for multiscale modelling: studies which couple events on the molecular level with those on the tissue level (Schnell *et al.*, 2007). In each case the processes involved include gene expression, cell differentiation, cell signalling, and biological clocks.

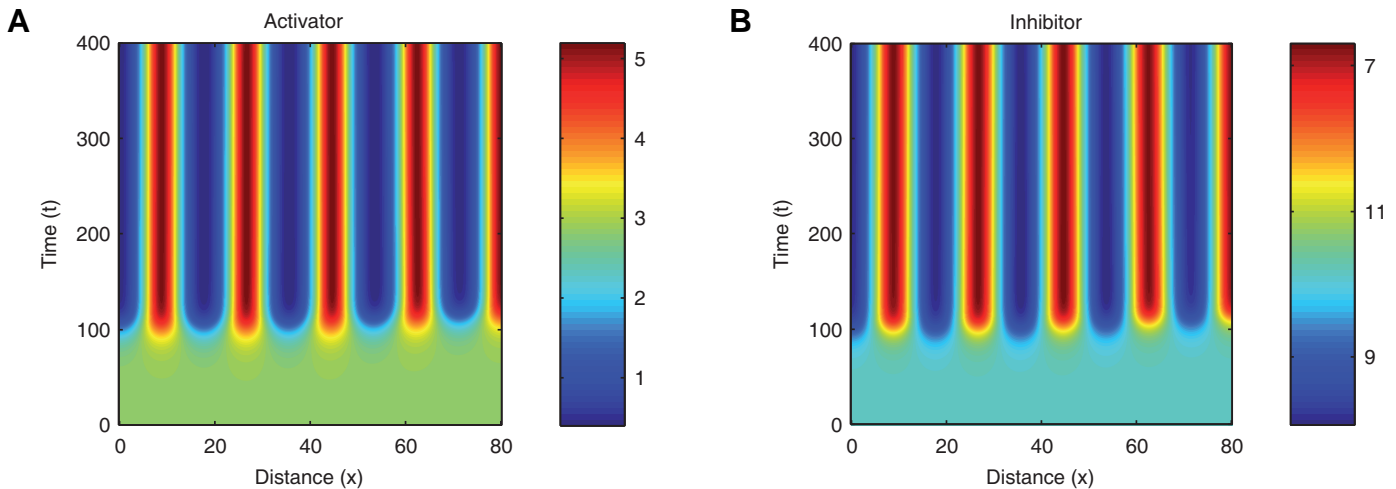
In this work, we are interested in how waves can interact with established patterning mechanisms to control spatial and temporal aspects of development. It is important here to draw a distinction between two different stages involved in patterning an embryo. The first is pattern specification, where cells become committed to following a specific developmental program by a pre-patterning mechanism that cannot be visualised simply by looking at physical properties of the cells, but may be driven by the creation of a genetic pre-pattern. The second stage involves the morphological events that take place during pattern formation. For example, changes in adhesion molecule concentration, cell motility, and the process of cell rearrangement. Often these stages cannot be separated; the processes are completely coupled by feedback mechanisms. However, an example of a process in which these two steps are clearly defined is somitogenesis. A wave of FGF signalling is present along the AP axis with a threshold level of FGF8 required for segmentation (Dubrulle *et al.*, 2001). First a genetic pre-pattern irreversibly marks the position of presumptive somites, and then cells undergo epithelialisation and somite boundary formation occurs (Gossler and Hrabé de Angelis, 1998). In feather bud formation, the initial pattern row forms along the midline of the embryo, with subsequent rows added on either side until the field is fully patterned – although no molecular players have been identified with this phenomenon, it is easy to imagine that a similar wave (be it in cell density, chemical concentration or adhesion molecules, for example) could play a role similar to that of FGF signalling in somite formation.

Below we outline a subset of models for patterning in developmental biology and describe their applications to somitogenesis and avian integument formation. The models include the reaction-diffusion model first postulated by Turing (Turing, 1952), the cell-



**Fig. 3. Schematic illustration of the interactions between activator ( $u$ ) and inhibitor ( $v$ ).** Arrowheads indicate catalysis whilst arrowtails indicate inhibition. Dashed lines indicate diffusion, with the length representative of the diffusion rate.





**Fig. 4. Numerical solution of a reaction-diffusion model in one spatial dimension.** Small initial fluctuations in an otherwise homogeneous system are amplified into a series of peaks and troughs in chemical concentration. See Appendix B for more details.

chemotaxis model of Keller and Segel (Keller and Segel, 1970) and Patlak (Patalak, 1963) and the mechano-chemical model of Oster, Murray and co-workers (Oster *et al.*, 1983). The models can be distinguished by those that simply have the ability to describe a pre-pattern for subsequent development, and those in which pattern specification is intricately connected to morphological events.

#### Reaction-diffusion models

In a seminal paper in 1952, Alan Turing first proposed the reaction-diffusion model for pattern formation (Turing, 1952). The mechanism, often termed *diffusion-driven instability*, involves the diffusion and interaction of two chemicals, known as morphogens. In short, Turing showed that without diffusion the system settles to a homogeneous steady state, but in the presence of diffusion, small fluctuations can become unstable and amplification of these instabilities leads to a spatial pattern in chemical concentration. Reaction-diffusion models have been postulated for a number of instances of pattern formation in biology, including fish and butterfly pigmentation (Kondo and Asai, 1995; Nijhout *et al.*, 2003; Painter *et al.*, 1999; Sekimura *et al.*, 2000), hair follicle initiation (Mooney and Nagorcka, 1985; Nagorcka, 1983-1984; Nagorcka and Mooney, 1982, 1985), feather germ formation (Jung *et al.*, 1998) and somitogenesis (Meinhardt, 1982, 1986).

One type of reaction-diffusion model works as follows: one chemical should be an activator ( $u$ ) and the other an inhibitor ( $v$ ). The activator should stimulate its own production (auto-catalysis) and production of the inhibitor, whilst the inhibitor should downregulate both its own production (auto-inhibition) and that of the activator (see Fig. 3). Both activator and inhibitor diffuse, but the inhibitor should diffuse more quickly. A system of non-linear partial differential equations (PDEs) can be constructed which describes the evolution of both chemical concentrations in space and time. A detailed outline of such a model is given in Appendix B.

Patterns form when a number of constraints are satisfied by the parameters describing activator and inhibitor kinetics (Gierer and Meinhardt, 1972; Murray, 2003). In this case, small disturbances to the system result in a pattern of peaks and troughs in chemical

concentration. Analytical results can be derived to show the range of parameter space in which patterning occurs, and also indicate the range of possible patterns *i.e.* the number of peaks in chemical concentration in each spatial direction. The equations can also be solved numerically – an example numerical simulation of a reaction-diffusion system in one spatial dimension is shown in Fig. 4. In this case, the field is initially homogeneous and small, random fluctuations are added to the  $u$  concentration profile. Over time, the fluctuations are amplified into a stable spatial pattern, with the wave length consistent with the range predicted by mathematical analysis (see Appendix B). An example of a numerical simulation of a reaction-diffusion system in two dimensions is shown in Fig. 5. In this example the homogeneous field breaks up into a pattern of spots of high chemical concentration.

In this model, there are two parameters, one which controls the rate of activator decay relative to inhibitor decay ( $b$ ) and one which controls the rate of inhibitor diffusion compared to activator diffusion ( $D$ ). In order for spatial patterning to be possible the following constraints must be satisfied:

$$0 < b < 1 \text{ and } Db > 3 + 2\sqrt{2}. \quad (1)$$

If  $b$  is decreased below  $(3 + 2\sqrt{2})/D$  then the spatially homogeneous steady state does not become unstable when diffusive effects are added and no patterning occurs. This is an effect that could be tested experimentally: if the activator removal rate could be increased by the addition of some drug, then the model predicts a loss of spatial patterning.

A reaction-diffusion model for feather bud formation can be supposed by assuming  $u$  is an activator/promoter of feather germ formation, FGFs 1,2,4 or Shh for example, and that  $v$  is an inhibitor of feather germ formation, BMPs 2,4,7 or retinoic acid, for example (Jiang *et al.*, 1999, 2004). In fact, Jiang and co-workers have suggested a discrete cell formulation of the model (Jiang *et al.*, 2004) which they term the *digital hormone model*. In both models, interactions between the activators and inhibitors lead to bud specification: a spatial pattern in chemical concentration that directs future cell movement and/or differentiation. For example, we may either suppose that cells migrate preferentially to areas of high activator concentration, or that cells with activator con-

centration above a certain threshold become specified as bud whilst others become specified as interbud. In either case, one would expect to visualise a chemical pre-pattern before a physical pattern is manifest, and the cells themselves not to play a role in shaping the pattern.

A basic reaction-diffusion model results in virtually simultaneous pattern specification, rather like that which occurs when skin cells are removed from the chick embryo, dissociated and cultured *in vitro*: all feather primordia appear simultaneously (Jiang *et al.*, 1999). In order to generate a sequential pattern we suppose that there must be some wave of maturation in the skin such that patterns are unable to form until the wave has passed and the field has, in some way, been designated as mature. The wave could be linked, for example, to cell motility, ability to respond to chemical cues or produce adhesion molecules. It may also be that the initial pre-pattern forms throughout the feather-forming domain and morphological structures can only form once the wave has passed, or that control of the pre-pattern is linked to wave propagation. These questions remain to be answered.

In the 1980's Meinhardt proposed a reaction-diffusion model for somitogenesis (Meinhardt, 1982, 1986), although it was not a reaction-diffusion model in the Turing sense. Similar to the ideas for feather bud formation, Meinhardt proposed that a gradient of positional information is present in the embryo which controls somite formation. He assumed that cells can be in one of two possible states, denoted by  $a$  and  $p$ , and that these states correspond, respectively, to the anterior and posterior halves of somites. If a cell is in state  $a$  then the genes responsible for synthesis of a substance A are turned on, and similarly for  $p$  and a substance P. The states  $a$  and  $p$  are such that they locally exclude each other but stimulate each other over a long range. Cells switch from one state to another until they reach a steady state: in this way a stable pattern of *apap...* stripes is formed. If cells are initially all in state  $p$ , the positional information gradient can be used to control the initial switch of cells to state  $a$ . The mathematical basis of Meinhardt's model is reviewed in (Baker, 2003, 2005).

### Cell-chemotaxis models

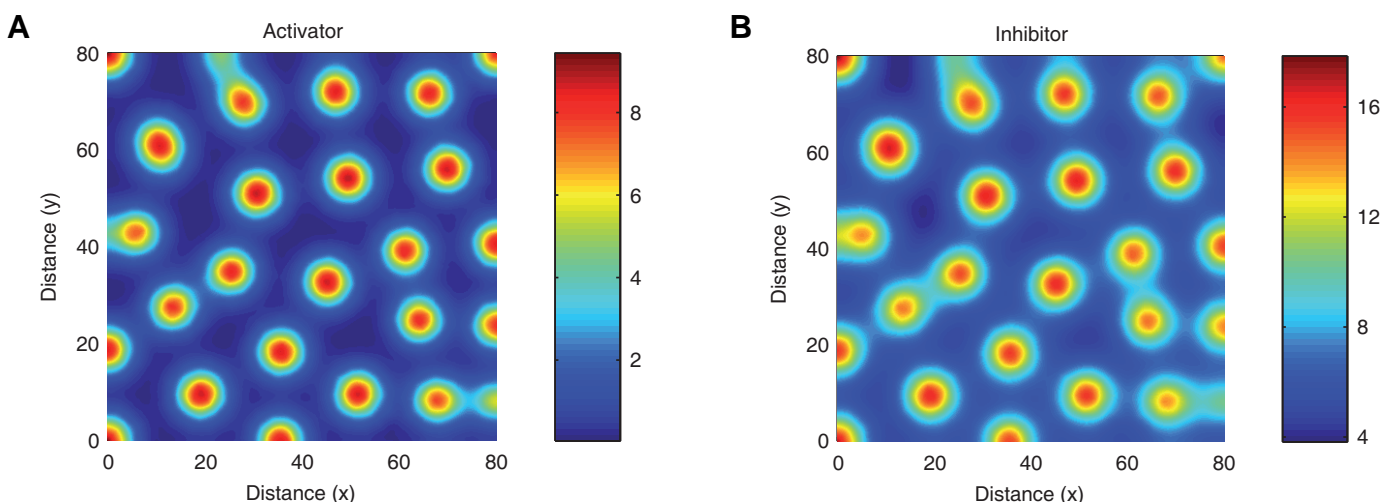
The most well known models describing pattern formation via

chemotactic movement are those of Patlak (1963) and Keller and Segel (Keller and Segel, 1970). The basic mechanism involves the differential movement of cells up gradients in chemical density (chemotaxis) and amplification of these gradients by localised chemical secretion. Analogous to the reaction-diffusion model, initial fluctuations in chemical concentration or cell density can become unstable and amplification of these instabilities leads to the formation of spatial patterns. However, the biological basis of the model is rather different: here cells play a role in shaping the pattern and the stages of pattern specification and cell re-arrangement cannot be decoupled.

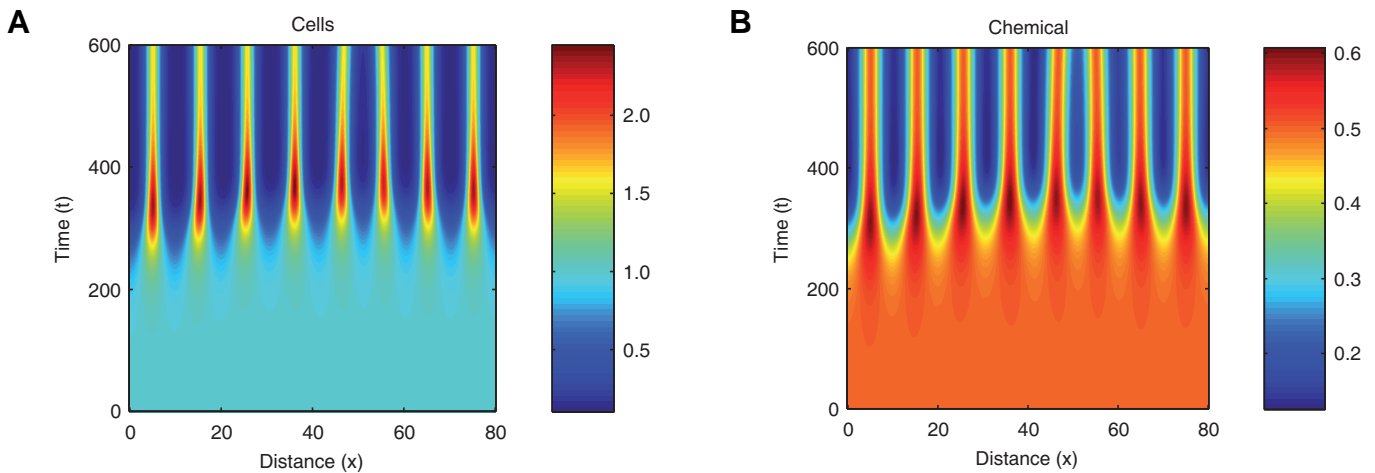
Since the original publications of Patlak, Keller and Segel, there has been much theoretical investigation of cell-chemotaxis models, see, for example, (Hillen, 2002; Newman and Grima, 2004; Othmer and Stevens, 1997; Schaaf, 1985). Chemotaxis models have been used extensively in developmental biology: from feather patterning to primitive streak formation (Painter *et al.*, 2000), enteric nervous system development (Landman *et al.*, 2003) and skin pigmentation patterns on the snake (Murray and Myerscough, 1991).

A general class of cell-chemotaxis models can be described as follows: we consider cell density ( $n$ ) and chemical concentration ( $c$ ). Cells move via random diffusion and also preferentially up gradients in chemical concentration. They may undergo proliferation and apoptosis, with these rates being dependent on local cell density and chemical concentration. The chemical diffuses randomly and is secreted locally by cells in the field. Once again, a system of non-linear PDEs can be written down which describe the aforementioned dynamics in terms of cell density and chemical concentration over space and time. A detailed outline of a cell-chemotaxis model is given in Appendix C.

As with reaction-diffusion models, spatially heterogeneous patterns in cell density and chemical concentration may form if certain constraints are satisfied by the parameters of the PDE model. Analytical results can be derived to uncover the range of parameter space in which patterning occurs, and also to indicate the range of possible patterns which may form. Numerical simulation of a cell-chemotaxis model in one spatial dimension is shown in Fig. 6: once again, spatial patterns arise from an initially



**Fig. 5. Numerical solution of a reaction-diffusion model in two spatial dimensions.** Small initial fluctuations in an otherwise homogeneous system are amplified into a spotted pattern of peaks and troughs in chemical concentration. See Appendix B for more details.



**Fig. 6. Numerical solution of a cell-chemotaxis model in one spatial dimension.** Small initial fluctuations are amplified into a series of peaks and troughs in cell density and chemical concentration. See Appendix C for more details.

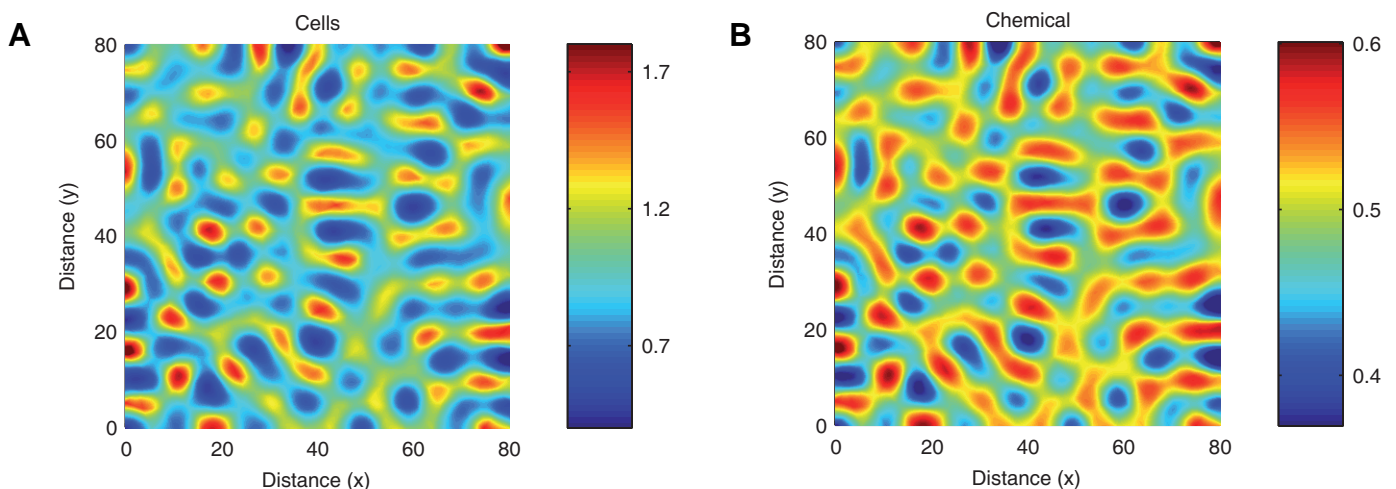
homogeneous field. As with the reaction-diffusion model, numerical simulation of the model can be carried out in two spatial dimensions, allowing us to display more complex patterning: this is shown in Fig. 7.

A cell-chemotaxis model for feather bud formation *in vitro* has recently been postulated by us. In this model the chemoattractant is supposed to be a member of the FGF family. Small random fluctuations in cell density or FGF concentration are amplified by a feedback loop of FGF production and cell movement. For example, a small peak in FGF concentration causes cells to move preferentially in the direction of the peak, where FGF production increases (due to increased cell density), which in turn induces more cells to move in the direction of the peak. Positive feedback competition between neighbouring peaks results in a pattern of cell density. Some peaks are eliminated due to competition while others stabilise and later become dermal condensations.

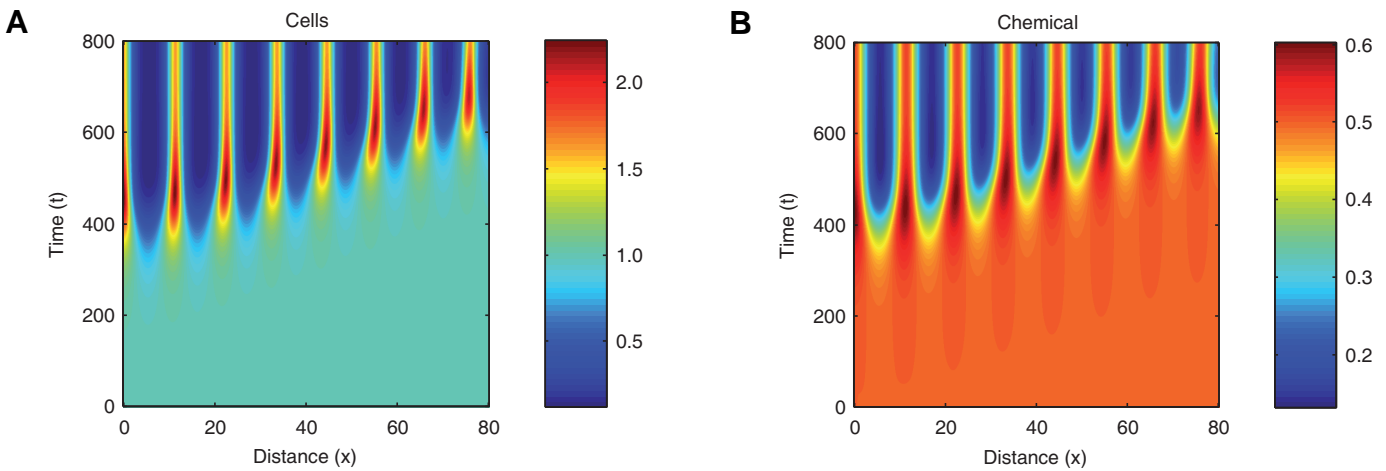
The question now to be asked is; how is sequential bud formation achieved *in vivo* with this model? It is well known that cell-chemotaxis models can produce propagating patterns of cell density if the initial disturbance in either cell density or chemical

concentration is localised to a specific point in the domain (Dee and Langer, 1983; Myerscough and Murray, 1992). Fig. 8 shows the results of a numerical simulation in one spatial dimension in which the initial cell and chemical fields were completely homogeneous, except for a small perturbation in cell density at  $x = 0$ . In this case, the disturbance appears to travel along the developmental axis, with regions close to the initial disturbance forming peaks and troughs in cell density and chemical concentration before those which are further away.

One could postulate that this is a sufficient mechanism for generating the morphologies seen *in vivo*. However, this mechanism is not robust in the sense that should a random perturbation arise in either cell density or chemical concentration in a region where the wave of patterning has not passed, a new wave of patterning will spread out from this point, destroying the order of pattern generation. Since all biological processes are subject to stochastic effects, from random movements to gene transcription and translation rates, it is likely that this scenario may arise. A more stable mechanism for patterning might involve a maturation/competency wave, which travels across the tissue and confers



**Fig. 7. Numerical solution of a cell-chemotaxis model in two spatial dimensions.** Small initial fluctuations are amplified into a complicated pattern of peaks and troughs in cell density and chemical concentration. See Appendix C for more details.



**Fig. 8. Numerical solution of the cell-chemotaxis model in one spatial dimension.** Initially, the field is supposed to be homogeneous throughout, with a small perturbation made to the cell density at  $x = 0$ . In this case, the pattern propagates across the domain, from left to right.

upon cells some patterning ability. This may be related to the ability of cells to produce chemoattractant, or to direct their movement towards such an attractant. These possibilities in the context of somitogenesis are the subject of current research by the authors and findings will be reported elsewhere.

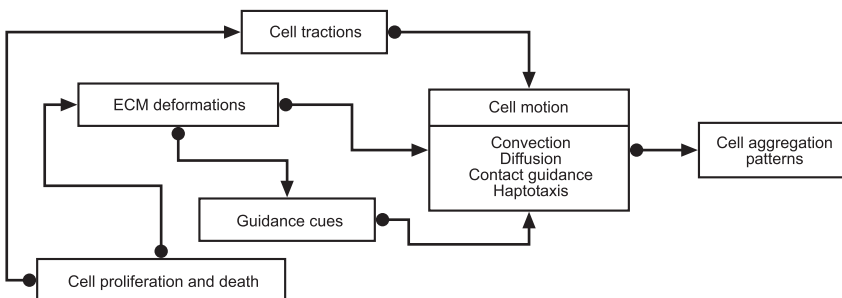
#### Mechano-chemical model

The mechano-chemical model was first proposed in 1983 by Oster, Murray and Harris (Murray *et al.*, 1983; Oster *et al.*, 1983). The model is based around the interactions between cells and the ECM, and the resulting forces that are generated as cells extend filopodia and adhere to the ECM. In much the same way as for a reaction-diffusion or cell-chemotaxis model one can show that small perturbations in cell or ECM density can become unstable and that amplification of these instabilities leads to a spatial pattern. However, the mechano-chemical models have a very different biological basis. As with the cell-chemotaxis model, cell rearrangement and pattern specification are coupled, but the prominent mechanisms driving cell movement are force-generated. Mechano-chemical models have been used to describe feather bud formation (Oster *et al.*, 1983), somitogenesis (Schnell *et al.*, 2002), limb chondrogenesis (Murray *et al.*, 1994; Murray and Maini, 1986; Oster *et al.*, 1985), wound healing (Murray *et al.*, 1988) and angiogenesis and vasculogenesis (Manoussaki, 2003).

Mechano-chemical models consider cell density ( $n$ ), ECM

density ( $\rho$ ) and the displacement vector of the ECM ( $u$ ), such that a material point in the ECM initially at  $x$  undergoes a displacement to  $x + u$ . The basic equation for cell density is in balance form: the rate of change of cell density in any small volume element is equal to the flux of cells in and out of the volume, plus cell proliferation and decay. Cell transport (flux) may arise due to convection, diffusion, haptotaxis (motion up adhesive gradients), chemotaxis and galvanotaxis (motion up gradients in electric potential) *etc.* It is also assumed that the presence of filopodia may allow cells to sense spatial gradients outside their local neighbourhood. A subsequent equation considers the mechanical interactions between cells and the ECM: it is supposed that the traction forces generated by cells are in equilibrium with the viscoelastic restoring forces of the ECM. The final piece of the model is a conservation equation for the ECM: in general it is only assumed that the ECM moves passively, by convection. For more details of the mathematical models see Appendix D. Fig. 9 illustrates the mechanisms that lead to patterning: a feedback loop arises since the guidance cues and ECM deformations, which lead to cell movement, are themselves controlled by cell movement.

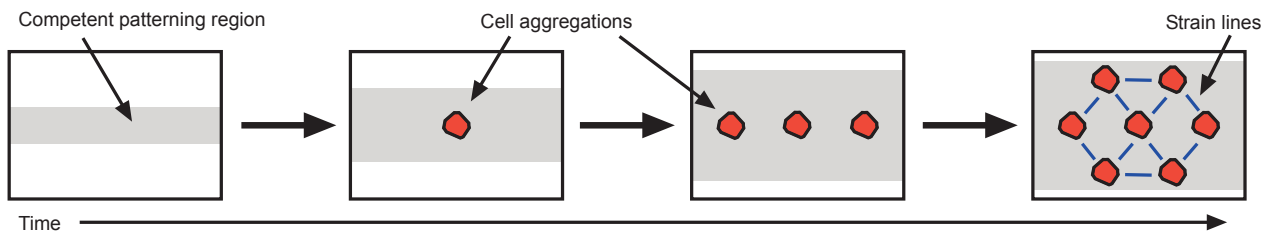
The mechano-chemical models are similar to the cell-chemotaxis models in that random fluctuations throughout the initial field lead to simultaneous patterning, whilst a small perturbation at a particular spatial location leads to sequential pattern formation.



**Fig. 9. The feedback mechanisms that lead to patterns in cell density.** Reproduced with slight modifications from (Murray, 2003; Oster *et al.*, 1983).

Once again, however, this is not a stable mechanism for sequential pattern formation *in vivo* as the homogeneous steady state is not stable to perturbation. It has been suggested that a stable, sequential patterning mechanism may arise if a change in traction force occurs as cells mature (Murray *et al.*, 1988). One can show mathematically that for low values of the parameter measuring traction force ( $\tau$ ) patterns cannot form – the spatially homogeneous steady state is stable. As the traction parameter rises above a certain threshold ( $\tau_c$ ), the steady state becomes unstable and we have a bifurcation from the homogeneous steady state to the patterning regime. If immature cells initially have  $\tau < \tau_c$  but  $\tau$  increases with age such





**Fig. 10. An illustration of the kind of propagating patterns that arise during pattern formation with the simplified model.** The grey shading represents the competent region of the patterning domain ( $\tau > \tau_c$  in our model) which expands as time proceeds. Patterning occurs as follows: (i) initially the competent region is too narrow to permit pattern formation; (ii) as the domain expands an initial row of aggregations starts to form; (iii) as the competent domain expands even further, a second row forms, offset from the first. Reproduced with slight modifications from Perelson *et al.* (1986).

that eventually  $\tau > \tau_c$ , the pattern may be obtained in a stable manner. See Appendix D for details of the mathematical analysis. We refer the reader to the work of Perelson and co-workers for detailed analysis of a similar system, including numerical simulation (Perelson *et al.*, 1986).

Mechano-chemical models have been used to describe feather bud formation (Murray *et al.*, 1988; Oster *et al.*, 1983). In each case the feather forming field is initially restricted to a homogeneous steady state; a wave of competency across the field renders the homogeneous steady state unstable and sequential patterns are generated. In the models the competency wave is either linked to a change in the cell traction parameter (as discussed here) or arises as a result of cell proliferation or a change in ECM properties. The patterns are initiated from the midline and so the pattern of ECM strains set up by the initial bud row biases the formation of secondary buds at positions displaced from the first row by half a wave length (Murray *et al.*, 1988). Fig. 10 illustrates pattern formation using the mechano-chemical model for feather bud formation.

There has only been one model for somite formation which considers the patterns generated when cells move up adhesive gradients (Schnell *et al.*, 2002). This model is based upon the stable signalling model for somitogenesis proposed by Maini and co-workers (Collier *et al.*, 2000; McInerney *et al.*, 2004) and, as such, does not employ the same mathematical mechanisms which lead to patterning as the mechano-chemical models described here. We are currently investigating an implementation of the mathematical framework discussed here in a new model for somitogenesis.

## Discussion

In this article we have outlined two patterning phenomena during development of the embryo in which a wave of competency appears to sweep over the pattern-forming field, conferring the ability upon cells to be recruited into a pattern. We presented a number of models which can be used to explain the phenomena, with the mathematical details and analysis presented as appendices.

Each of the three models presented – reaction-diffusion, cell-chemotaxis and mechano-chemical – are mathematically similar and so the same analytical tools are needed to study them. In each case, we start out with an initial field, close to a homogeneous steady state, and see a bifurcation, the onset of instability and transition to a spatially heterogeneous state, as one of the

model parameters is moved through a critical threshold. The resulting patterns depend on the specific parameter choices, initial and boundary conditions and domain size. In particular, each model displays a pattern that is critically dependent on scale and domain size: the larger the domain, the richer the patterning potential of the field. We now proceed by discussing the validity of each model as a pattern forming mechanism.

### Reaction-diffusion model

The main criticism that has been made of this model since it was first proposed in 1952, is that it is too simplistic and subsequently, the search for real biological examples of Turing morphogens has been neglected. The mechanism of diffusion-driven instability in chemistry has long since been documented (Castets *et al.*, 1990; Lee *et al.*, 1994; Lee *et al.*, 1993) but in biology there has even been evidence to dispute the mechanism (Akam, 1989). In a recent publication, Sick and co-workers (Sick *et al.*, 2006) provide the first real evidence for the Turing reaction-diffusion model in biology. They investigate the regulation of hair follicle patterning in developing mouse skin, proposing that the protein WNT and its inhibitor DKK constitute the activator and inhibitor, respectively, of a reaction-diffusion model.

A reaction-diffusion model specifies a pre-pattern which then guides subsequent morphogenetic events. One must assume, for example, that a certain path of cell differentiation is triggered in regions where the morphogen concentration lies above a threshold, or that cells respond chemotactically to the pre-pattern. In either case the process becomes multi-step but there is no feedback between cells and the chemical pre-pattern, which may increase the robustness of the system to environmental perturbation. One may also argue against the reaction-diffusion model on account of its sensitivity to changes in initial/boundary conditions and parameter values.

### Cell-chemotaxis model

Numerous chemoattracting and chemorepelling factors have been identified as acting during embryonic development – see, for example, (Affolter and Weijer, 2005; Keller, 2005; Yang *et al.*, 2002). Chemotaxis models integrate cell movements with the evolution of chemical concentration so that pattern specification is driven by cell rearrangements. As such, they are more able to adapt to changes in their environment and should be more robust. However, similarly to reaction-diffusion models, they suffer from an acute dependence upon initial and boundary conditions, and further, they also have a tendency to exhibit *blow up* – where

solutions become infinite in finite time.

### Mechano-chemical model

The mechano-chemical models are constructed from physico-chemical principles, which play a huge role in shaping the embryo. They characterise known cellular properties and as such, deal with easily measurable quantities such as cell densities, forces and tissue deformations. Even more so than in the chemotaxis models, these models couple changes in morphology with chemical patterning.

### Conclusions

We have described a number of models in this paper, each of which can be used to model a number of developmental phenomena and describe different types of pattern formation. At present, none of the prototype models that we have discussed explicitly incorporate the presence of a developmental wave which controls the timing of pattern onset. However, we have outlined ways in which this may be achieved within the modelling frameworks we consider. One of the challenges of mathematical modelling is to construct hypothetical experiments which allow one to distinguish between each model, and rule some out, if possible. They may range from the simple hypothesis that patterning ceases to occur in a reaction-diffusion model if the activator is removed, to hypotheses which concern the roles of different properties of the ECM or variation in the developmental wave speed or shape.

The other great challenge from a theoretical perspective is to begin to construct multiscale models for the patterning processes that take place during development. Such models incorporate important events at the cellular level with those taking place on a tissue and organ level; the result being models which capture the salient details of a mechanism without being overly complicated. For example, in a model for chemotaxis, one might consider events on a cellular level, such as signal transduction and cell polarisation, and how to integrate them into traditional models of chemotactic movement (Erban and Othmer, 2005, 2007; Firtel and Chung, 2000). Models should also contain biologically quantifiable parameters, so that experimental data can be integrated into the model in the form of parameter values or initial conditions. Alongside developments in mathematical techniques, numerical methods and algorithms need to be developed for simulating increasingly complex models.

The biological challenges, on the other hand, are to develop experimental techniques which allow for further identification and characterisation of the players involved in biological pattern formation and to design experiments in a systematic way that allow for the robust measurement of parameters such as diffusion and proliferation rates.

The future of truly interdisciplinary research in this area lies in the attitudes of both communities, theoretical and experimental: being committed to communication across specialist boundaries, and the development of tools and methods to facilitate the achievement of common goals. Mathematical modelling can be used as a tool to drive knowledge: to elucidate pertinent questions, test hypotheses and concepts in a rigorous framework and to devise experiments. In turn, the results from

these experiments must be used to refine the models, thereby establishing a feedback loop essential to biologically accurate model development.

### Acknowledgements

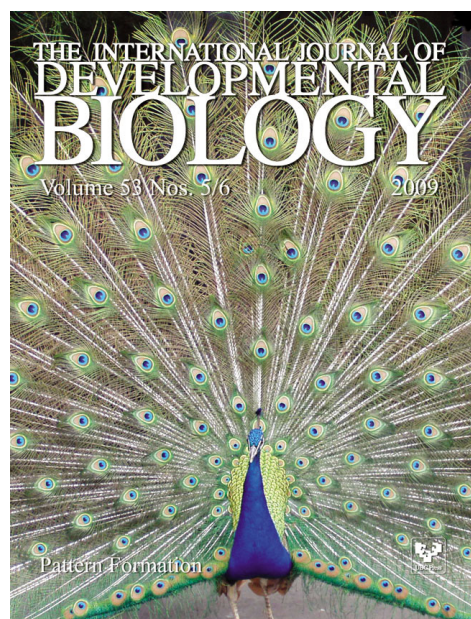
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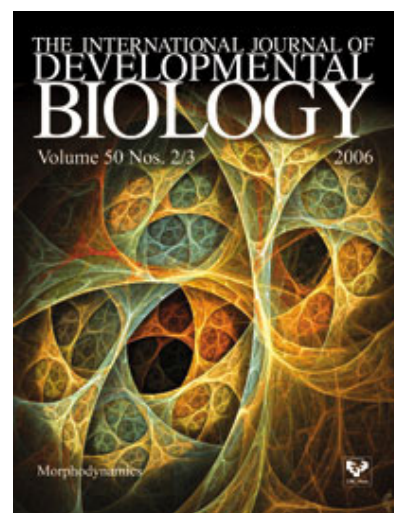
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