Spatially periodic patterns like zebra stripes or insect segments are mysteriously precise. How do they form during development? The deeper question, which lies at the heart of developmental biology, is: How do cells in different places within an embryo acquire different states of gene expression? In principle, there are three possible answers: either a cell's position causes its state, or its state causes its position, or both are caused by some third agent. These alternatives provide a framework for classifying theoretical models: positional information models, the most widely used today, belong to the first category. Dozens of rival schemes have been proposed, but until now they have been overlooked because they were obscured by confusing jargon, impenetrable mathematics, or tortuous logic.

In order to facilitate comparisons, this book (1) strips away all ad hoc assumptions to expose the central tenets of each model, (2) traces the historical roots and familial relationships among different types of models, and (3) illustrates the rules of each model in terms of how it would solve the basic problem. It also reexamines the computer metaphor in developmental biology: Are embryonic cells robots in disguise? The surprising answer, based upon their ability to perform Boolean logic, store and process information, execute iterative subprocesses, and malfunction in predictable ways, is yes!

This valuable guide to the menagerie of models about striped embryos and periodic patterning is required reading for biologists and zoologists interested in the pattern subfield of developmental biology.
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Preface

Lord Kelvin once remarked that he never fully understood a process until he could make a mechanical model of it [431]. This is a book full of models, some of which have profoundly influenced the history of developmental biology. The particular themes of the book, its iconoclastic style, and its focus on cybernetics are attributable to my own academic odyssey. My first exposure to model-building was when I worked as a computer programmer (under Seymour Papert and Marvin Minsky) in the Artificial Intelligence Laboratory at MIT where I was an undergraduate. The lab group was interested in how humans think, and most of the members designed programs to enable computers to 'converse', 'see', play chess, etc. Others, including myself, wrote interactive programs for teaching children about scientific principles. I created a simulated microworld of reptilian evolution where the user would select an environment, and the reptile population would attempt to adapt by randomly changing the ranges of variation of its anatomical parameters (via 'mutations'). What amazed me was how easy it was to reduce seemingly complicated structures (e.g. feathers) to simple equations that could be drawn by graphic algorithms. Only later did I encounter D'Arcy Thompson's 'On Growth and Form' [882] which makes this same point for anatomical shapes in general.

Had it not been for an apprenticeship with David Botstein and Ira Herskowitz, where I first encountered the joys (and sorrows) of biological research (my project concerned T7 phage genetics), I might never have left computer science. After applying to and being accepted in the graduate program in UC Berkeley's Molecular Biology Department, I was faced with the choice of an advisor and a project. One day, in our core course, John Gerhart delivered a fascinating guest lecture on Escherichian symmetries in virus heads, and soon thereafter I asked to work in his lab. At the time, he was investigating both frog and fly development, and I chose to work on flies because of their intriguing patterns of bristles. Chiyoko Toku-naga, an expert geneticist, had recently joined the lab after collaborating for many years with Curt Stern, a pioneer in the field of developmental genetics. She would tell me stories of the early days in Stern's lab, when he
was formulating his Prepatterning Hypothesis. His theory was rapidly being eclipsed by Lewis Wolpert's Positional Information Hypothesis, and it was illuminating to debate the pros and cons of the two paradigms with Chiyo and John. One of the essay questions in John's developmental biology course (co-taught by Gunther Stent) asked about a corollary of Wolpert's 'French Flag Problem': how could a concentration gradient of a chemical cause cells to produce blue, white, or red pigments in different zones? My answer was that if cells had vesicles containing pigments of different colors, then the chemical could cause osmotic swelling, and different colors could be released due to different bursting thresholds. I never liked that answer (though I did get an 'A' on the test) nor any of the then-popular explanations for how cells interpret positional information. I finished my dissertation in 1977, still skeptical about the ability of the new theory to explain patterns containing large numbers of identical elements such as bristles.

My postdoctoral years were spent in the think tank where the Polar Coordinate Model was born: the Developmental Biology Center at UC Irvine. The Center was a cauldron of ideas concocted by the faculty (including my sponsors Peter Bryant and Howard Schneiderman), the postdocs and graduate students, ... and repeatedly stirred and spiced by a parade of visiting scientists (cf. the book 'Cellular Basis of Morphogenesis' [223] to get a feeling for the ongoing ferment in this field). To make sense of the panoply of different ' patterning' theories, I began classifying them using a framework that I had devised as a graduate student. This book is the culmination of that effort, tempered by years of trying to teach squirming undergraduates about the wonders of gradients and clockfaces. I offer it as a field guide for others who have also felt lost in the wild menagerie of strange models that have lately seemed to multiply without limit.

Constructive comments on the manuscript were kindly furnished by Larry Blanton, Richard Campbell, John Gerhart, Kent Rylander, and Helmut Sauer (series editor). Technical jargon has been avoided wherever possible, but a college-level understanding of embryological principles is essential. newcomers to developmental biology may find the book's numerous citations useful as entry points to the field's vast literature. Finally, although I have endeavored to achieve an ecumenical scope, my parochial background as a fly researcher colors the text in many places. Do not interpret this slant as bias. The next breakthrough in this field could as easily come from a creature with green leaves as one with six legs.

Lubbock, Tex., December, 1991

Lewis I. Held, Jr.
A clever idea that solves this riddle was proposed by Hans Driesch [186, 1011] in 1901, and in 1969 Lewis Wolpert [997] formalized the concept as the 'positional information' hypothesis: (1) embryonic cells differentiate based upon information that they receive about their positions within a coordinate system, and (2) the boundary coordinates are specified independently of physical dimensions so that the entire system adjusts to the size of the field that it spans (i.e. the embryo or a part thereof). The details of how the mechanism works will be discussed later. What is important here is that the school of thought that has evolved from the concept of positional information is rooted in – and constrained by – the property of scaling invariance.

An alternative starting point for theorizing is the property of periodicity. Consider the following 'American Flag Problem' (fig. 1): how can an array of cells generate periodic patterns of elements like the stars and stripes of the American Flag? Positional information can also solve this riddle, but so can a host of other theoretical mechanisms that will be discussed in this book. A subordinate problem is how some periodic patterns routinely develop a precise number of elements (e.g. 10 fingers, 24 ribs, 32 teeth) or, in terms of the American Flag: how can an array of cells generate exactly 13 stripes and 50 stars? Numerical constancy is related to size independence and will be treated as a separate issue (chap. 7).

Types of Patterning Mechanisms

Embryonic cells can be committed to different pathways of development without manifesting any tissue-specific differences. For example, the wings and legs of fruitflies develop from separate groups of cells inside the larva. Prior to metamorphosis, wing and leg cells are indistinguishable cytologically, but when transplanted into the abdominal cavity of a host larva they form only wing or leg structures, respectively. They are said to be ‘determined’ [408] for different fates and to possess distinct ‘states of determination’ [339].

Given the notion of ‘states’ [795, 924, 941, 948] the problem of pattern formation can be reduced to simple mathematical terms. If \( \phi \) designates the position of a cell and \( s \) is its state of determination or differentiation, then any pattern can be represented as a set of ordered pairs \( (p, s) \). For example, if the positions along a line are numbered from 1 to 6, then an alternating pattern of black (B) and white (W) cells can be symbolized as

\[
[(1, B), (2, W), (3, B), (4, W), (5, B), (6, W)].
\]

The general problem thus becomes: ‘What causes the correlation of particular values of \( p \) and \( s \)?’ Whenever two entities are correlated in nature (e.g. fire and smoke, thunder and lightning, winter and spring), either one causes the other or both are caused by a third force. For \( p \) and \( s \), the possibilities are:

1. \( p \rightarrow s \): The position of a cell causes it to adopt a particular state.
2. \( s \rightarrow p \): The state of a cell causes it to adopt a particular position.
3. \( x \rightarrow p \& s \): Some third agent (\( x \)) causes the correlation of positions...
and states. An example of x is cell lineage, since a mother cell can divide directionally (causing the p of each daughter) and bestow instructions (x) asymmetrically.

These causal relationships define distinct classes of mechanisms, and most published models can be assigned unambiguously to a single class (table 1). (This same taxonomic scheme has been advocated by Steinberg and Poole [822]). By contrast, actual developmental pathways typically employ multiple strategies [180, 369, 459, 541, 646, 967] (see chap. 7). Operationally, the type of mechanism that is used at a given time and place should be discernable by transplanting a cell from one position to another. If the cell changes its fate (adopting a fate appropriate for its new position), then its position is causing its state. If the cell moves back to its original position, then its state was causing its position. If the cell neither changes

<table>
<thead>
<tr>
<th>Model or category</th>
<th>Derivative or related models</th>
<th>Distinguishing features</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Position-dependent (p→x) class</strong></td>
<td>The position of each cell (relative to field boundaries or neighboring cells) dictates its state of differentiation. The coordinate system allows the patterns to regulate.</td>
<td></td>
</tr>
<tr>
<td>Positional information subclass</td>
<td>Cells know where they are via coordinates which they 'interpret' as particular states of differentiation. The coordinate systems establish (independently of growth) by a scalar variable with fixed boundary values.</td>
<td></td>
</tr>
<tr>
<td><strong>Gradient Model</strong></td>
<td>Source-Sink Model [24, 149, 996, 997] Double-Gradient Model [14, 156, 157, 395, 744, 997, 998] Phase-Shift Model (wavelength = field length) [124, 44, 310] Gierer-Meinhardt Model (wavelength = field length) [279]</td>
<td>The coordinate system is established (independently of growth) by a scalar variable with fixed boundary values.</td>
</tr>
<tr>
<td>Progress Zone Model [867, 868]</td>
<td>Coordinates are assigned temporarily as cells exit a growth zone.</td>
<td></td>
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</tbody>
</table>

**Table 1 (continued)**

<table>
<thead>
<tr>
<th>Model or category</th>
<th>Derivative or related models</th>
<th>Distinguishing features</th>
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</thead>
<tbody>
<tr>
<td>Prepattern subclass</td>
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<td></td>
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<tr>
<td>Physical Force models</td>
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<tr>
<td>Traction Model [36, 351, 662]</td>
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<tr>
<td>Periodic Buckling Model [37, 38, 495, 835, 959]</td>
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<td>Physicochemical Model [288]</td>
<td></td>
<td></td>
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<tr>
<td>Reaction-Diffusion models</td>
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</tr>
<tr>
<td>Turing Model [31, 219, 907]</td>
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<tr>
<td>Gierer-Meinhardt models (wavelength &lt; 0.5 field length) [279, 570, 571, 573, 579, 580]</td>
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<td></td>
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<tr>
<td>Induction Model</td>
<td>Template Model [105, 106, 433, 434]</td>
<td></td>
</tr>
</tbody>
</table>

**Determination wave subclass**

<table>
<thead>
<tr>
<th>Chemical Wave models</th>
<th>Bicuspoid-Zhabotinsky Reaction [227, 470, 619, 643, 976, 979, 980]</th>
<th>Traveling (or standing) waves in the concentration of a diffusible molecule (or precipitate) arise through chemical reactions.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sequential Induction Model</td>
<td>Dityrosine cAMP-Signal Model [171, 172, 600, 635, 707, 729]</td>
<td>Each cell induces a neighboring cell to adopt a particular state.</td>
</tr>
<tr>
<td>Clock and Wavefront Model [134, 146, 1020]</td>
<td>Peridinin-Chlorophyll A Oscillator Model [136, 574]</td>
<td>Cells oscillate between two states and cease oscillating when a wavefront reaches them.</td>
</tr>
<tr>
<td>Inhibitory Field and Competence Wave Model</td>
<td>Fixation in the Antenna 1 Oscillator Model [1011]</td>
<td>Cells are not 'competent' to differentiate before a wavefront reaches them.</td>
</tr>
<tr>
<td>Poincare Oscillator Model [659]</td>
<td>Phyllotaxis models [593, 721, 883]</td>
<td>Cells that have adopted a 'preferred' state do so and inhibit neighboring cells from doing so.</td>
</tr>
</tbody>
</table>
its fate nor moves back, then the third type of mechanism would be indicated, though this result would also be expected after a p→s or s→p mechanism has been completed.

Vignettes of models from the first category are presented in chapters 1-4. Chapters 5 and 6 discuss models from the second and third categories, respectively. To facilitate comparison, the same linear array of six alternately black or white cells is used in all of the model illustrations, and each model’s rules for cellular decisions are listed. Extrapolation to 2 or 3 dimensions is usually left to the reader, as is generalization to patterns where the elements and intervals are multicellular, rather than unicellular. In chapter 7, an attempt is made to distill from the foregoing models a cybernetic ‘deep structure’ [111, 590] that directs development and constrains evolution. Though far from encyclopedic (for other compendia cf. Meinhardt [573], Murray [619], and Ransom [706]), the sample of models discussed in this book will acquaint the reader with the diversity of possible explanations for patterning phenomena. If there is a single lesson to be gleaned from this survey, it is ‘Wolpert’s Maxim’: do not infer process from pattern, since so many processes can produce the same pattern [998]. An awareness of theoretical alternatives can also serve as an antidote for the tendency to shoehorn new data into ill-fitting old paradigms, including the currently reigning theory, which was sired by Wolpert himself.
Chapter 1: Positional Information Mechanisms

Positional information models belong to the $p \rightarrow s$ category because they postulate a position-dependent assignment of differentiated states. Cells are supposedly informed of their positions, and this information causes them to select particular states according to predetermined rules. Individual models differ only in how they specify positional information.

The Gradient Model

The archetype of positional information mechanisms is the Gradient Model (Fig. 2a) [486, 798, 858, 881, 996, 997, 1003]. In the familiar ‘source-sink’ version [24, 149, 996, 997] the axioms are as follows. A diffusible chemical ‘morphogen’ (signaling molecule) is produced at one end of an array (the source) and consumed at the other (the sink). When a steady state is reached, the concentration has a gradient profile. Each cell records the concentration at its location as a coordinate and ‘interprets’ this ‘positional value’ as a state of differentiation. In figure 2a, the rule is that odd-numbered cells become black and even-numbered cells become white (even numbers) states of differentiation, though it is unknown whether cells can actually compute ‘even’ vs. ‘odd’ (see text). b Intercalation mechanism, based upon the Polar Coordinate Model [242, 245]. The outer coordinates are established first. Numerical gaps cause mitosis. Mitosis (thin arrows) are asymmetric since one daughter retains the parent number while the ‘intercalated’ daughter does not. Naïve cells compute a coordinate by averaging the coordinates of immediate neighbors. c Progress Zone Model [867, 868]. The gray semicircle represents the ‘progress zone’ where positional values increase incrementally. Extruded cells keep their values. d Conditional ‘IF-THEN’ rules (phrased in the second person as cellular commands) implicit in one or more of the models depicted above.
white, though, as discussed below, it is uncertain whether cells can actually perform such computations. The utility of the model is that it easily explains how patterns can regenerate after parts have been removed: either (1) cellular memories are erased and the mechanism starts over with fewer cells (‘morphallactic regulation’), or (2) mitoses at the cut edge produce cells having successively lower coordinates (‘epimorphic regulation’) [997].

The precepts of this model were based upon regulative phenomena in numerous developing systems, and its predictions have subsequently been tested in many of those same systems [1008]. Proof of its operation has been adduced for the anterior-posterior axis of Drosophila embryos, where the anterior morphogen has been identified as a DNA-binding protein encoded by the bicoid gene [187, 243, 649, 854, 856]. The first periodic patterns to arise along this axis are 7-striped arrays within which individual ‘pair-rule’ segmentation genes are transcribed [5, 397, 410]. For each gene, the stripes manifest a 2-segment periodicity (hence the name ‘pair-rule’), but the arrays for different genes are out of phase relative to one another, permitting an overlapping combinatorial code for cell states [269, 411, 492, 745]. Pair-rule gene expression is controlled via an intermediate echelon of ‘gap’ genes, so-named because mutant larvae are missing multisegmental swaths of cuticle [282]. Moreover, each stripe of every pair-rule gene appears to be under the control of a different set of maternal, gap, and other pair-rule genes [398, 668, 669, 722, 799, 815, 937], implying that this periodic pattern is merely an illusion created by a highly aperiodic mechanism [6]. However, other models (to be discussed later) challenge this conclusion.

In other versions of the Gradient Model, various amendments have been proposed: (1) all cells act as weak sinks, instead of one end acting as a strong sink [573, 866]; (2) all cells act as sources and establish a gradient by pumping the morphogen in one direction [134, 278, 486, 997]; (3) there are two opposing source-sink gradients, and cells measure the ratio of the two different morphogens [14, 156, 157, 395, 744, 997, 998]; (4) the concentration gradient is not created by sources or sinks but by a reaction-diffusion mechanism [279] or by a cell-signaling process that is formally equivalent to reaction-diffusion [22]; (5) instead of a concentration gradient, cells differ in the degree to which two oscillating chemical reactions are out of phase [124, 144, 310]; and (6) a gradient stage is avoided altogether by having two opposing wavefronts of cell-surface interactions directly establish a step-function concentration profile for the morphogen [566].

Positional Information Mechanisms

If an organ regulates epimorphically, then removal of the gradient's high point should cause the remaining tissue to produce a mirror-image duplicate of itself, whereas excision of areas lacking the high point should allow complete regeneration [441]. Hence, the high point should be locatable by systematically amputating various parts of an organ. When such experiments were performed with the developing wing of Drosophila, all four quadrants underwent duplication [245]. None regenerated. This paradoxical inability to find a high point led to a new model where positional information is not specified by conventional gradients.

The Polar Coordinate Model

Unlike the Gradient Model, the Polar Coordinate Model [81, 245] assumes that cells assess their positions by observing the coordinates of their immediate neighbors (presumably by contact between their cell surfaces [242]), rather than via a long-range diffusible signal. The model invokes polar (radial and angular) instead of Cartesian (perpendicular gradient) coordinates. Excision of a sector supposedly leads to (1) healing together of normally nonadjacent cells; (2) local proliferation in response to the positional disparity; and (3) an 'intercalation' zone which bridges the gap of coordinates via the shorter of the two possible routes (clockwise or counterclockwise) around the circumference. Pieces containing less than half the circumference (e.g. a quadrant) would therefore duplicate, regardless of their location in the organ as a whole (i.e. there would be no high point). Interactions between developing limb buds and regenerating limb blastemas in amphibians suggest that the same mechanism is used for both the development of the original pattern and its regenerative responses to surgical manipulations [612–614].

How would the coordinate system arise during normal development? If the peripheral coordinates are established first, then newborn cells could adopt intermediate coordinates until all gaps have been eliminated (fig. 2b) [153, 242, 245, 614]. An intercalation mechanism can explain several phenomena which the basic Gradient Model cannot, including: (1) why some organs never exceed a definite size even when given additional time to grow (because growth should stop automatically when all coordinates are present; double-gradient models can also explain determinate growth) [14, 79, 80, 245, 997]; (2) why defects in cell adhesion can lead to overgrowth (because adhesion should be crucial for contact-mediated com-
munication between adjacent cells, less so for diffusible signals) [78, 427, 548]; and (3) why cell death during the growth phase can cause pattern duplications (because removal of more than half an organ should cause the remaining piece to duplicate at any stage, not merely after growth has ceased and all coordinates have been specified) [75, 286, 287, 420, 690, 694, 749, 790].

Tests of the Polar Coordinate Model in insect, salamander, and chick limbs have yielded extensive supportive evidence, implying that vertebrates and arthropods use a common strategy to construct their appendages [81, 415, 421]. However, many regulative properties of chick wing buds can also been interpreted in favor of a gradient mechanism [866], raising doubts about the monophyly of the process, and questions remain about other phenomena which the model cannot easily explain [242, 441].

In an effort to alleviate some of the model's shortcomings, other authors have proposed (1) specifying the angular and radial coordinates via the ratio of two morphogens [1, 786]; (2) retaining the rules for intercalation but computing positional disparities from Cartesian coordinates [153, 305, 441, 445, 448, 747, 977]; (3) partitioning the coordinate system into discrete territories which intercalate only when an entire territory is missing [311, 575, 785, 793]; or (4) dispensing with coordinates altogether and using a 'smoothing' rule to control intercalation [509, 510, 597, 977, 978].

The Progress Zone Model

Though sometimes categorized as a gradient model [1008, 1009], the Progress Zone Model is uniquely different from any of the mechanisms discussed thus far. It emerged from experiments on the wing rudiments of chick embryos. The chick wing develops from a bud which grows mainly in a 'progress zone' at its tip. Reciprocal grafts between young and old buds led to the idea that the proximo-distal coordinate of a cell reflects the length of time that it spends in this zone [867, 868]. The cells would thus acquire positional information via temporal information (they presumably can measure time and stop their 'clocks' when they exit the zone; fig. 2e) in contrast to conventional gradient models (where positions are specified independently of growth) and intercalation mechanisms (where signaling is correlated with growth but not with time per se).

Puzzles and Paradoxes

Can Cells Actually Perform Mathematical Calculations?

All positional-information models assume that cells record their positions as a numerical quantity ('positional value') [997, 1004]. Epimorphic regulation requires that new numerical states be computed from old ones, and the Shortest Intercaletion Rule [245] demands that cells choose the smaller of two numbers. Whether cells can actually perform such computations is an interesting question. Precedents do exist for elementary mathematical capabilities in some cells. Thus, animal neurons can compute the sum of positive and negative inputs via effects on their membrane potential [525, 724], and Drosophila and nematode cells can assess the ratio of X chromosomes to autosomes during sex determination [382]. However, some patterning models require cells to compute cosines or even more complicated functions [153, 747, 786]. Must cells have the equivalent of a secondary school education in order to participate in pattern formation? Chapter 7 explores the limited abilities of embryonic cells to store and process information, and the general conclusion is that cellular 'intelligence' is closer to a kindergarten level.

How Does Interpretation Work?

The issue of cell 'intelligence' is especially troubling with regard to how cells interpret positional information [134, 1008]. For the Gradient Model, the orthodox view is that morphogen-sensing genes have different concentration thresholds that are sharpened by autocatalysis and intergenic repression [14, 34, 229, 456, 512, 540, 572, 577]. Evidence supporting this view has come from studies of Drosophila gap genes [402, 474, 499, 854] and Xenopus mesoderm inducers [322, 323]. (Unorthodox interpretive schemes have been proposed by Babloyantz [23, 24] and Goodwin [306].) However, the level of detail in most anatomical patterns is orders of magnitude greater than a French Flag (or a striped Drosophila embryo) and the theoretical precision of concentration sensing is crude by comparison [136]. Conceivably, organs could be subdivided into hierarchies of nested gradients [134, 321, 558, 997], permitting a serial combinatorial code of positional values, but decoding would still be a problem [1008, 1011]. To appreciate the dilemma, consider that each of the million-or-so hairs in your skin would have to possess a unique 'area code' [277] and decipher its code by looking it up in the genetic equivalent of an enormous 'area-code/differentiated-state' directory [134, 138, 146, 540, 706] (cf. Beardsley [48]
and Davidson [162]). A similar dilemma pesters the cognitive maps that are supposed to control human limb movements [21, 61].

**Collective Amnesia?**

An embryo can theoretically use a single coordinate system to specify anatomical patterns in many different organs by merely changing the rules for how the coordinates are interpreted [997]. In that case, cells would need to identify their 'organ state' (e.g. leg vs. wing) before deciding how to translate their coordinates into cellular states of differentiation (e.g. neuron vs. myocyte). Mutations in genes that encode organ shapes should cause the cells of one organ to construct an anatomy typical of another, as opposed to *cell type* interconversions [63, 65, 92, 369, 599, 652, 891, 916, 938]. Indeed, many such 'homoectic' mutations have been found in Drosophila [7, 450, 488, 547, 666]. For example, mutations in several Drosophila genes can cause a partial transformation of the antenna into a leg. Strangely, a clonal analysis of leg-tissue islands in the antennae of Antennapedia flies showed that the transformations occur concomitantly in groups of neighboring cells, i.e. via *proximity*, not via pedigree [692]. Similarly, another type of homeotic transformation (termed 'transdetermination' [339]), which is routinely encountered during long-term culture of nonmutant Drosophila tissues, also affects nonclonal groups of cells [263] (cf. Karlsson [432] for distinctions between transdetermination and ordinary homeosis). The perplexing implication is that organ states such as 'legness' or 'wingness' may not be properties of single cells, but rather of *cell clusters* (cf. Chambon [105]). Xenopus muscle differentiation likewise appears to involve a 'community effect': a cell will only shift its fate (in response to an inducing signal) if a sufficient number of its neighbors also does so [334].

**The Antenna-Leg Paradox**

Within the Antennapedia antenna, specific leg structures always develop in predictable positions, allowing a mapping of corresponding antennal and leg domains [265, 693]. The map has been construced as evidence that the antenna and second leg interpret the same coordinate system according to different sets of rules (cf. Haynie and Bryant [362]). Given this apparent homology, however, their regulative behavior is difficult to understand. In Drosophila the external structures of the head and thorax develop from separate pockets of epidermis called 'imaginal discs' [266]. The second leg develops from its own disc, but the antenna comes from part of a disc that also forms the eye. When the eye-antennal disc is bisected, the eye portion regenerates an antenna, and the antennal portion duplicates [262, 264], implying — according to the Polar Coordinate Model — that the antenna comprises less than half of a larger eye-antennal regulative 'field' [408, 929]. By contrast, when parts of the second-leg disc are removed, it behaves as an entire field [283, 749]. How can less than half of a coordinate system equal a whole coordinate system? Conceivably, the eye may constitute an outer annulus of the coordinate system, rather than an oversize sector. A related riddle is: if the arista (the feather tip of the antenna) is homologous to the tarsus according to the Antip map and a comparable map inferable from spineless-arista-pedia phenotypes [86, 851], then why does the boundary line separating the anterior and posterior 'compartments' [151] of the eye-antennal disc bypass the arista entirely [604] (cf. Brower [73]), whereas it bisects the tarsus all the way down to the claws [493, 824]? The importance of this question stems from the supposed significance of the anterior-posterior compartment boundary in general [151, 491, 556] and its presumptive role in causing appendage outgrowths in particular [126, 267, 574, 576, 578].

**Warped Coordinate Systems?**

Another peculiarity of Drosophila leg discs is that, unlike the wing discs, one quadrant (the upper medial one) can regenerate the remaining three-quarters of the disc [769]. The Polar Coordinate Model explains this oddity by assuming that more than half of the leg's circumferential coordinates are crowded into this quadrant [245]. The difficulty with this ad hoc remedy, however, is that it creates a new problem. If growth stops when all discrepancies between adjacent positional values have been eliminated [80], how can one part of an organ acquire a three-fold higher density of coordinates?

**How Do Body and Limb Coordinate Systems Mesh?**

The Drosophila embryo apparently uses a Cartesian (anterior-posterior and dorso-ventral) coordinate system [649], but the imaginal discs, which arise as inpocketings of the embryonic body wall [44, 123] use a polar coordinate system [245]. Do disc cells algebraically convert one system into the other, or do they erase their positional memories and create a polar system de novo [242]? (The same dilemma applies to body vs. limb axes in amphiobians [441].) Attempts to reconcile the two coordinate-system models [267, 575, 578, 786, 971] have not been wholly satisfying.
Chapter 2: Prepattern Mechanisms

Models Involving Physical Forces

Physical forces can create periodic patterns in inert matter, e.g. the rings of Saturn (gravity); ocean waves, sand dunes, mackerel clouds (fluid mechanics); and the harmonic waveforms of musical instruments (vibrations) [16, 230, 703, 734, 735, 949]. The physical properties of cells and their extracellular materials (e.g. viscosity and elasticity) can also produce local deformations in response to internal or applied forces [30, 36, 288, 351, 619, 637, 650, 661, 662, 944, 950], and these distortions could theoretically promote the development of structures.

Before the ascendency of the positional information paradigm, the dominant idea in the field of pattern formation was the concept of ‘prepatterns’ proposed by Curt Stern [831, 832] in 1954. Stern imagined that epithelial folds could cause ‘stress points’ (fig. 3a) where structures such as bristles might be induced:

The larval imaginal discs (of Drosophila) are made up of cell layers that are folded in complex ways. Let us postulate that differentiation of bristles occurs at those points at which folds cut across each other. According to this hypothesis, an allele that leads to differentiation of a specific bristle would be involved in provoking the formation of specific folds. Another allele whose phenotypic effect does not include formation of the bristle would be responsible for a different kind of folding of the imaginal disc. The different types of folding of the discs would constitute different patterns. Since these patterns would precede the appearance of their corresponding bristle patterns, I refer to them as prepatterns [835].

The foundations for the prepattern hypothesis were laid by Stern’s mentor Richard Goldschmidt, who (1) showed that some color patterns on lepidopteran wings are preceded by latent prepatterns in the rates of wing

Fig. 3. Prepattern mechanisms. Prepattern models postulate that the visible pattern of structural elements (in this case, black cells) is preceded by a nearly congruent (isomorphic) array (‘prepattern’) of sites (‘singularities’) which prompt the cells to differentiate as pattern elements. Singularities may differ from the background either physically or chemically (or both). a Physical Force Model, based upon Stern’s original notion of prepatterns [835]. A layer of cells is compressed by an external force (schematically depicted as a spring) which causes it to buckle. The stresses at the apices of the folds then function as singularities, causing the cells there to become black. (Buckling forces have been implicated in causing corrugations in the cerebral cortex [495], in the ciliary body of the bird eye [37, 38], and in insect tracheae [959] and tarsi [931]). b Reaction-Diffusion Model, of the kind proposed by Turing [907]. Chemically reactive molecules diffusing at different rates cause the concentration of a product – the ‘morphogen’ (M) – to peak at ‘wavelength’ intervals. Wherever the concentration exceeds a threshold (dashed line), the cell at that site becomes black (cf. Waddington [931]). c Induction Model, analogous to neural induction in vertebrates. An underlying ‘template’ pattern induces corresponding states of differentiation in an apposed layer of naïve cells. d Conditional ‘IF/THEN’ rules implicit in the models depicted above.
scale maturation [293, 294, 803]; (2) reviewed the literature on pattern formation in 1938, including a conjecture by Krieg (1922) that the striped arrangements of hair follicles in the tiger embryo (another example of a prepattern) are attributable to ‘tensions within the time of pattern determination [294]; and (3) speculated on a primary role for ‘growth tensions’ in tissue patterning [294].

Because a prepattern supposedly arises from forces within a whole tissue, it should be resistant to local perturbations. Thus, if a mutation alters a prepattern, then a few mutant cells in a wild-type background should not disturb the overall (wild-type) pattern but rather should acquiesce ‘nonautonomously’ [609, 859] in its formation. Given this logic, Stern was surprised when the first Drosophila mutation that he tested in this way behaved autonomously. In homozygous flies, the mutation achaete causes the absence of a specific thoracic bristle. In genetically mosaic flies where most of the thorax consists of wild-type tissue, achaete cells typically fail to form a bristle when they reside at that site—in effect ignoring the surrounding majority of cells [831]. From this result Stern reasoned that prepatterns cannot be sufficient for the induction of structures: cells must also be ‘competent’ to respond to signals from the prepattern, and achaete must be affecting the competence of cells at a specific site. For example, cells might have a threshold of strain above which they become bristles, and achaete raises the threshold at one location. When more than a dozen other pattern-affecting mutations were similarly tested in genetic mosaics, most were found to also behave autonomously [888]. Among them was a homeotic mutation—spineless-aristapedia (ss) — which (like Antennapedia) causes a partial transformation of antenna into leg. According to the prepattern hypothesis, the autonomy of ss indicates a hidden prepattern for leg structures in the developing antenna (to which only the mutant cells can respond) [691, 727], and because other homeotic mutations also manifest autonomy [505, 506, 603, 887, 889], one is led to the absurd inference that each imaginal disc must contain a hidden prepattern for every other disc [441]. Positional information provides a more plausible hypothesis since such mutations could simply be altering the rules by which an invariant system of coordinates is interpreted [888, 997]. Consequently, prepattern models have waned as explanations for adult cuticular patterns. However, they have recently experienced a revival vis-à-vis embryonic body segmentation because of reaction-diffusion schemes that can explain the striped patterns of segmentation gene expression [353, 478, 480, 539, 624, 625].

Reaction-Diffusion Models

Two years before Stern published his hypothesis, Alan Turing, a founder of modern computing [381], described a clever model based upon chemical prepatterns [907]. By postulating imaginary chemical reactions between molecules which diffuse at different rates, Turing showed that a homogeneous distribution of the molecules is unstable under certain conditions [219, 615]. Statistical fluctuations become amplified into peaks and troughs of concentration (fig. 3b) at wavelengths which depend upon the relative rates of reaction and diffusion. Only recently have actual chemical reactions been found which do indeed behave in this way [500, 667, 689, 981]. In two dimensions, Turing’s equations can produce periodic patterns reminiscent of zebra stripes or leopard spots [31]. Virtually any periodic pattern in any dimensional space can be simulated by the reaction-diffusion schemes of Alfred Gierer and Hans Meinhardt [279, 570, 571, 573, 579, 580] and others [71, 619, 625, 626, 642, 643], who have modified the parameters of catalysis and diffusion in Turing’s model and added new assumptions. Generally, these types of models predict certain pattern modulations as a function of shape [288], and the predictions are strikingly confirmed in the coat markings of mammals [32, 33, 616-618] and the fruiting buds of slime molds [90, 567, 568].

A curious property of reaction-diffusion models is ‘stochastic indeterminacy’ [313, 642, 643, 1013]: the final configuration of the pattern elements cannot be exactly predicted from the starting conditions. (Positional information models are deterministic: they should yield identical patterns from trial to trial.) Indeed, many anatomical patterns that manifest a certain regularity at one level are indeterminate at another level [948, 955]. For instance, human fingerprint patterns manifest a uniform ridge spacing, but ridge configurations are not identical in identical twins [154] (cf. freckles). Such patterns are ‘epigenetic’ [408] insofar as their features are not specified genetically. Presumably, the genes merely establish the starting conditions (e.g. reactant concentrations), with the outcome being dictated by the same random perturbations (e.g. concentration fluctuations) that initiate the process [307, 308]. Mechanochemical models, which combine aspects of both physical-force and reaction-diffusion mechanisms, have been designed and, in some cases, augmented with further assumptions to make them less indeterminate [288, 619, 660, 663].
Induction Across Layers

A trivial explanation for the origin of a pattern is that it is imprinted from a "template" prepattern in an apposed layer of cells (Fig. 3e) [105, 106, 409, 433, 434, 754, 927, 928, 1001]. Such inductions are common in vertebrate skin. Thus, bird feathers and mammalian hair are primarily epidermal in construction, but their positions are determined by underlying clusters of dermal cells [776, 805]. In Drosophila wings, the veins in the ventral layer are induced by those in the dorsal layer [251]. A peculiar variation on this theme is the long-distance induction of "neurobarrels" in the rat trigeminal system by aferents from the facial whiskers [767]: the whisker-vs.-neurobarrel patterns are remarkably isomorphic, and the cauty of particular rows of whisker follicles in a neonate causes a rostrocaudal cascade of abnormalities in the corresponding neurobarrels of the brainstem, thalamus, and somatosensory cortex [45, 457]. In an analogous manner, optic cartridges of second-order laminar neurons are induced by afferent retinal axons in Drosophila [586] and other insects [569].

The Preformationist Paradox

The induction of one pattern by another begs the question: How does the inducing pattern arise? Indeed, the entire prepattern school of thought has been criticized for implying an infinite regression of patterns induced by prepatterns [756, 803, 927, 997]. Historically, this criticism was justifiably leveled against the antiquated notion of preformationism, which argued that eggs (or sperm) contain preformed homunculi which, in turn, must have eggs bearing smaller homunculi ad infinitum [408, 602, 655]. The objection would be legitimate in this case if prepatterns could only be established by induction, but they can also arise de novo via physical forces or chemical reactions as discussed above. Comparable misunderstandings have repeatedly arisen from a failure to appreciate the cardinal distinction between the rules that generate a pattern and the information content of the pattern itself [14, 15, 69, 504, 826, 828, 1010].

Prepatterns vs. Positional Information

All prepattern models employ 'singularities' [834, 835] (sites where physical or chemical parameters differ from the background) as cues for inducing structures. A structure should form wherever a singularity is present unless certain cells are unable to respond. Thus, a prepattern and its subsequent pattern can be isomorphic [625], or the prepattern may have extra 'cryptic' singularities [561, 833]. Prepattern models differ from positional information models in several key respects [136, 625, 888, 998, 999, 1007, 1008]:

1) Identical structures use identical signals. For a pattern of 3 bristles, there would be 3 identical singularities, all of which would directly signify 'Make a bristle', so the cells would never know their positions. In contrast, positional information would use 3 different positional signals as bristle commands—e.g. 'You're at (x1, y1)', 'You're at (x2, y2)', and 'You're at (x3, y3)'—and hence the cells would know where they are.

2) Cells can be 'stupid'. If a cell is located at a prepattern singularity, then all it must do is switch its state relative to the cells of the background. There is no true interpretation stage: a nudge suffices. With positional information, however, cells must not only be 'bilingual' (able to translate positional coordinates into differentiated states) but their vocabulary must be as large as the number of pattern elements, since each location uses a different signal even if the elements are identical. The distinction is analogous to bitmaps vs. vector representations in computer graphics: in bitmaps (= coordinate systems) the states of all pixels (= cells) must be specified regardless of the type of image, whereas most geometric patterns can more economically be encoded by a vector format (= prepattern) [673, 726].

3) The number of pattern elements is size-dependent. Whereas coordinate-system models are designed to ensure pattern constancy regardless of pattern size, prepattern mechanisms inherently lack this ability [558, 621], though ad hoc amendments can be added to enable them to do so [24, 352, 665, 670, 671]. Since size can usually be altered easily, the demonstration of size-dependence in a given system can serve as a convenient operational criterion for ruling out the sole involvement of positional information mechanisms. It does not prove a prepattern mechanism, however, since other types of mechanisms, e.g. Darwinian ones, are also size-dependent. Because absolute size is a function of both cell size and cell number, it is possible to vary cell size (e.g. by polyploidy) and cell number (e.g. by starvation) separately to observe whether the pattern responds to either or both of these factors. Structures which vary in number as a function of organ size, cell size, or cell number include: zebra stripes [32, 33], Drosophila bristles [367, 758], Hydra tentacles [64], melanophores and ciliated epidermal cells in Bombina (a frog) [211, 212], whorls of fruiting bodies in...
Polyphondylum (a cellular slime mold) [809], wing veins in Ephesia (a moth) [559], pigment stripes in alligators [620], and ocular dominance columns [894] and lateral line primordia [984] in Xenopus. Notable patterns that do not change with cell size are the number of somites in Xenopus [133, 135, 346] and the number of 'fiz' stripes in Drosophila embryos [863].

**Hybrid Models**

The greatest strength of positional information mechanisms is their regulative ability; their greatest weakness is the amount of information processing they require [134, 1008]. Prepattern mechanisms have a complementary strength and weakness. It was inevitable, therefore, that hybrid mechanisms would be proposed which exploit the best features of the two types of models [1007]. Examples include:

**The Gradient/Reaction-Diffusion 'Superposition' Model [573]**

In many periodic patterns the structures are similar but not identical. For example, your fingers resemble one another but differ in length. Finger positions could be designated by a reaction-diffusion prepattern, with each finger growing to a different length based upon a gradient along the distal edge of the palm [134, 138, 1008]. Hybrid models of this kind are economical because (1) positional information need only be interpreted by a few rudiments (thereby minimizing information processing), and (2) positional signals need only be accurate enough to distinguish the rudiments (thereby minimizing the demands on the signal-to-noise ratio). The slight differences among human fingers would not require any interpretation of positional information per se since they could arise directly from a gradient in tissue growth (cf. Child [110], Huxley [407], and Thompson [882]), but there are many instances where one member of an anatomical series is greatly exaggerated, e.g. the wing strut of the pterodactyl (an enormous fourth digit), elephant tusks (enlarged incisors), and the 'sabers' (canines) of saber-toothed tigers. Such cases have been marshaled as evidence for a 'Principle of Non-equivalence' [508, 515, 1000], which postulates that (1) all cells have unique positional values, and (2) the ability to change the interpretation of those values genetically allows the independent evolution of formerly identical structures. Hybrid models, while not violating this principle, demand that any changes in the number or arrangement of structures must be explained otherwise—namely, in terms of the prepattern portion of the mechanism.

**The Gradient/Reaction-Diffusion 'Tuning' Model**

The network of genetic interactions that governs the expression of Drosophila's pair-rule segmentation genes appears to be highly complex—involving both positive and negative signals, reciprocal and nonreciprocal interactions, redundant functions, and stripe-specific sets of enhancer elements [5, 47, 98, 99, 350, 397, 410, 413, 675, 774]. However, Lacalli and Harrison [479] have argued that some of the complexity may be illusory: pair-rule gene products may be participating in reaction-diffusion mechanisms whose parameters (e.g. diffusion rates) are merely 'tuned' by gap gene products. The notion that diffusion rates are important for striping is supported by the polarized release of pair-rule gene transcripts from the apical ends of syncytial blastoderm nuclei, where the translated proteins would experience markedly greater diffusional impedances than gap gene proteins, which are not so confined [165].

**The Progress-Zone/Oscillator Model**

A peculiar feature of the development of the chick wing is that the time required for the specification of each bone rudiment within the progress zone is uniform despite a huge range in eventual lengths (e.g. wrist elements vs. humerus) [513, 514, 542, 867, 868]. The surprising implication is that the limb skeleton may begin as a periodic pattern of identical elements that diverge in size through subsequent growth. Wolpert and Stein [1011] have devised a hybrid model (cf. Meinhardt [574]) where oscillating chemical reactions within the progress zone produce a periodic prepattern of concentration peaks as they leave the zone. If each peak becomes a bone, then the different shapes and sizes of the bones could be controlled by the duration of time spent in the zone, as in the original Progress Zone Model. Thus, the rudiments would be created by a prepattern, with positional information steering them into different fates, as in the Superposition Model above. Analogous cases where a meristematic growth zone undergoes cyclic changes—leaving a periodic pattern in its wake [949]—include: tree rings [347] (alternating xylem and phloem), Xenopus tailbud somites [133], terminal segments in short-germ insects [244, 755, 757, 770], mouse molar teeth [538], barred feathers [294, 601, 641, 917], agouti hairs [294], and the spiral series of chambers in nautilus shells [132, 882, 935].
The gross anatomy of the vertebrate body is constructed by branched induction. The sequential induction model [72] can be illustrated by a model for the induction of the neural tube. In this model, the neural tube is induced by a gradient of inducing activity that is highest at the center and decreases gradually towards the periphery. The neural tube is then divided into a series of segments, each of which is induced by a different set of inducing factors. The sequential induction model is supported by experimental evidence, such as the induction of the neural tube by the notochord and the induction of the eye by the optic cup. The sequential induction model is also consistent with the idea that the neural tube is induced by a series of sequential waves of gene expression, each of which induces the expression of a different set of genes. The sequential induction model is an important concept in the study of neural development. It is also relevant to the study of the development of other organs, such as the limbs, where the induction of each limb bud is sequential and involves the induction of a series of different sets of genes.
ple - by the optic cup inducing the lens and then the lens inducing the cornea [341, 644, 807]. Comparable cascades may establish the states of individual cells within tissue monolayers. The clearest demonstration is the development of the Drosophila retina, where the inductive cascade within each cluster of 8 photoreceptors begins with photoreceptor R8 and ends with R7 [340]. Two different 'recruitment' scenarios are consistent with the available data [42, 986]: (1) The inductions might only involve one inducing cell at a time, as in the scheme (→* denotes an induction) R8→R2, R2→[R3 and, later, R1], R8→R5, R5→[R4 and, later, R6], and R8→R7 [28]; or (2) a combination of signals from several adjacent cells may be required, according to rules such as 'IF you are next to R1 and R6 and R8, THEN become an R7' [741, 890, 892, 893], and this 'epigenetic combinatorial code' [713] could extend to other ommatidial cell types (pigment cells, cone cells, etc.) as well [91, 92, 715]. In either case, the inductions do not iterate: the R7 of one cluster does not induce another R8 as the 'seed crystal' for an adjacent cluster [496, 1012], even though the clusters do develop sequentially across the retina. In theory, iterative inductions can construct periodic patterns [105, 106, 580, 581]. For the array depicted in figure 4a, a unidirectionally transmitted signal instructs black cells to induce white neighbors, and vice versa. In two dimensions, either parallel or concentric stripes could arise, depending upon whether the original source of the signal is a line or a point.

In monolayer cultures of Dictyostelium amoebae, scattered 'pacemaker' cells emit regular pulses of the diffusible signaling molecule cyclic AMP, which induces nonpacemaker cells to move toward the signal source [171, 172, 600, 635, 707, 729]. The signal is relayed throughout the cell population because each cell emits its own pulse upon stimulation, and reverse-propagation of the wave (i.e. cells responding to reflected signals from outlying cells) is prevented by a transient refractory period. Curiously, stripes (either concentric or spiral) of alternately moving vs. stationary cells arise because the refractory period includes two phases: a burst of movement followed by a stationary period [231, 914, 915]. Thus, this alternating pattern is not caused by alternating signals (i.e. moving cells telling neighboring cells not to move and vice versa) but by one signal which leads to an automatic succession of two states within every cell. Such dual wavefronts have been termed 'primary' and 'secondary' because one causes the other but not vice versa [765, 1020].

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### Determination Wave Mechanisms

Fig. 4. Determination wave mechanisms. Determination wave models create patterns sequentially from one end of a cellular array to the other, utilizing a single 'determination wave' whose propagation mode (and cellular effect) depends upon the assumptions of the particular model. **a** Sequential Induction Model. Alternating signals ('become white' or 'become black') are relayed along the array, causing cells to adopt states unlike their neighbors. **b** Clock and Wavefront Model [146]. All cells synchronously oscillate between black and white states (period - Δt) until a wavefront (striped semicircle) reaches them, causing them to keep the state they happen to have at the time. **c** Inhibitory Field and Competence Wave Model [196]. A cell is only competent to become black when it resides in the wavefront (striped semicircle). Upon becoming black a cell immediately establishes an inhibitory field ('IF', black oval), within which no other cell can become black. **d** Conditional 'IF/THEN' rules implicit in the models depicted above.
The Clock-and-Wavefront Model

Many types of cellular oscillators are known [58, 124, 125, 291, 304, 305, 619, 664, 977], the most familiar being the mitotic cycle. Theoretical mechanisms that explicitly employ such oscillators in patterning include the Phase-Shift Model of Goodwin and Cohen [124, 144, 310], the Clock-and-Wavefront Model of Cooke and Zeeman [134, 146, 1020], and the previously discussed Progress-Zone/Oscillator Model of Wolpert and Stein [1011]. In all of these models, there are two essential components: (1) a two-state 'clock' which oscillates synchronously (or nearly so) in all cells, and (2) a unidirectionally traveling 'wavefront' which is capable of stopping (or being modulated by) the cellular clock. In the Progress-Zone/Oscillator Model, the wavefront is associated with a terminal growth zone, whereas the other two models can operate within static (nongrowing) arrays. If the cells within an array are synchronously oscillating between two states, then a wavefront that stops the oscillations will leave behind a periodic pattern of alternating states. In figure 4b, the wavefront travels at a rate of two cells per oscillation cycle. A faster rate of travel would yield more than one black or white cell in each 'bandwidth' of the final pattern, and the widths of the black vs. the white bands could be made unequal by supposing that the cells spend unequal amounts of time in the two states during each cycle.

Although the wavefront could be propagated by a relayed signal as in the sequential induction mechanism, Cooke and Zeeman [146] present an intriguing alternative: in addition to its oscillator clock, each cell might have what is tantamount to an alarm clock [cf. refs. 12 and 801]. If the array is spanned by a morphogen gradient, and each cell schedules its alarm clock for a time corresponding to its morphogen concentration, then the alarm clocks could later ring in sequence from one end of the array to the other (causing the oscillator clocks to stop and resulting in a periodic pattern) without any intercellular communication. Whereas a 'relay wave' can theoretically be blocked surgically (e.g. by removing a cell or inserting a barrier, as one might stop a chain of falling dominos), a 'schedule wave' cannot [135, 170, 370, 496, 524, 958, 1012]. *Xenopus* somites, which arise sequentially in the typical vertebrate manner, behave according to the schedule wave scenario [143, 215, 679]: (1) the wave cannot be physically halted (no matter how early the operation is performed) [170], and (2) somite number is independent of both body size and cell number (as expected for a gradient mechanism) [133, 135, 224, 346]. In chick embryos, heat shocks cause spatially periodic somite abnormalities that are correlated with the duration of the cell cycle [696, 697], suggesting that somitic oscillators may be mitotically coupled [836]. Other systems where the cell cycle may gate cells into different pathways (cf. Reinert and Holtzer [719]) are *Dictyostelium* (where amoebae become prestalk vs. prespore based upon their cell-cycle phase at the commencement of aggregation [297]), the chick limb bud (where the proximodistal skeletal elements arise at a rate of one element per cell cycle [513]), and the ferret brain (where a cell's cell-cycle phase causes it to migrate to a particular layer of the neocortex [564]). Slack [795] has outlined a hybrid 'Clock-and-Wavefront/Gradient' Model, in which cells progress through an entire 'clockface' of scalar states per cycle (instead of oscillating between only two states), and the effect of the wavefront is to stop each cell's clock, resulting in a sawtooth series of gradients.

Inhibitory Field Models

Since antiquity, gardeners have known that a plant's apical meristem prevents nearby axillary buds from forming other apices. Surgical experiments by Child [110] and others [408, 731] circa 1910–1930 showed that many animal organs (e.g. a hydra head or a newt limb bud) can similarly prevent organs of the same type from arising in their immediate vicinity. Moreover, Child used this concept of an 'inhibitory field' to explain a periodic pattern: he argued that colonial hydroids acquire a regular spacing of hydranths along the stolon because a new bud sprouts whenever the tip grows beyond the inhibitory field of the previous hydranth [109] (cf. Pickett [683, 686]). Sir Vincent Wigglesworth was the first (in 1940) to apply the inhibitory field idea to the patterning of structures within an organism [963, 964]. He studied the positions of abdominal bristles in a hemipteran insect. Like the hairs on a human forearm, these bristles are spaced fairly regularly but are otherwise randomly arranged. Wigglesworth found that the new bristles at each successive molt arise in the largest gaps within the previous pattern. He conjectured that bristle cells consume a diffusible substance needed for bristle development, so that the first cells able to form new bristles would be those that are furthest from pre-existing sites.

In contrast to Wigglesworth's imaginary factor, which would act as a positive regulator, most subsequent 'lateral inhibition' [280, 579, 660, 788, 839] models postulate inhibitor molecules that would function as negative regulators. Although the production of an inhibitor is formally equivalent to
<table>
<thead>
<tr>
<th>Feature</th>
<th>Anaabaena heterocysts</th>
<th>Drosophila bristles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Specialized (S') cell</td>
<td>Heterocyst (nitrogen-fixing, non-mitotic) [357, 847, 913, 992]</td>
<td>Bristle mother cell, which divides to produce a 4-cell (mesoautotrophic) or 8-cell (autotrophic) bristle organ [325, 431]</td>
</tr>
<tr>
<td>Background (B') cells</td>
<td>Vegetative cells (photosynthesizing, mitotic) [814, 991]</td>
<td>Epidermal cells (nonsensory) [688]</td>
</tr>
<tr>
<td>Dimensions of the pattern</td>
<td>One-dimensional (filamentous) [814]</td>
<td>Two-dimensional (monolayer) [688]</td>
</tr>
<tr>
<td>Arrangement of S cells</td>
<td>Evenly spaced [995]</td>
<td>Some bristles are evenly ('isotropically') spaced; others are arranged in rows or aperiodic 'constellation' patterns (fig. 5) [369]</td>
</tr>
<tr>
<td>Number of B cells in each S cell interval</td>
<td>Approximately 10 [596, 967]</td>
<td>Approximately 5–10 [555, 367]</td>
</tr>
<tr>
<td>Frequency of 'incipient doublets' (pairs of S cells that commence development closer than a normal interval)</td>
<td>4% [967]</td>
<td>Occasional [400]</td>
</tr>
<tr>
<td>Outcome of 'sibling rivalry' between members of the incipient doublet</td>
<td>One of the 'probriocysts' completes differentiation; the other regresses to a vegetative state [596, 967, 968]</td>
<td>One of the 'probriocysts' completes differentiation; the other presumably regresses to an epidermal state [400]</td>
</tr>
<tr>
<td>Method used to artificially suppress S cell development</td>
<td>Puncture of probriocyst [596] or breakage of filament [967, 990]</td>
<td>Construction of genetic mosaic whose mutant tissue cannot form bristles [835]</td>
</tr>
<tr>
<td>Effect of suppressing S cell development</td>
<td>Nearby vegetative cell commences heterocyst differentiation [596]</td>
<td>Nearby epidermal cell commences bristle differentiation [831–834]</td>
</tr>
<tr>
<td>Genes whose mutant alleles (or scabs) reduce S intervals</td>
<td>hsr [83], Multiple 7 [969], Terminal 7/4 [969]</td>
<td>Delta, Enhancer-of-split, Noch, polechaetoid, scabrous, shaggy, shibbere [103, 369, 788]</td>
</tr>
<tr>
<td>Chemicals or other conditions which increase or decrease S intervals</td>
<td>Increase spacing: fixed-nitrogen compounds [913, 995] Decrease spacing: 7-azauraytophan [955], rifampicin [995]</td>
<td>Increase spacing: triploidy [367] Decrease spacing: haploidy [738]</td>
</tr>
<tr>
<td>Putative inhibitor</td>
<td>Glutamine or glutamine derivative [83, 993] (but cf. Wolk [994])</td>
<td>Product of the Delta [365] (or scabrous [27]) gene</td>
</tr>
<tr>
<td>Putative mode of inhibition</td>
<td>Diffusion [83]</td>
<td>Cell contact [788] (or diffusion [27, 598])</td>
</tr>
<tr>
<td>Additional mechanisms supposedly involved in the patterning of S cells</td>
<td>Cell-lineage rule; each vegetative mitosis is asymmetric, and only the smaller daughter can become a heterocyst [594, 967]</td>
<td>Equivalence groups: only cells in certain areas can become bristles [369, 789]</td>
</tr>
</tbody>
</table>

the consumption of an inducer in terms of the types of patterns that are formed [114], a failure of the mechanism (e.g. caused by mutations) would lead to different 'default' outcomes: either complete absence or ubiquitous differentiation of the pattern elements, respectively. Both lateral inhibition and 'medium-range' distances (of several cell diameters) activation [580] are thought to control vulval cell fate in Caenorhabditis elegans [327] and the expression of 'segment-polarity' genes in Drosophila [376, 554]. Inhibitory fields need not be mediated by diffusible chemicals: they could theoretically be caused by mechanical forces [324, 325], by direct cell contact, or by cellular extensions (e.g. filopodia) [372]. Table 2 compares two systems where evidence exists about the molecular basis of the inhibitory agent: the spacing of heterocysts (nitrogen-fixing cells) in Anaabaena (a cyanobacterium) and the spacing of bristles in Drosophila. The various bristle patterns that are supposedly created by inhibitory fields are depicted in figure 5.

The type of pattern studied by Wigglesworth can theoretically be generated de novo – rather than gradually through growth – by having naive cells randomly choose to adopt a particular state and immediately prevent their neighbors from doing so [114]. When the array becomes saturated with inhibitory fields, then the nearest-neighbor distances between the sites will range from 1.0 to 2.0 inhibitory-field radii. To produce a more uniform spacing of pattern elements, a 'competence wave' has been included in several models [116, 579, 659, 720, 995]. The term refers only to the effect of the wavefront – not its mode of travel, which could theoretically be via either a relay or a schedule. The idea, as illustrated in figure 4c, is that a cell is only 'competent' [925] to become black when it is within the wavefront. Upon becoming black, this cell immediately establishes an inhibitory field, so that the next cell that can become black will lie just outside the field. In the final pattern all of the nearest-neighbor distances would equal 1.0 inhibitory-field radius.

Inhibitory fields have been invoked not only to account for the patterning of heterocysts [596, 967, 968] and insect bristles [114, 115, 369, 370, 425, 487, 720, 831] as discussed above, but also insect neuroblasts [178, 180] (where laser-ablation of an incipient neuroblast leads to its replacement by an adjacent ectodermal cell) and pheromone glands (whose pores develop only at the vertices of polygonal epidermal cells) [813]; axo-lotl dermal glands (where nearest-neighbor distance is proportional to gland size) [385]; shark scales [718]; shark [717], reptile [657, 658], and mammal [89] teeth; sheep hairs [114, 117]; and leaf stomata [84, 85, 471–473, 752, 753].
Models that include a competence wave (either linear or annular) in conjunction with inhibitory fields can produce amazingly precise patterns, including hexagonal lattices (bird feathers [158, 159, 196, 524], fly ommatidia [369], and, possibly, vertebrate photoreceptors [483, 965]) and arithmetic spirals (plant leaf primordia [593, 721, 768, 818, 883, 884, 936, 1016]) (fig. 6). The inhibitory-field/competence-wave mechanism thus provides a plausible solution to the 50-star portion of the ‘American Flag Problem’ posed in the Introduction.

Specific Inhibitor Model
Hierarchies of qualitatively different inhibitors could create an orderly series of different cell states [137–140, 580] – an idea first proposed by Rose [730–732]. He argued that cells having state ‘A’ could produce inhibitor ‘A’, which would diffuse locally, allowing cells outside the inhibitory field to assume state ‘B’ (the next state in the hierarchy), produce inhibitor ‘B’, and the process would thus continue until the tissue becomes partitioned into a series of stripes (A, B, etc.) whose relative widths could be controlled by the diffusion parameters of the inhibitor molecules. Since this model could easily produce alternating red and white bands, it shows how the 13-stripe portion of the ‘American Flag Problem’ could be solved without a coordinate system. Embellishing upon this simple model, Green and Cooke [321] have proposed a ‘Serial Diversion’ Model, which, interestingly, can scale a pattern in proportion to its size. Instead of inhibitory fields, they use a reaction-diffusion mechanism. A competence wave enables cells to make a succession of different activator and inhibitor molecules (cf. Meinhardt [373]), which diffuse to the boundaries of the tissue. Because their diffusion is confined, their concentrations will increase if the area decreases, allowing them to reach the ‘diversion threshold’ for the next activator and inhibitor at an earlier time, hence causing the competence wave to compress the entire pattern to fit the area. Bard [33] had earlier proposed a similar hybrid mechanism to explain how parallel stripes develop in zebras.

Specific Activator Model
The opposite of a Specific Inhibitor Model would be a ‘Specific Activator Model’, and this type of mechanism was advocated by Weiss [952]. He imagined that the confrontation of two tissues A and B could cause reactions at their interface, leading to the creation of an intermediate layer C, which could then react with both A and B, thus continuing to stratify the tissue ad libitum. Similar arguments for sequential interfacial inductions

Fig. 5. Drosophila bristle patterns, supposedly created by inhibitory fields. On the fly surface there are roughly 3,000 bristles, a few dozen of which are much larger than the rest. These ‘macrorhachae’s (M’s) function as extended mechanoreceptors (like cat whiskers) and tend to occupy highly stereotyped aperiodic (‘constellation’) patterns. The smaller ‘microrhachae’s (m’s) tend to be evenly spaced and aligned to varying degrees. Most bristles on the dorsal abdomen are ‘isotropically’ spaced (i.e. no axial alignments; as in the insect studied by Wigglesworth [963]), while those on the thorax and legs are aligned in uniaxial rows, and those in the compound eye occupy vertices in a triaxial lattice of ommatidia. Bristles can be so close that they actually touch (the ‘tandem’ array). Based upon various lines of evidence (including mutant phenotypes), all bristle cells are thought to possess inhibitory fields, with the region-specific differences in bristle arrangements being due to temporal and spatial restrictions on exactly which epidermal cells are ‘competent’ to become bristle cells [365, 788]. From Hild [369].
have been advanced to account for progressive differentiation in sea urchins [161] and Xenopus [557, 800, 802], and for intercalations of cell states between stripes of segment-polarity gene expression in Drosophila [377, 393, 492, 554, 555]. Aspects of Weiss's model also resemble elements of Meinhardt and Gierer's scheme for the lateral activation of mutually exclusive states' [573, 580].

portion of the eye rudiment is diagrammed. Three successive stages (left to right) in the progress of the wavefront are illustrated. Though an uncommitted (white) cell must be in the competence wavefront zone in order to become an R8 (black) cell, its presence there is not sufficient. The cell must also reside outside the inhibitory fields (shaded ovals) of pre-existing R8 cells. Cells that satisfy both criteria become R8 cells and establish new inhibitory fields. Thus, a hexagonal lattice forms because each new row of R8 cells arises in the interstices of the inhibitory fields of the previous R8 row (after Held [369]). The sites in the original (leftmost) column, which function here as a 'seed crystal', could conceivably have been established by an earlier perpendicular wave traveling along that column (as in fig. 4c). This model was first proposed as an explanation for the development of hexagonal patterns of bird feathers, which also develop in a wave-like progression within discrete epidermal 'tracts' [196]. In other types of models for hexagonal patterning, the competence wave has been retained, but inhibitory fields have been replaced by either a reaction-diffusion [623] or a physical-force [351, 662, 681] mechanism. A competence-wave/chemotaxis strategy has actually been used in vitro to coax chemotactic bacteria to self-assemble into a lattice [82]. b. Cellular interactions supposedly occurring in a plant apical meristem. The leaves of many plants originate in a spiral pattern. Along the cylindrical stem created by the dome-shaped meristem, the spiral becomes a helix (not shown). Because the positions of new leaf primordia (small filled circles) can be altered by Surgically separating their presumptive sites from older adjacent primordia, it seems that older primordia possess inhibitory fields (shaded circles) [818, 936]. Furthermore, because primordia arise at a constant distance from the apex, there appears to be a competence zone (dark annulus) analogous to the morphogenetic wavefront in the Drosophila eye. However, in this case, successive ranks of cells move through the (stationary) wavefront zone (by centrifugal displacement away from the mitotically active apex), rather than the other way around, and the zone is circular instead of linear. Interestingly, mathematical simulations have shown that inhibitory fields, centrifugal growth, and a competence zone are sufficient to generate spiral arrangements - even without a 'seed crystal' to start the pattern [593, 721, 883]. Moreover, the simulations explain why successive primordia in so many plant species arise at the 'golden angle' of 137.5°. The explanation is purely astatic, as depicted here: given the relative overlap of the wavefront zone with the inhibitory fields of the three most recent primordia (No. 2, 3, and 4), the next primordium (No. 1) must arise at approximately this angle relative to its immediate predecessor (after Richter [721] and Wardlaw [936]). Hexagonal lattices and arithmetic spirals are also found in the realm of animal behavior, e.g., honeycomb and orb webs, and there too, only a few cue-driven behaviors seem to be involved [31, 194, 195, 682, 716, 922, 985].

Fig. 6. Precise patterns, presumably created by inhibitory fields plus competence waves. Surprisingly, researchers working with geometrically different patterns in organisms as different as flies, birds, and plants have independently devised similar models to explain the precise patterning of ommatidia, feathers, and leaves respectively. a. Cellular interactions which supposedly lead to the development of a hexagonal lattice of 'R8' photoreceptor cells in the Drosophila eye [27]. A 'morphogenetic wavefront' [714] (dark vertical bar) sweeps across the array, endowing cells with the 'competence' to become R8 cells. Cell columns are numbered so as to provide reference landmarks. Only a small
Chapter 4:
Darwinian Mechanisms

'Sibling rivalry', as exemplified by the contests between Anabaena proheterocysts or Drosophila bristle cells (table 2), is merely one form of intercellular competition—a phenomenon observed commonly in embryos [429, 588, 810]. In 1881, 22 years after Darwin's 'Origin of Species', a book entitled 'Der Kampf der Theile im Organismus' [738] ('The Struggle of Parts within the Organism') was published which asserted that embryos develop in a manner analogous to how species evolve—by selection among competing variants (in this case, cells) for an adaptive outcome (a functional anatomy) [653, 655]. The author was Wilhelm Roux, a student of Haeckel's and the founder of Entwicklungsmechanik (the science of developmental mechanics) [30, 602, 739]. Since then, much evidence has been adduced for selective strategies in development, and Gerald Edelman's recent book 'Neural Darwinism' [202] has rekindled interest in such mechanisms, especially as they pertain to the nervous system.

Cell Death Models

One of the most perplexing events in development is cell death [763]. Why should an embryo invest its precious energy in creating new cells, only to destroy them? Aside from serving a sculpting function during morphogenesis and metamorphosis [378, 530, 763], there appear to be several pattern-related reasons for why cells die:

(1) Being born into a 'pruned' lineage tree. Studies of cell death mutants in C. elegans have led to the conclusion that certain cells are autonomously 'programmed' to die [213]. Given the evolutionary conservatism of many of the lineage motifs [841, 842], such retroactive alterations may be genetically simpler than the redesigning of entire pathways. Comparable cell deaths are observed among bristle and scale lineages in insects [485], as well as among identified neuroblasts [531], perhaps for the same reason.

(2) Residing in a sagging gradient. According to hybrid 'gradient/reaction-diffusion' models, a constant number of structures can be generated within a field of cells if the size of the array is tightly controlled [134, 136, 138, 1008], and that control could be mediated by the slope of the gradient. Lawrence [490] has suggested that when the slope is too shallow, one way to steepen it is to contract the cell array by interspersed cell death, as does indeed occur in mutant Drosophila embryos whose segments are wider than normal [546] (cf. Klingensmith et al. [465]).

(3) Waiting too long to select a state. On average, 2 or 3 cells per ommatidium die during the construction of the Drosophila retina, and mutations are known which simultaneously rescue the cells and distort the lattice [987]. The implication is that the extra cells normally die because they 'fail to establish contacts appropriate to a specific cell type', and that their elimination is necessary to 'tighten' the lattice [91]. The existence of such a 'clean-up' mechanism may explain why so many genes can mutate to give reduced-eye phenotypes [246, 523]: slight timing errors [92] could easily lead to massive cell death.

(4) Matching or mismatching. There are two major vertebrate systems that employ the bipartite strategy of (1) creating an excess of cells, and then (2) selecting a subset thereof: the immune system and the nervous system. In the immune system, antibody diversity is generated by gene rearrangements within lymphocyte progenitors [197]. Those lymphocytes bearing 'anti-self' antibodies are then eliminated (negative selection) [742, 772, 923], while those bearing 'anti-invader' antibodies proliferate during each infection cycle (positive selection) [3, 974]. In the nervous system, excess neurons are produced in virtually every region, the competition for targets is intense, and many neurons die because they fail to find their targets [148, 654, 702]. For complex neural networks, the selective strategy is advantageous insofar as (1) it ensures numerical parity of the matching populations [112, 435, 436, 973], and (2) it can fine-tune the interconnections based upon functional criteria [74, 430, 701, 702]. The latter point is one of the central tenets in Edelman's Neural Darwinism Theory.

Given these examples where cell death is used as a patterning tool, the cell-death mechanism depicted in figure 7a is plausible, though no actual periodic patterns have been shown to originate in this manner thus far.

Most deaths in development are suicides rather than murders [113, 771]. A notable exception is the engulfment and murder of the C. elegans gonadal linker cell by the E.lp and E,rp killer cells. Laser ablation of the parent cell before E.lp and E,rp are born rescues the linker cell [213, 864].
(involving sequential influences among neighboring elements) is the centerpiece of Edelman's 'Topobiology' Theory [203-205], except in that case, the elements are cells on the one hand and inanimate extracellular substrates on the other. These two entities communicate via 'CAMs' [198, 199, 201] (cell adhesion molecules) on the former and 'SAMs' (substrate adhesion molecules) on the latter. In response to position-specific SAM signals, a cell can: (1) change its CAMs; (2) cause the SAMs to change; or (3) react by choosing any response in the accessible subset of its genetic or behavioral repertoire (accessibility is supposedly under CAM control), including moving (steered by the CAMs [200]) to another SAM environment, where a new dialog can begin [271]. Kauffman's and Edelman's models are ornate versions of 'Turing machines' [394, 906] and, as such, serve as testable incarnations of the 'Cybernetic Metaphor' for embryonic development (see chap. 7).

A state-change mechanism, such as the one diagrammed in figure 7b, may be involved in the patterning of lepidopteran wing scales. In several genera, the scales are arranged in parallel anteroposterior rows [1015], and within each row there are two types of scales: cover ('c') and basal ('b') scales which alternate (cbcbcbcb ... [1014]. In Pieris, each scale row develops from a single file of scale precursor cells (with no ordinary epidermal cells intervening within the file), which are also of two types: wide ('w') and narrow ('n'). However, the alternation of the w and n precursors is less perfect: 35% of the [w+b + [w+n] sequences have homotypic pairs [w+n + w+n], whereas only 4% of the total [cb + ccb + cbc] sequences include them [cc + cbc]. Cell death and cell movements have been ruled out as explanations for the fine-tuning process [1014], leaving state changes as a possibility (e.g. wnn → nwn → cbc, and wnn → wnw → cbc).

State-Change Models

There is evidence that morphallactic regulation may involve a 'ripple effect', wherein cells throughout a surgically reduced organ change their states in response to changes in the states of their neighbors until the original coordinate system is compressed to span a smaller cellular array [242]. In his 'Adaptive Antichaos' Model, Kauffman has devised rules for how the states of a cell's (or gene's) neighbors (or regulators) can influence its own choice of states; and by setting Darwinian criteria for certain types of mismatches, he has shown that Boolean networks of such elements can develop stable, orderly patterns [447]. A similar 'living network' scenario...
Chapter 5: Rearrangement Mechanisms

Tissue movements figure prominently in the early development of most metazoans, and individual cells frequently rearrange or migrate [247, 452, 899]. A cell's state of determination or differentiation can theoretically cause it to assume a specific position relative to other cells (by propelling it in a particular direction, or by stimulating it to move until a certain condition is met) [947]. Clear-cut evidence for such a mechanism has recently been found for identified motoneurons in embryonic zebrafish: after they were transplanted to new positions, the somata of many primary motoneurons moved back to their original positions [209].

Adhesion Models

Many viruses [10, 290, 516] and subcellular structures [10, 329, 412, 461, 921, 952, 953] are capable of 'self-assembly'. As exemplified by the Watson-Crick pairing of nucleotide bases, the process usually relies upon a 'jigsaw puzzle' fitting together of complementary binding sites on the participating monomers, which yields a configuration of minimum free energy. The idea of a self-assembling supracellular neuro-architecture, based upon the same jigsaw-puzzle metaphor, was crafted by Roger Sperry in a series of papers in the 1940s (it is traceable to Langley, 1895) [481, 700] and cogently summarized in 1963. It was Sperry's [808] conjecture that

... the cells and fibers of the brain and cord must carry some kind of individual identification tags, presumably cytochemical in nature, by which they are distinguished from another almost, in many regions, to the level of the single neuron; and further, that the growing fibers are extremely particular when it comes to establishing synaptic connections, each axon linking only with certain neurons to which it becomes selectively attached by specific chemical affinities.

A mechanism of this kind operates in the central nervous systems of insects, where different axon pathways express different surface antigens [43, 301]. Based upon ablation experiments, surface-labeling studies using monoclonal antibodies, and other lines of evidence, Goodman et al. [302] proposed a 'Labelled Pathways Model', which contends that neuronal pathfinding in the insect CNS is accomplished by qualitatively different labels that are 'read' by the growth cones of migrating axons.

Sperry's Chemoaffinity Model [314] has been eliminated as a viable explanation for the afferent connections between retinal axons and tectal neurons in the amphibian visual system. One item of counterevidence [702] is the ability of the 'projection pattern' (i.e. the pattern of retinotectal linkages) to be compressed or expanded to accommodate the relative sizes of the two organs when they are altered surgically [192, 193]. Such adjustments should be impossible if specific retinal axons are programmed to bind to specific tectal cells. In its place, many other models have been proposed, most focusing on single behaviors in the neural repertoire [128, 238, 239, 702]. An exception is the Synthetic Model of Fraser and Hunt [236, 237, 239], which has the virtues of being able to explain (1) minority as well as majority results from a number of experimental regimens, and (2) the formation of 'ocular dominance stripes'. These peculiar stripes develop when axons from an extra transplanted eye try to project onto the same tectal surface as axons from the in situ eye [128]. The Synthetic Model invokes three forces: (1) an adhesive 'C' force between retinal and tectal cells regardless of origin; (2) a position-specific retinotectal adhesive force, which is graded along the AP and DV axes; and (3) a repulsive 'R' force between retinal fibers (with the relative strengths assumed to be 'C > R > DV > AP' in Xenopus). If the R force depends upon correlated electrical activity in neighboring axons (i.e. 'fibers that fire together synapse together' [237]), then ocular dominance columns (which are uniformly about 200 μm wide [128]) are explicable because the 'xenophobic hatred' of the fibers exceeds their 'domestic distaste'. Neighboring retinal cells do indeed fire synchronously in mammals [582]. In the mammalian CNS, analogous naturally occurring stripes characterize intrinsic and descending projections of the neocortex, projections to the cerebellum, binocular retinal projections to the superior colliculus, and afferent projections from the lateral geniculate nucleus to the visual cortex [128, 131, 401], and activity-dependent rules may likewise explain the periodicities of these afferent segregations [113, 780, 1019]. Interestingly, these rules are formally the neural equivalents of the 'local-activation-lateral-inhibition'
rules in reaction-diffusion models for fingerprints and zebra stripes [589, 660, 872]. In an article entitled ‘Thinking about the brain’ [150] Crick once asserted that ‘there is something in embryology that likes stripes’, and the prevalence of such mechanisms in development may explain why.

Axial gradients of potential adhesion molecules have accordingly been found in the retinotectal systems of birds [902] and rats [129]. Elsewhere, adhesion gradients have been implicated in lepidopteran wings (proximal-distal axis) [631, 632], Drosophila wings (radial, as a function of distance from the wing margin) [725], and cotton bug tergites (anterior-posterior axis) [648], where they may provide haptotactic guidance cues [437, 900] for migrating axons [630], bristle cells [725], or myocytes [102, 972], respectively. Along the proximal-distal axis of the amphibian limb, graded adhesive differences have also been demonstrated, which may play a role in the communication of positional information [634], and an adhesivity gradient likewise appears to guide the migration of the amphibian pronephric duct [1017, 1018].

‘Sorting out’ is the process whereby cells of different types segregate after being mixed and aggregated in vitro. Many cell mixtures behave in this manner [225, 226, 898, 951], demonstrating that the state of a cell can indeed cause its position, at least under these artificial conditions. Sorting out per se does not prove an adhesive mechanism since differential chemotaxis – e.g. in response to an oxygen gradient caused by respiration of the aggregate – could also cause sorting out [541, 898]. Indeed, chemotactic sorting has been demonstrated for prostalk and prespore cells in Dictyostelium [846, 874]. Classic experiments by Townes and Holtfreter showed that amphibian cells from different germ layers can segregate into concentric layers that mimic their normal stratification in vivo [897] – suggesting that a ‘self-assembly’ strategy might be used during normal development (but in amphibia, neither gastrulation nor neurulation proceeds via cell sorting). Steinberg [819] conducted numerous similar experiments with a variety of different tissues and discovered an ‘Adhesive Hierarchy Rule’: if tissue A sorts to the inside of an aggregate when mixed with tissue B, and tissue C likewise sorts inside C, then A will sort inside C. Based upon this rule and other evidence, he proposed a ‘Differential Adhesion Model’ [820]. The model assumes that cell types vary in the degree of their adhesivity, and the ‘stickier’ the cell type (A > B > C) the more it will tend to form a tight and homogeneous clump [821, 823]. Investigations of neuronal pathfinding in the insect peripheral nervous system led Berlot and Goodman [57, 300] to propose a similar ‘Adhesive Hierarchy Model’.

Rearrangement Mechanisms

Our results suggest that the filopodia of the pioneer growth cones in the auditory and limb buds of the grasshopper embryo express an adhesive hierarchy, whereby the surfaces of neurons are preferentially adherent over the surfaces of the epithelial cells. (1) Given only the epithelium, the growth cones extend proximally along its surface, appearing to follow an epithelial adhesive gradient. (2) Given a choice in the periphery, however, of neurons versus epithelium, the filopodia preferentially adhere to the neural surfaces and thus guide the growth cones onto these neuronal cell bodies and axons. (3) Given a choice in the CNS of different axon bundles, certain neuronal surfaces appear to rank higher in the adhesive hierarchy than others; they invariably choose a particular axon bundle on which to extend, similar to the observation of selective fasciculation by central neurons that led to the labeled pathways hypothesis [57].

An ‘Adhesivity/Determination-Wave’ hybrid model has been proposed for somite formation in chick embryos based upon observed increases in the adhesivity of mesodermal cells before and after somite formation [35, 53, 108]. The idea is that paraxial mesoderm cells (perhaps gated into groups by the cell cycle [455]) increase their adhesivity when a competence wave reaches them, and clumps of them (nascent somites) consequently pinch off sequentially from the segmental plate.

Both homophilic and heterophilic adhesion molecules have been discovered [41, 176, 198, 282]. The patterning scheme in figure 8a employs heterophilic molecules at the poles of each cell to assemble an alternating chain of two cell types. Head-to-tail associations of this kind are critical for differentiation of Myxococcus bacteria [458]. A dramatic illustration of self-assembly in vitro is the spontaneous alignment of human fibroblasts [216] or keratinocytes [320] into ‘fingerprint-like’ [154] patterns containing loops, whorls, and triradii (cf. Nübler-Jung [647] and Seul et al. [777]).

Repulsion Models

The movements of chromatophore cells emigrating from amphibian neural crest explants in vitro (and the spreading behavior of such cells from transplants in vivo) led Twitty and his collaborators [908, 910, 911] (but cf. Erickson and Oliver [217]) to conclude that:

...the cells move in response to a mutual stimulation, or 'repulsion', probably mediated through the action of diffusible substances involved in or produced by the metabolism of the cells (themselves)... with the cells gradually spreading peripherally until they are spaced beyond the effective range of such mutual influences [908].
One of their experiments was especially informative [912]. Individual chromatophores were sucked into a capillary tube (full of coelomical fluid) and observed for several days. When there was only a single cell in the tube, it moved little. When two cells were placed together, they moved in opposite directions; and when there were three cells, they spread out until their spacing was uniform — all of which are behaviors consistent with Twitty's 'Mutual Repulsion Hypothesis' (fig. 8b).

In certain urodele amphibians (e.g. Taricha torosa) the pigment cells secondarily (i.e. following their initially uniform 'primary' distribution) aggregate into bands on the dorsal flank of the tadpole [904, 905]. The aggregation behavior also occurs in vitro and is attributable to direct cell contacts — via filopodia — which pull neighboring cells together [909]. In the frog Bombina, a gossamer network of cruciform filopodia develops from one class of (adepidermal) melanophores. Ellinger described the stages in its formation:

To this stage [6 days postfertilization], the melanophore distribution and orientation of cell extensions appeared to be random. This arrangement was altered markedly during the seventh day of development. A pattern became established in which the epidermal melanophore extensions frequently left the cell body at right angles to each other ... As differentiation proceeded, a 'grid' of interwoven epidermal melanophore processes became apparent within the integument [211].

Based upon the behavior of the filopodia at their intersections, Ellinger argued that the filopodial interactions cause the rearrangement of the lattice.

A similar causal role has been ascribed to filopodia on the scale cells of moth wings [633]. The scale cells in Manduca sexta are initially randomly arranged. They gradually align and become evenly spaced by rearranging, during which time the distances between neighboring scale cells are spanned by axially oriented filopodial extensions ('epidermal feet'). Whether the filopodia push or pull or simply serve as 'phone lines' for the communication of intercellular navigational signals is not known. In Drosophila melanogaster, mutations in several unlinked genes cause the misorientation of bristle cells, and the intervals that separate neighboring bristles (which are uniform in wild-type flies) are correlated with the relative orientations of the cells (bristles facing one another are far apart, while those facing away from one another are close together), suggesting a repulsive interaction via unidirectional filopodia [371]. (Bristle and scale cells are thought to be homologous [485, 917, 960]).

Fig. 8. Rearrangement mechanisms. Rearrangement models assume that the states of cells cause them to relocate. The starting configuration could be the outcome of an earlier round of patterning (via a different mechanism) or simply random. Whether intercellular communication is required depends upon the mode of rearrangement. a Adhesion Model. Cells have complementary binding sites on their surfaces, represented here as jigsaw-puzzle protrusions and indentations. The cells move until their binding sites are occupied. Depending upon the dimensions (1, 2, or 3) of the final array, its geometry (striped, checkerboard, etc.), and the numbers of cells in each homotypic domain, different types of binding (homophilic vs. heterophilic, quantitative vs. qualitative, etc.) might be needed. b Repulsion Model. One of the two cell types (black) is assumed to repel cells of its own kind. The mechanism of repulsion could be electrical (as depicted here, though galvanotaxis seems to play little role in development [282]), mechanical (via filopodial extensions like growth cones [372]), or chemical (via chemorepellent molecules [912]). The model is more easily visualized for a two-dimensional array (where cells can jostle within a fluid monolayer) than for the one-dimensional situation depicted here (where cells must move out of alignment in order to rearrange, and a separate mechanism, not shown, must be invoked to realign them). c Interdigitization Model. Different cell types that are originally segregated into separate files, merge into a single file. Here the interdigitization is accomplished by choreographed rotations of heterotypic pairs. d Conditional 'IF/THEN' rules implicit in the models depicted above.
Growth cones of navigating axons (an extreme version of a filopodium) can also respond to inhibitory (repulsive?) signals from other axons [164, 434]. The possibility that attraction and repulsion utilize the same signaling mechanism is suggested by the phenotypes of *egl* mutants in *C. elegans*, where a normally attractive target becomes repellent [838].

**Interdigitation Models**

In addition to being able to migrate individually (e.g., neural crest cells) and jostle within a moving monolayer (e.g., mesodermal cells during archenteron invagination) [247, 452, 899], cells can apparently also 'dance', i.e. perform choreographed maneuvers with one or more partners. For example, as preclusters of photoreceptors emerge from the morphogenetic furrow in the developing *Drosophila* retina, the cluster cells are arranged in an arc-shaped single file, but shortly thereafter they close ranks and form a rosette [986]. Also in the same tissue layer, the four cells of each bristle organ align themselves before enveloping one another concentrically [91]. Neither of these rearrangements actually produces the larger periodic patterns to which the cell groups belong. However, there is evidence that some periodic patterns do form via 'minuets' wherein the cellular partners pivot and interdigitate (fig. 8c). Thus, there is a row of bristles (row 8) on the tarsus of Hawaiian *Drosophila* species where the bristles simultaneously alternate with 'bracts' in a single file [367]. Bracts are uninucleated cuticular protuberances, which develop from epidermal cells that have been induced by the bristle-cell complex [370]. Based upon what is known about tarsal cell lineage in *D. melanogaster* [366, 493], it is likely that the bristle and bract cells originate lateral to each other and subsequently interdigitate, perhaps by the pivoting of bristle-bract cell pairs. Pivoting may likewise occur in the wings of certain lepidopteran species, where the precursor cells for the basal scales apparently insert themselves (at regular intervals) into homogeneous rows of cover-scale precursor cells [862, 1014]. Quarter pirouettes (90° rotations) have been observed in two other systems:

1. The hexagonal array of photoreceptors in *D. melanogaster*. Photoreceptor clusters in the ventral half of the eye swivel 90° clockwise and those in the dorsal half swivel 90° counterclockwise, so that a new (somewhat zigzag) plane of mirror symmetry is established at the equator [893].

Rearrangement Mechanisms

(2) Blocks of myotomal cells in *Xenopus*. Somites swivel 90° clockwise on the right side of the body and 90° counterclockwise on the left side [346]. Comparable rotations occur in chick embryos, though in that case they happen after the somites have segregated from the paraxial mesoderm [52]. (In *Xenopus* they occur during segmentation.)

In neither the *Drosophila* eye nor the vertebrate spine do the rotations cause the periodicity of the patterns that they modify. The rotations may serve an optical function in *Drosophila*. (Other insect eyes are highly stratified along the dorsoventral axis [296].) In *Xenopus* and chick embryos, the purpose of the pirouettes is less apparent. (Somites in most vertebrates do not rotate [135].)

**Chemotaxis Models**

The chemotaxis models of Oster, Murray, and others [620, 622, 663] behave in a manner that resembles the aggregation of *Dictyostelium* cells [172], except that the diffusible signal is neither pulsed nor delayed: all cells are assumed to emit a chemoattractant continuously. If the cells are initially scattered within a linear stripe, then the stripe will automatically dissolve into a series of clusters, each of which becomes more attractive as it forms, depleting the surrounding areas of motile cells. Such models are formally equivalent to reaction-diffusion schemes insofar as they also rely upon local autocatalysis and lateral inhibition [663]. A developing system where a narrow stripe actually does fragment into separate islands of cells (perhaps by this sort of mechanism) is the supraorbital lateral line primordium of *Xenopus* [983, 984], and a peculiar dispersal of tissue fragments has been described for pigment cell clusters in the fish *Blechnius pohli* [901]. In accordance with the predictions of this sort of model, a suspension of chemotactic bacteria can indeed coalesce into a precise lattice of clumps if the stimulus that initiates the chemotaxis spreads centrifugally (like a competence wave) through the array [52].
Chapter 6: 
Cell-Lineage Mechanisms

Until about 1900 it was customary to characterize embryos as 'mosaic' or 'regulative', based upon whether isolated blastomeres form only their normal portion of the anatomy (mosaic) or a larger portion (regulative) [583, 602, 998]. Gradually, the value of the distinction waned, as it became clear that the outcomes of such experiments depended critically upon the stage at which they were performed [160, 408, 945]. However, a version of the dichotomy has persisted until only recently – namely, the notion that embryos with 'determinate cleavage' (precise sequences of cleavage planes) assign cellular fates by 'cytoplasmic determinants' (substances that are asymmetrically partitioned to daughter blastomeres), whereas those with indeterminate cleavage assign fates via cellular interactions [160, 161, 602, 975]. In the last few years, the conceptual wall separating these two categories has begun to crumble, thanks mainly to nematodes and sea urchins, where determinate cleavage is combined with regulative ability. Ablations or transplantations in these species can alter cell fates, thus implicating cellular interactions (rather than the rigid cellular pedigrees) as the causative agents in the assignment of those fates [161, 459] (cf. Dohle [183]).

Citing such cases, which are heretical exceptions to the old stereotypes, Davidson has devised a more pluralistic classification scheme for incorporating the role of cell lineage in embryonic development [163], plus a set of basic models for the spatial regulation of histospecific genes [162] (cf. Holiday [388]).

In the wake of the old paradigm's demise, the cautionary lesson is that a ritualized cell lineage per se does not prove that cells are assigned their fates by means of that lineage [161, 459, 779, 829, 1000]: 'a determinate lineage may precisely locate cells, which then go on to adopt highly predictable fates because of extracellular cues' [244]. Only when relative cell positions are experimentally altered can the mechanism be ascertained. By contrast, it is easy to disprove a cell-lineage mechanism. For instance, by using a pigment mutation to genetically mark individual cells in the developing retina of Drosophila and then charting the locations of their descendants in the photoreceptor array, Ready et al. [714] were able to reject the hypothesis that each ommatidium is descended from a single mother cell. (They also showed that the equator of the eye is not established clonally.) Indeed, as a rule, organisms whose embryos manifest indeterminate cleavage do not construct their organs as clonal modules [151, 861].

Exceptions to this rule include certain miniature organs, e.g. insect sensilla [485] and leaf stomata [684, 816, 817]. These structures do develop according to strict pedigrees, though here too, some of the lineages may not be causal in assigning fates. Thus, each mechanosensory bristle in Drosophila is constructed from four cells – a shaft cell, a socket cell, a neuron, and a thecogen [451, 688] – all of which are descended from a single mother cell via two differentiative divisions [355]. The shaft and socket cells are sisters, as are the neuron and thecogen [91, 355], and the orientations of the mitotic spindles during these divisions are highly reproducible from one individual to the next [400]. Such rigid cell lineages are theoretically sufficient to assign fates, but in fact they can be bypassed: along the wing margin, where hundreds of bristle cells jostle into alignment, sister cells can contribute to separate bristles and adopt the same fate (e.g. socket cells) [355]. It is their relative positions that appear to determine their fates – a supposition which is supported by the changes in cell fate that accompany mutational alterations in bristle cell positions [29, 498].

Three types of possible lineage mechanisms are discussed below, along with various developing systems whose cell lineages conform to one or another of the schemes. Unfortunately, in most cases the critical experiments have not been performed to ascertain whether the lineage plays a causal role.

The Quantal Mitosis Model

The term 'quantal cell cycle' was introduced by Holtzer [390–392] to designate an asymmetric cell division in which the parent cell differs from at least one of its daughters, in contradistinction to a 'proliferative' cell cycle, where parent and daughters are identical. An example of a quantal cycle was mentioned for Anabaena in chapter 3 (table 2): every mitosis along the filament is asymmetric, yielding a small daughter that is capable of becoming a heterocyst, and a large daughter that is not [594, 967]. The asymmetric mitoses in this case, however, are not solely responsible for the
regular spacing of heterocysts, since inhibitory fields play a crucial role. Other periodic patterns where quantal mitoses are involved in the spacing of pattern elements include:

(1) The alternating pattern of hair cells ('trichoblasts') and ordinary epidermal cells in the root epidermis of various plant species [155, 184], including the monocotyledon Phleum pratense. The last division of the precursor cells in a plane perpendicular to the longitudinal axis of cell polarity gives rise to two daughter cells. The division is unequal, a smaller cell with denser cytoplasm being cut off toward the apical (distal) end, and a larger cell at the basal (proximal) end. The smaller cell (trichoblast) remains strongly meristematic and soon forms ... a root hair [62].

(2) The spacing pattern of gonial precursor cells in Volvox (a colonial green alga [462]). At the sixth cleavage in Volvox carteri, 'all 16 (or under suboptimum conditions only a portion) of the cells, which were derived from the two anterior tiers of the 16-cell embryo, divide unequally to yield a small-anterior/large-posterior pair of sister cells. The larger member of each pair becomes a gonial initial; the smaller, a somatic initial' [463]. Surgical experiments with a related species, Volvox obversus, have shown that size alone is not the determining factor [705]. The distances between the gonial initials are subsequently increased by further divisions of the intervening somatic cells and by several additional asymmetric mitoses of the gonial cells themselves.

In both of the above examples, the cellular array constitutes a 'tessellation' [288, 330] pattern, since it fills space using a single type of tiling unit. Here, the unit is a clone containing one pattern element plus one or more 'background' cells (which are generated by the same compartmentalized mechanism that produces the pattern element itself). Other systems where pattern elements appear to be positioned by a clonal tessellating mechanism include: wing scales in certain butterflies [487, 948] and leaf stomata in certain plant species [84, 152, 471, 751, 753]. In the hypothetical illustration in figure 9a, every cell (at a definite time in development) undergoes a mitosis which is both asymmetric (the daughters are different colors) and polarized (black is on the left and white is on the right), resulting in a periodic pattern.

**The Stem Cell Model**

The stem-cell strategy is a variation on the quantal mitosis theme [1006]. In this case, only one cell undergoes asymmetric divisions, and it does so repeatedly (fig. 9b). Many organisms employ stem cells as pluripotent progenitors for a variety of terminally differentiated cell types [342, 534, 695]. In order for a stem-cell mechanism to directly generate a periodic pattern, it must create alternatingly different daughters. Such a process produces certain chains of cells in leech [779] and earthworm [849] embryos. Both of these annelids have determinate lineages in which "band-
lets' of ectodermal and mesodermal precursor cells arise from iterated mitoses of huge stem-cell 'teloblasts'. There are 5 teloblasts on each side of the midline, and 2 of them (N and Q) produce (n or q) bandlets within which every other 'blast cell' eventually manifests a different cell-lineage pattern. In the n bandlets of the leech, the alternate blast cells (n\textsubscript{i} and n\textsubscript{j}) also show a different staining intensity of a tracer dye when it is injected into the N teloblast, suggesting a difference in diffusibility through the cytoplasmic bridge to the parent teloblast when they are born [59, 60]. Bissen [59] incorporated this observation into a 'Flip-Flop Feedback Model', in which good diffusibility allows a blast-cell daughter to signal the teloblast to change its state (e.g. from 0 to 1) while at the next mitosis poor diffusibility would block the feedback signal and cause the teloblast to revert to its ground state, resulting in a flip-flop alternation of states: n\textsubscript{i}, n\textsubscript{j}, n\textsubscript{i}, etc. Interestingly, segment number — which is extremely precise in leech species — is apparently not controlled by cell lineage [778] (cf. Pfannenstiel [683]).

An organism whose lineage has been analyzed completely is the nematode C. elegans [843, 865]. Kimble [459] has shown that even the most complicated lineage trees can be reduced to 2 elemental components — a 'stem cell' motif and a 'symmetrical mitosis' motif — that have been embellished in 5 possible ways: (1) a new switch in cell fate at some point in the pedigree; (2) a polarity reversal where fates are conserved but transposed; (3) a duplication of a cell near the beginning of a tree; (4) a duplication of a cell near the end of a tree; or (5) an iteration of one of the previous alterations.

L-Systems and Fractal Geometry

In many developing systems, cells 'self differentiate' [343, 408], automatically passing through a sequence of states in which each state change (of gene expression or cellular activity) is elicited by the previous state: state A → state B → state C, etc. [19, 95, 106, 428, 533, 592, 811, 885, 945, 948]. By combining this notion of internally motivated state changes with a stem-cell style of meristemistic growth, Aristid Lindenmayer invented a class of mathematical models that has since come to be called 'L-systems' [518–520, 522]. Like 'Turing machines' [174, 394, 906], L-systems postulate transition rules of the form 'IF you are in state x, THEN adopt state y'. By allowing more than one cell to be a stem cell, an L-system can cause the main filament to form branches that continue to grow and change states independently. (Because multiple growth points require parallel computations, such L-systems resemble cellular automata [886].) Furthermore, because the transition rules can lead to recursive cycles, the branches can undergo further branching on a smaller scale, resulting in 'fractal' patterns [40, 426, 550], which possess the property of 'self-similarity' (i.e. their structure remains the same at different levels of scale). Many biological patterns (e.g. trees, fern leaves, and blood-vessel networks) are fractals and can be simulated in this manner [520, 521, 656, 699, 740] (cf. De Reffye et al. [167]).

The Cortical Inheritance Model

In C. elegans, mutations in the gene lin-17 cause asymmetric mitoses to become symmetric throughout various lineages [363, 844], implying that the assignment of cell fates may be controlled by 'switch genes' that are regulated in a cell-cycle-dependent manner [940]. Asymmetric mitoses are generally thought to result from qualitatively or quantitatively unequal allocations of gene-regulatory molecules to the two daughter cells [228, 363, 795, 797] (cf. Gober et al. [289]), though the control of mating-type interconversion in yeast (where each daughter inherits a different 'cassette' gene at a particular locus) shows that other mechanisms are also possible [375]. The regulatory molecules could either reside in the cytoplasm or the cortex [998]. Many examples of 'cytoplasmic determinants' are known [106, 549, 860]. By contrast, the eukaryotic cell membrane is usually viewed as a 'fluid mosaic' [281, 792], incapable of reliably partitioning regulatory signals because it cannot rigidly hold an array of molecules. However, epithelial cell polarities challenge this notion [247, 526, 528, 529]. The anisotropically pigmented cortices of many fertilized eggs (e.g. the dark animal hemisphere in Xenopus) are a vivid demonstration of the ability of cell surfaces to maintain large discrete domains, and there is evidence that they can stably maintain microdomains as well [329].

Aside from the remarkably periodic patterns of cilia in protists [235, 804], there is at least one known case of a eukaryotic cell that generates a periodic pattern of molecules in its cortex. This is the Drosophila syncytial blastoderm, which is technically one cell despite its thousands of nuclei. It produces pair-rule stripes just beneath its cell membrane. The stripe widths and intervals do not depend upon nuclear spacing, since both
remain uniform in haplodiploid mosaics where nuclear sizes vary [492, 863]. Conceivably, comparable scaffolds of molecules could be assembled within any cell (not just an egg), and future division planes could partition the cortex in a coordinated fashion (fig. 9c). In that case, the descendant array would constitute a cellularized version of the ancestor cell's cortex. Evolutionary reversals in the polarities of nematode lineages have been explained using this same rationale: the assumption is that the progenitor cell has at least three domains (A, B, and C) that can be segregated in different sequences (ABC→A/BC→A/B/C, or ABC→AB/C→A/B/C) by eccentric placement of successive division planes [841].

Chapter 7:
The Computer Metaphor in Developmental Biology

Watching time-lapse movies of embryonic development can be fascinating, especially when blastomeres are cleaving or motile cells are 'feeling' their way along. Embryonic cells seem more like active robots than passive 'bricks' [196, 249, 752, 946]. The automatic nature of development has invited comparison with programmable computers ever since the latter were invented. Indeed, two of the founders of modern computing — Alan Turing and John von Neumann — speculated extensively on embryological problems (specifically the puzzles posed by morphogenesis [907] and self-reproduction [924], respectively). Historically, the computer metaphor [14, 15, 106, 268, 423, 482, 926, 1010] has provided some valuable heuristics ... plus, unfortunately, many vacuous clichés and muddled debates [308, 605, 962]. Part of the problem is that arguments can sink into a semantic quagmire over how to define developmental 'control' [952, 954, 955] or developmental 'program' [504, 645, 826, 828] (cf. Sternberg [840] for useful definitions of both terms in the context of nematode development). Given the illustration of so many developmental mechanisms in cybernetic formats in the previous chapters, plus a new conceptual framework for thinking about patterning, there is now a fresh opportunity to reexamine the metaphor. In this chapter, an attempt is made not only to tease out new insights, but also to begin raising the metaphor to the level of a testable theory (cf. Goodwin [308] and Huszagh [405]).

Local vs. Global Information

If anatomy is indeed built by cellular 'robots', then the engineering of development might best be seen from a cell's point of view [196, 336]. Whereas a cell can 'feel' and 'taste' its neighbors and 'smell' molecules that waft in from the distance, its sense of what is happening elsewhere is nec-
Chapter 7

essarily limited. Its predicament is like that of an ant walking on a billboard, trying to read the message by deducing each letter from its curvature, vertex angles, and crossbars [591]. Fortunately, development seems designed for a lilliputian perspective. Thus, for example, limb regeneration (in insects and vertebrates) obeys "Bateson's Rule" [977], which governs the arrangement and handedness of triplicated limbs that share a common shoulder or hip joint (fig. 10a). Such limbs arise spontaneously only rarely but can be easily produced by grafting operations [245] or by mutationally induced cell death [76, 283–285, 287, 749]. When two extra limbs bracket a normal limb, they have a handedness opposite to it. At the cellular level, Bateson's Rule is attributable to a 'Local Continuity Rule' [77, 510, 977] (inherent in the Polar Coordinate Model and related schemes), which requires that cells restore pattern continuity (by moving, dying, or intercalating new cells) whenever they are beside an inappropriate neighbor. Branched limbs and other regenerative outcomes may seem so illogical as the equation '1 + 1 = 3', but they are perfectly natural to the participating cells because no local rules are violated, and the same holds true for other abnormal anatomies (e.g., Siamese twins) that have new internal planes of mirror symmetry [240, 311, 494, 541, 748]. Indeed, many large-scale geometries and topologies can easily be reformulated mathematically in terms of small-scale behavioral rules [2, 361, 672].

Cellular perceptions may be as confined in time as they are in space. For instance, cells that participate in multiple inductions only acquire the competence to proceed to the next step if the previous step has been completed [282]. To the extent that development proceeds by a cascade of events at the cellular level, it is 'epigenetic' [341] – i.e., genes need not be directly involved (see below). Where genes do seem to play crucial roles is in (1) initiating such chain reactions [307] and (2) encoding states of cellular determination [442].

**Binary Codes and Boolean Logic**

Digital computers use a binary code for both memory storage and information processing [934], and there is mounting evidence that cells do likewise to a limited extent. Cells can react to simple cues such as 'divide', 'move', or 'become a neuron' [196] (cf. ritualized animal behaviors that are also triggered by simple cues [25, 328, 406, 535, 562, 607]). The IF/THEN rules that link stimuli to responses are binary insofar as a cell can either

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**Fig. 10. Local harmony vs. global discord.** This figure is intended to allow the reader to experience development from a 'cell's-eye view'. a Schematic diagram illustrating 'Bateson's Rule' (cf. Bateson's [46] fig. 153), as applied to the special case where a three appendages (represented by hands) develop from a left-limb blastema transplanted onto a right stump (black base) in salamanders. The two extra limbs (shaded) that grow out from the graft interface are always mirror images of the central limb (black), which retains its original handedness [245]. 'M1' and 'M2' indicate planes of mirror symmetry. The actual limbs that Bateson studied arose spontaneously and had both extra limbs on the same side of the original; cf. French [240]. Such regenerative behavior is attributable to 'Local Continuity Rule' [77, 510, 977], which ensures that all cells ultimately (after intercalary regeneration) reside next to the types of cells that would normally be their neighbors. Notice that thumb cells reside next to other thumb cells across plane M1, and 'little finger' cells confront one another across plane M2. Thus, although the anatomy at this gross level is bizarre, it violates no rules from the perspective of the individual cells. After Frensch et al. [245]. b 'Impossible Triangle' Illusion [609], upon which M.C. Escher based several of his famous lithographs [220, 879]. Here too, there is an obedience to rules of local connectedness, as each vertex is architecturally valid, but the whole triangle is topologically heretical because its mutually orthogonal edges should not be able to achieve closure. This comparison between the triplicated limb and the Impossible Triangle is more real than metaphorical, since our retinal cells gather information in much the same local manner as salamander limb cells, and it is only when the higher-order centers in our visual system attempt to compile the image fragments into a self-consistent mental object that difficulties arise [301, 352]. After Escher [220].
execute the behavior or ('0') not do so. The stepwise nature of differentiation in most organisms means that particular states constrain future choices, so that cells follow a branching pathway of binary decisions until they reach particular final states [341, 466, 796, 945, 948] (but cf. Roth [737]), at which point their 'potency' equals their 'prospective fate'. For example, the sequence of cellular decisions leading to a cholinergic motor neuron would be: 'a' commitment first to ectoderm rather than to mesoderm, then to nervous tissue rather than to skin, then to neuron rather than to glia, then to motor neuron rather than to sensory neuron, and finally to synthesis of acetylcholine rather than \( \gamma \)-aminobutyric acid' [830].

The idea that cells might encode their determined states as ordered series of their previous choices (e.g. a final state of '11001' based upon 5 earlier decisions) was first proposed by Stuart Kauffman [438, 442, 443, 449] (cf. Slack [794]). He argued that cells could 'remember' each decision by adopting either an ON or an OFF state for individual regulatory genes (e.g. a gene being transcribed vs. not being transcribed [272, 941]), and particular combinations of the 'memory gene' products would ultimately activate other ('structural') genes that would implement the expression of the final differentiated state (e.g. hemoglobin genes in an erythrocyte).

Kauffman's original hypothesis was based upon frequencies of 'transdetermination' exhibited by cells belonging to different parts of the Drosophila body [440]. Certain types of interorgan transformation (e.g. leg cells becoming wing cells) occur more frequently than others during long-term tissue culture. If the possible transformations (leg \( \rightarrow \) wing, leg \( \rightarrow \) eye, eye \( \rightarrow \) wing, etc.) are diagrammed as a network, then certain paths are favored over others. (Homeotic transformations in human epithelia also obey a 'Weighted Network Rule' [797].) Such biases are understandable if each binary register is controlled by a separate 'switch' gene, and mutations (or spontaneous epigenetic errors) are sufficiently rare that they typically affect only one gene at a time. For example, a code of 11111 could easily change to 10111 if the gene for the second register became defective, but two separate events would be needed to change 11111 to 10101, thereby making the first type of transformation (e.g. leg \( \rightarrow \) wing) more likely than the second (e.g. leg \( \rightarrow \) eye).

Combinatorial codes are efficient because they utilize a minimum number of genes to specify a maximum number of states, so evolution should have favored them [277, 443]. Genetic and molecular evidence implicates the involvement of a combinatorial code for differentiated states in Drosophila body segments and parasegments [7, 87, 98, 411, 414, 424, 492, 745, 853] (but cf. rebuttals [269, 298, 490]), compartments [248, 252, 443, 449] (but again cf. rebuttals [72, 331, 332]), CNS neurons [179, 181, 182], photoreceptors [713], and bristle cells [356, 369]; C. elegans neurons [228, 939], vulval cells [840, 845], and other cell types [363]; yeast mating types [812]; Dicystosteleum prespore vs. prestalk cells [56]; and flowering vs. vegetative states of plant apical meristems [687]. Particular histospecific genes in various organisms are likewise thought to employ combinatorial control mechanisms [101, 175, 232, 477, 532, 880].

Convincing evidence for a combinatorial code has come from analyses of homeotic mutations that affect flower development in two distantly related plants: the crucifer Arabidopsis thaliana and the snapdragon Antirrhinum majus [67, 97, 121, 585, 698, 773]. In each of these species, there are 3 genetic functions - a, b, and c - that specify the identity of 4 types of organs (each of which occupies a separate whorl along the axis of the flower): sepals, petals, stamens, and carpels. The code has been deciphered by studying the phenotypes of double and triple mutant combinations of null alleles which cause particular organs to develop as the 'wrong' type. If the states of the a-, b-, and c-type genes are represented as a triplet code - with '1' indicating an active gene (or functional allele) and '0' indicating an inactive gene (or null allele) - then the codes for the various organs have been shown to be: sepals (100), petals (110), stamens (011), and carpels (001) [359].

In order for a cell (or a computer) to act upon particular combinations of 1's and 0's in a binary code, it must be capable of executing Boolean logic [305, 1010]. Continuing with the Arabidopsis example, if there were a regulatory gene that specifies the petal state, then it would only be expressed 'if a is on AND b is on AND c is off'. 'Decoding' operations of this kind are usually thought to involve multimeric complexes of transacting regulatory proteins, which control the transcription of histospecific genes by binding at upstream regulatory sites [54, 55, 162, 234, 277, 374, 676]. In the case of the 'petal gene', an upstream site(s) would presumably bind dimers of a and b proteins, and only when that site(s) is occupied would the gene be transcribed. (Evidence for regulation of Drosophila genes by helix-loop-helix protein dimers is discussed below.) Given that some genes have as many as 20 upstream regulatory sites, which can bind as many as 12 different trans-acting factors [880], the potential capacity of each gene to act as an information processing device becomes appreciable, leading to the notion that cells may indeed be smarter than we think [48, 162].
Chapter 7

Default States

Surprisingly, '000' in *Arabidopsis* and *Antirrhinum* does not encode any organ of the flower, but rather specifies 'leaf'; in the *abc* triple mutant, all organs of the flower are transformed into leaves [121, 585]. In his 'Metamorphosis of Plants' published in 1790, Goethe had used morphological evidence to argue that the leaf represents the 'ground state' for all floral organs [17]. A colloquial version of the Ground-State Problem is [318]: 'Is a zebra a white horse with black stripes or a black horse with white stripes?' (It is the latter [33]; cf. Carroll [98] and Lawrence [490].) Much significance has generally been attributed to such 'default' states [4, 50, 396, 845, 857], i.e. the fates that cells adopt in the absence of signals that would normally direct them into one pathway vs. another. The usual assumption is that ground-state anatomies are 'atavisms', i.e. anatomies which existed in evolutionary ancestors prior to the origin of the genes necessary to encode the 'higher' states [7, 248, 250]. Thus, when the entire bithorax complex of *Drosophila* is deleted, nearly all body segments transform into a (normally leg-bearing) second-thoracic segment [507, 787], implying that this 'centipede' anatomy is the ancestral phenotype. However, null mutations at other homeotic loci (which interact with the complex) transform nearly all segments into legless eighth-abdominal segments [169, 759, 850, 852, 855] – a wormlike condition [704]. Are both anatomies ancestral, with one more ancient than the other (cf. French [241])? Evolutionary interpretations of this sort can be misleading until more is known about both the genetic regulatory network and homologous networks in outgroup species [50].

Linguistic Hierarchies

As biologists, we are accustomed to thinking about symbolic languages and information processing [70, 303, 422, 504, 678, 1013]. Chromosomes use 'DNA language', whose alphabet contains 4 letters (the nucleotide bases), which are arranged in 3-letter words (codons), which in turn are grouped in sentences (genes) that are punctuated by start and stop codes (level I). Ribosomes then translate the text of each gene into 'protein language' (level II), whose 20 letters (amino acids) are arranged in a variety of motif domains (zinc fingers, leucine zippers, homeoboxes, etc. [10, 234, 503]) which dictate the protein's function as an enzyme, gene regulator, structural component, etc. Control mechanisms at the level of gene regulation [439, 446, 791] (e.g. operon circuitry [305]), metabolism (e.g. activation of one enzyme via phosphorylation by another [404, 606, 875]), and cytoarchitecture (e.g. filamin proteins cross-linking an actin network [10]) would constitute a third echelon (level III) of command structure [127], though here the linguistic analogy weakens since there is no one-to-one correspondence with units or sequences at the lower level [308, 645, 955, 1013].

If there is an alphabet of development at the cellular level, then it must consist of unitary cellular behaviors such as mitosis, movement, signaling, adhesion, repulsion, state-change, shape-change, polarity-change, and suicide [20, 484, 663, 956]. Each of these behaviors (level IV) is implemented by the underlying network of gene-regulatory, metabolic, or cytoarchitectural factors (level III). By combining specific cellular behaviors in particular sequences, various gadgets (including any mechanism considered in chaps. 1-6) could be built. At this echelon (level V), the 'words' would be these modular 'subroutines' (see below), e.g. a 'word' with three 'letters' might be: 'Divide twice, then rearrange in such-and-such a manner, then differentiate in such-and-such a way based upon your new positions'. The linguistic analogy regains its power at this level, since it is not merely the combination of subunit commands but their permutation that matters, as in English syntax [590]. Ultimately, the entire program of embryonic development (level VI) would be translated into a final anatomy (level VII). (For a different stratification scheme, cf. JI [423].) Computer languages are designed in this same layered fashion [878]: words in the highest-level language (which the user employs to input instructions) are implemented by strings of operations at successively lower levels, which are ultimately based upon a binary machine code analogous to the base sequence in DNA. DNA may not be the only 'hard disk' inside a cell: an intriguing recent idea is that the conformational states of tubulin dimers within intact microtubules may function as bits for the storage of cellular memories and the processing of cellular information [344, 345, 708].

The hierarchical nature of development was popularized by C.H. Waddington, whose 'epigenetic landscape' metaphor envisions developmental pathways as valleys in a landscape (fig. 11a) [380, 764, 766, 925, 926, 930]. The contours of the landscape are stabilized by an underlying framework of interconnected guy wires, which ultimately are pegged to individual genes. The genes function like puppeteers, with mutations manipulating the landscape and causing cells to follow deviant paths that
lead to novel phenotypes. Paul Weiss [952, 954, 955] argued for the importance of emergent properties at all levels of the embryological hierarchy, especially during ‘self-organizing’ phases (cf. [308, 527]). Ergo, he reasoned, a purely reductionist molecular approach (comprising levels I–III) to development, which neglects cellular mechanisms (levels IV–VI), is bound to miss important clues that might help solve the puzzle — an opinion shared by many others [299, 349, 645, 877, 970, 997] (but cf. Caplan [961]).

Stent’s Cat, Brenner’s Virus, and the ‘Homeobox Homunculi’

In thinking about the question of how genes ‘compute’ anatomy, we confront the entire span of levels from genotype to phenotype: from linear DNA to three-dimensional morphology. Whether a ‘Make-a-Hand’ program [69], for example, actually exists as such in our DNA depends upon which cellular mechanisms are actually involved. To the extent that the mechanisms are self-organizing and not deterministic, the correspondence between anatomical parts and genetic counterparts will be lessened [273, 309, 341, 536, 537]. In any event, the execution of a definite sequence of steps at a higher (cellular) level does not necessarily imply a colinear sequence of clustered elements (regulatory genes) at a lower level [504].

An epigenetic view of development has been advocated by two eminent developmental (cum molecular cum neuro-) biologists — Gunther Stent and Sydney Brenner — who present different clues as evidence. Stent’s witness is the Siamese cat [825, 826]. The crossed eyes of this cat are attributable to a disturbance of the contralateral projection of its retinal axons, which is ultimately traceable to the same defective tyrosinase gene that gives the cat its peculiar coat color [333, 302, 781, 782]. Tyrosinase catalyzes the synthesis of the dark pigment melanin, and the retinal neurons are descended from the same cells that form the pigmented layer of the retina, which may explain their deviant afferent paths. Hence, there is apparently a cascade of errors from the mutant gene to the phenotype, but in no sense, Stent maintains, can the tyrosinase gene be imagined to ‘specify’ the neural circuitry of the visual system. By implication, there may be no genes whatsoever that directly specify that circuitry — or any other complicated anatomy. (However, it is wise to try to deduce how normal genes control development based upon the effects of mutations that affect development [961, 962].) Stent cites Waddington’s epigenetic land-
scape as the appropriate metaphor here: it is the interactions among many genes that collectively determine the outcome of dynamic processes at a higher (cellular) level. This same point is affirmed by Sydney Brenner, whose witness is the bacteriophage T4 [69]:

We now know that the shape and structure of something like phage T4 is not explicitly and uniquely represented in its genome ... The key process is self-assembly which depends on the bonding properties of many different protein molecules so that the representation of the structure is distributed over many genes in the DNA.

There is no gene in T4, Brenner surmises, that specifies 'Make an icosahedron' [504]. Again, by extrapolation, one could imagine that the more complicated anatomies of multicellular organisms are built skyscraper-style from so many intermediate levels of interactions – each with its own emergent properties – that the edifice is virtually entirely epigenetic, soaring far above its genome.

Development starts with a few ordered manifoldnesses; but the manifoldnesses create, by interactions, new manifoldnesses, and these are able, by acting back upon the original ones, to provoke new differences and so on. With each effect immediately a new cause is provided and the possibility of new specific action. – Driesch (1894) [185], as translated by Stern [832].

Early development (may be) regarded as a series of defined morphological and physiological stages, each with its own pattern, albeit difficult to discern, and each pattern serving as the spatial condition for the transformation of that stage to the next by a limited set of morphogenetic mechanisms ... There would be no long-range homing of the embryo toward the adult ... – Gerhart (1980) [274].

Opposing this argument are the 'homeobox homunculi': clusters of homeobox-containing genes in both insects and vertebrates, whose order along their respective chromosomes is colinear with the head-to-tail order of anatomical regions where they are expressed [168, 188, 260, 387, 511]. The conservation of this colinearity through some 600 million years of divergent evolution implies that the order must be important mechanistically (but cf. Holland [386]). The clusters may be positional-value 'memory boards', which record morphogen concentrations in the primary gradient field [259, 261, 403] (cf. Hanscombe et al. [348] and Sander [757]). While such one-gene-one-structure correspondences may be rare in embryos (cf. macrochaete genes [275, 743]), there is growing evidence (next section) that the genome is functionally (if not physically) fragmented into subunits which play multiple roles in development [1021].

Subroutines and Modules

A 'subroutine' is an algorithm that can be accessed merely by specifying its name once its sequence of instructions has been defined elsewhere [467, 934]. Subroutines can be repeatedly incorporated as modular building blocks in larger programs, which in turn can be used as units in still larger programs. An advantage of subdividing a complicated process into modular units is the efficiency it affords in the face of errors [290], as illustrated by a parable of Herbert Simon's:

The parable concerns two watchmakers, Hora and Tempus. Both make watches consisting of a thousand parts each. Hora assembles his watches bit by bit; so when he pauses or drops a watch before it is finished, it falls to pieces and he has to start from scratch. Tempus, on the other hand, puts together sub-assemblies of ten parts each; ten of these sub-assemblies he makes into a larger sub-assembly of a hundred units; and ten of these make the whole watch. If there is a disturbance, Tempus has to reset at worst nine assembling operations, and at best none at all. If you have a ratio of one disturbance in a hundred operations, then Hora will take four thousand times longer to assemble a watch – instead of one day, he will take eleven years [468].

Aside from error correction, the biological advantages of constructing anatomy in a modular manner [142] theoretically include: (1) economy of genetic specification, since a relatively small number of components needs to be encoded, and (2) acceleration of anatomical evolution, since changes in one subsystem need not affect other subsystems [69, 504]. A developmental subroutine would be a ritualized series of genetic or cellular actions that is used in multiple places or at multiple times during development. In the realm of morphogenesis there appear to be common subroutines for the folding of epithelia into tubes, pockets, or vesicles (regardless of which particular organ's epithelium may be involved in any given species) [30, 222, 335, 336, 628, 629, 651]. At the genetic level, the lexicon of patterning subroutines may include the following:

1) A neural-patterning machine. In Drosophila, a single basic mechanism constructs both the embryonic central nervous system (ECNS) and the adult peripheral nervous system (APNS) [94, 369]. Two types of genes are involved, and they function sequentially. First, 'equivalence group' (EG) genes establish territories within the ectoderm (ECNS) or epidermis (APNS), presumably in response to a coordinate system of positional information [93, 276, 789]. Then, following the stochastic inception of neuroblasts (ECNS) or bristle cells (APNS) within these areas, the 'inhibitory
field' (IF) genes enable these cells to inhibit neighboring cells from differentiating in a like manner [356, 365, 369, 788]. Subsequently, the neuroblasts and bristle cells (1) delaminate from the epithelial layer, and (2) undergo a stereotyped series of mitoses leading to a clone of ganglion neurons or bristle-organ cells (shaft cell, socket cell, sheath cell, and one or more neurons), respectively [354, 355, 903]. It is noteworthy how many different schemes are concatenated in this single pathway [180, 369]: positional information, random selection, inhibitory fields, cell rearrangement, and cell lineage (though, as discussed earlier, the final cell-lineage step may not directly assign bristle-cell fates). Because the sets of EG genes used by the ECNS and APNS overlap to a large extent - as do the sets of IF genes - it is likely that the two nervous systems are built by the same machine [18, 94, 369], with the idiosyncratic features of each neuroanatomy being crafted by the unshared genes (which may 'tune' the parameters of the EG and IF subroutines to different settings).

(2) An analog-to-digital transducer. Some of the same genes that are used to establish equivalence groups during Drosophila neurogenesis (as described above) also function in embryonic segmentation and sex determination - processes that are seemingly unrelated to each other as they are to neurogenesis. What all three systems do share is a dichotomous cellular choice: each cell must decide to become: (1) neural or non-neural; (2) part of the segment (parasegment [490, 491, 556]) or the intersegmental membrane (parasegment boundary); or (3) male or female. Although each decision is inherently 'digital' (i.e. two distinct alternative states), the factors that influence the choice may vary continuously in an analog manner. Hence, there is a need for some sort of analog-to-digital molecular transducing device. The protein products of the equivalence-group genes (hairy, extramacrochaetae, daughterless, and the four achaete-scute genes) possess a helix-loop-helix binding domain that allows them to form homo- or heterodimers [11, 100, 746] (thus integrating positive and negative analog signals), and a separate DNA-binding domain in some of them enables the dimers to bind (and regulate) other genes that are directly responsible for encoding states of cellular determination (thus converting the signals to the digital ON or OFF state of a target gene) [39, 214, 257, 258, 918]. The situation is best understood for sex determination, where each cell first measures a ratio between 'numerator' (X-chromosomal) and 'denominator' (autosomal) gene products, and then switches on or off a master gene (the Sex-lethal locus) based upon these inputs [218, 382, 674, 895]. (Helix-loop-helix dimerization cannot be the whole story since run, a segmenta-

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The whole ensemble of pair-rule genes may constitute a separate analog-to-digital device [207]. The ability of a homologous vertebrate gene - MyoD - to transform fibroblasts into myoblasts [652, 942, 943] implies that the same device may be used for specifying certain cell states in vertebrates.

The above examples illustrate how sets of genes can be used repeatedly in different pathways during development. Indeed, an emerging theme in developmental biology is that small numbers of genes are often 'plugged into' large numbers of developmental circuits [845]. Cases where the same genes are successively used for different functions within a single pathway include: (1) the re-usage of Drosophila IF genes to specify cellular fates within bristle organs (e.g. a shaft cell vs. a neuron) [356, 369] and ommatidia (e.g. a cone cell vs. a pigment cell) [92]; (2) the re-usage of Drosophila segmentation genes to encode neural cell identities within each segment [177, 179, 181, 182, 190, 677]; and (3) the re-usage in Arabidopsis of one of the c genes (that functions in the organ identity code) to limit the number of whorls along the flower axis [121]. The efficient utilization of a limited number of genes makes sense from an evolutionary standpoint, and it helps explain why so many mutations have pleiotropic phenotypes [20, 723]. A related - and still unresolved - question is whether genomes allocate separate genes for patterning and housekeeping (i.e. vital metabolic or physiological functions) [69, 70, 141, 834].

There appear to be many situations where single subroutines are employed in similar roles at multiple locations during development. All of them entail 'serially homologous' [136, 166, 737, 920, 932, 933] organs. As mentioned in the Introduction, such patterns are common in multicellular organisms. Their naturally occurring variants were the subject of Bateson's 'Materials for the Study of Variation' (cf. Kellogg and Bell [453] and Maynard Smith [558]). Consider your hands versus your feet. Because they have the same basic skeletal anatomy (as do your entire arm and leg), they may be outputs of a single subroutine, whose variables (e.g. bone lengths and articulation angles) assume different input values (see, below; cf. Riedl [723] and Tabin [873]), depending upon where the limb develops [736]. Indeed, although the vertebrate limb has evolved into structures as superficially different as elephant legs and bat wings, the essential pattern of bones has remained constant [166, 312, 379, 384, 783], suggesting that the subroutine has been highly conserved while the settings of its input variables have changed drastically. If the subroutine itself were to be altered (either muta-
tionally or evolutionarily), then the two pairs of limbs should change their morphologies coordinately. Numerous examples of such correlated changes are known [379, 419, 543, 1003]. For instance (1) the brachydactylyous anomaly 'A2' in man concomitantly shortens the middle phalanges of the index finger and the second toe [1003], and (2) the pan has evolved a thumb-like sesamoid bone on both its hindfeet and forefeet, though only the bone on the forefoot has any apparent function [736]. Comparable coevolutionary trends have been found for tooth patterns in the upper versus the lower jaw in mammals [88]. Perhaps the clearest example of how evolution can tinker with developmental subroutines is in nematodes, where lineages vary among species according to the same few rules that govern differences among lineages within the species *C. elegans* [326, 459, 840–842].

**Iteration and Halt Conditions**

An intriguing aspect of many periodic patterns is their exact numbers of pattern elements [136, 233, 558]. Why, for instance, do humans typically have 10 fingers, 24 ribs, and 32 teeth (cf. the American Flag Problem)? What mechanisms ensure such constancy? This question was posed as the 'Counting Problem' by John Maynard-Smith in 1968 [559]. He envisioned two types of possible 'counting machineries’, which could perform an operation *n* times: (1) 'ratio' counters, where *n* is the ratio between two quantities, and (2) 'digital' counters, where *n* is the number of interdependent events in a finite series. An example of a ratio-counting mechanism is the triggering of blastular events by the nucleocytoplasmic ratio during cleavage. Studies of haploid and polyploid embryos (which have smaller or larger nuclei, respectively) in *Xenopus, Ambystoma* (an axolotl), and *Drosophila* indicate that the number of cleavages (and the onset of transcription known as the 'mid-blastula transition') are determined by the nucleocytoplasmic ratio [206, 319, 399, 460, 464, 639, 640, 762]. The most popularly hypothesized digital counting devices are 'chemical counters' [558], where different gene products are synthesized at each step of a process (A, B, C, etc.) until the last product terminates the process. Such models have been invoked to explain the number of cleavages in *Volvox* [463, 869, 870]; the numbers of segments in horseshoe crabs [416], short-germ insects [757], and leeches [210] (including leech segmental identity [553]); and the timing of embryonic events in general (with or without a causal link to mitosis) [760, 761] (cf. Cooke and Smith [145]). (As for how mitotic coun-

ters might trigger differentiation events, cf. Temple and Raff [876] and the flowchart in figure 16–38 of Alberts et al. [10].)

What would happen if the counting mechanism for an iterative process were to malfunction because of a failure of its 'halt condition' (i.e. the rule that dictates 'IF condition *x* prevails THEN stop')? As programmers alas know well, such default 'bugs' can cause repetitions to occur ad infinitum. Mutations that cause comparable 'infinite loop' phenotypes include: unc-86 mutations in *C. elegans*, which force certain lineages to repeat their pedigree rules many times over [104, 228, 229] (cf. related nematode mutations [12, 750]); the *floricaula* mutation in the snapdragon *Antirrhinum* [120, 122], which replaces flowers with iterated inflorescence meristems (cf. comparable mutations in other plants [871]); and *bag-of-marbles* mutations in *Drosophila* [565], which increase the number of cells per ovarian cyst from 16 (a precise number in wild-type flies, which is due to exactly 4 mitoses per germ cell) to between 50 and several hundred (cf. Lifschytz [517]). Such 'bugs' may have been instrumental in the evolutionary origin of extremely high numbers of vertebrae in snakes and body segments in millipedes.

'Counters' are only applicable to patterns whose elements are specified sequentially. The issue of how element number is controlled in 'synchronous' [927] patterns would seem to call for entirely different mechanisms [368, 558, 728] (but cf. Meinhardt [573]). As discussed in chapter 1, positional information models are designed to ensure constant (size-independent) patterns, including fixed numbers of pattern elements. For pre-pattern mechanisms, some authors have envisioned the triggering of the prepattern algorithm only when the developing organ reaches a critical size [134, 136, 138, 1008]. For determination wave mechanisms, the rate of the wave movement along the tissue could be established earlier by a positional information scheme [146]. Interestingly, there are patterns whose element number varies along one axis but not another. Examples include mouse whiskers [191] and fly bristles [367]. In the latter case, the number of rows of bristles is size-invariant, whereas the number of bristles within each row varies in proportion to the size of the organ [367].

**Input-Output, 'Morphospaces', and Evolutionary Constraints**

The notion that developmental pathways can constrain anatomical evolution was championed by D'Arcy Thompson [882] (e.g. his 'grid transformations') and Richard Goldschmidt [295] (e.g. his 'norm of reactivity'; cf.
Kaufman [444]). It has experienced a revival lately, sparked by Stephen Gould's book 'Ontogeny and Phylogeny' [315] and fueled by Alberch et al.'s [9] 'ontogenetic trajectory' technique for operational analysis [466, 608, 704]. A vivid illustration of the relationship between development and evolution is provided by Raup's studies of the shapes of mollusk shells [709, 710, 712]. Only three variables (the rate of growth of the shell's mouth, the mouth's distance from an imaginary axis around which it revolves, and the mouth's rate of translation along that axis) are required to simulate most shell shapes. The variables define a 3-dimensional 'morphospace' [316, 560] within which the actual shapes of living and fossil families of molluscs can be plotted as contiguous domains [711]. Evidently, all molluscs share an ancient 'SHELL' algorithm, which can produce a variety of output shapes (e.g. corkscrew, planar spiral, or bivalve), given different species-specific settings of the input variables. Shapes outside the space should never arise unless the mechanism itself is altered. Moreover, if each variable is under separate genetic control, then the most likely pathways of evolutionary change from any given point in the space would be along vectors parallel to the axes.

Comparable morphospaces for patterns can be imagined for many of the mechanisms discussed earlier [646, 718, 840]. One example would be the morphospace of possible phyllotactic patterns (spiral, distichous, and whorled) defined by inhibitory field [593, 721, 883] or physical force [325] models. A generic evolutionary trend that is expected for any mechanism which lacks the ability to regulate (e.g. prepattern devices) is a proportionality between the number or pattern elements and the size of the pattern [367] (chap. 3).

Positional information mechanisms are not usually helpful in predicting how patterns should change with evolution [540, 1002]. However, one prediction concerning digit patterns has led to some provocative questions. A complication in trying to apply the Polar Coordinate Model [81, 245] (chap. 1) to vertebrate limbs (vs. insect appendages) is that the coordinate system is supposedly centered on the tip of the limb, but the tip branches into digits, which raises two questions: (1) which digit, if any, lies at the origin, and (2) what role, if any, does the coordinate system play in specifying digit positions? Based upon the results of digit amputation experiments with newt limbs, Stock and Bryant [848] proposed that (1) none of the digits resides at the origin, and (2) each prospective digit inherits a different subset of coordinates which are lateral to the origin. Because the subsets contain less than half of the circumferential coordinates, they automatically (due to the Shortest Intercalation Rule) undergo duplication. Such duplication explains why vertebrate fingers are internally mirror-symmetric (i.e. the left half of each finger is a mirror image of the right half). Moreover, because those digits closest to the origin would contain more coordinates, they should grow to greater length. Hence, the 'Stock-Bryant Rule' for digit evolution dictates that the lengths of the digits within a limb must decrease linearly from a single 'high point' (defined by the origin of the coordinate system). That point could be centrally located (e.g. the human hand, where the middle finger is longest) or more lateral (e.g. the human foot, where the longest digit is typically the hallux), but there should never be a pattern where two long digits flank a shorter one (since this would imply two high points) [848]. Contrary to expectation, such patterns do exist, though they are rare [383, 627]. Alternative models, based upon sequentially branching condensations of cartilage cells, have been proposed by Oster et al. [660, 663, 783] and others [636, 638, 966], and they predict an entirely different set of developmental constraints [8, 208, 619, 663, 783]. The reader should consult Gardiner and Bryant [253] for what may be a definitive answer to the classical problem of the 'metapterygial axis' and the puzzle of 'Gregory's Pyramid'.

Game Theory

Every scientist dreams of discovering a universal law of nature that could crystallize a collection of seemingly unconnected facts into a glorious gem of insight. Biologists have been frustrated in this quest because living beings, unlike atoms, are only comprehensible in the context of their histories, and different species have evolved differently [272, 417, 563, 946]. There is, unfortunately, no periodic chart of cell types, nor any equivalent of 'F = ma' that governs developmental trajectories. However – and here is where hope may still lie for some fulfillment – there may be a finite number of elemental patterning strategies [162, 489]. The rainbow spectrum of species-specific developmental pathways could then be unwoven into a limited number of primary strands. The collection of schemes surveyed in the earlier chapters, together with the kinds of devices discussed in this chapter, may constitute a large portion of the contraptions that evolution has invented to solve the problem of embryo engineering. If that problem is envisioned as a game, then this book represents an attempt to devise a pluralistic 'game theory'. Even if all the models and metaphors cannot unite development's Gordian Knot, they at least portray possible solutions that can inspire us as we grope.