From DNA to Diversity
To our families

“From DNA to Diversity is written for a general audience, including undergraduates, with an interest in developmental and evolutionary biology, and it is a joy to read. Using striking examples, the authors summarize the current state of thinking on the interconnectedness between developmental genetics and evolutionary diversification.”
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“This book helps to fill a gap in the teaching of evolutionary theory that arose because developmental biology was not a direct participant in the evolutionary synthesis. . . . This is an outstanding account of the latest findings in molecular developmental biology.”
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Martin J. Cohn, University of Florida; Evolution and Development

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“From DNA to Diversity can be, and should be read by College and University students as well as scientists out of the field, who want to be informed of what is new and promising in biology.”
Jean Deutsch, Universite Phillippe et Marie Curie, Paris; BioEssays
From DNA to Diversity

*Molecular Genetics
and the Evolution of Animal Design*

2nd edition

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The Earth is now populated by between 1 million and perhaps as many as 20 million animal species, which represent probably less than 1% of all animal species that have ever existed. An even more remarkable fact is that all of this diversity—aardvarks and ostriches, butterflies and pythons, dinosaurs, and earthworms—descended from a common bilaterally symmetrical ancestor that lived in Precambrian seas more than 540 million years ago. Traditionally approached through paleontology, systematics, and comparative anatomy, the story of animal evolution has, until recently, been sorely missing one huge chapter—namely, genetics.

Animals diverge from common ancestors through changes in their DNA. The major question, then, is, Which changes in DNA account for morphological diversity? The answer to this question has eluded us for the half-century since the Modern Synthesis was proposed and the structure of DNA was discovered. Although many reasons exist to explain this omission, foremost among them is that biology first had to address another central genetic mystery—that is, which genes out of the thousands in any species control morphology?

One of the most important biological discoveries of the past two decades is that most animals, no matter how divergent in form, share specific families of genes that regulate major aspects of body pattern. The discovery of this common genetic “toolkit” for animal development has had two major implications for researchers. First, it has enabled biologists to uncover widely conserved molecular, cellular, and developmental processes whose existence was concealed by previously incomparable anatomies. Second, it has focused the study of the genetic basis of animal diversity on how the number, regulation, and function of genes within the toolkit have changed over the course of animal evolution.

The genetic picture of morphological diversity presented in this book is highly influenced by the legacy of previous successes of genetic logic. The mysteries of enzyme induction in bacteria and bacteriophage life cycles were, through formal genetic logic and molecular biology, ultimately reduced to elegant genetic switches that determined the on/off state of groups of genes. This success laid the foundation for understanding the regulation of genes in different cell types of multicellular organisms and, in turn, the regulation of genes in space and over time during the development of individual organisms. Similarly, recent advances in understanding how the toolkit operates in the design of just a few model species has laid the foundation for studies of the evolution of a wide variety of animal structures and patterns.

The presentation in this book lies at the intersection of evolutionary biology with embryology and genetics. Comprehensive treatment of any of these long-established, fast-growing disciplines can be found in full textbooks dedicated to each. Because our goal is to elucidate general principles about the genetic basis of morphological change, we will focus on those
genes, developmental processes, and taxa that are best known and best illustrate these principles. The book is organized into two parts. The first part (Chapters 1–3) focuses on the history of animals and on animal developmental genetics and regulatory mechanisms. We first examine some of the major trends in animal design and evolution illustrated in the fossil record and by modern forms (Chapter 1). Next, we take an inventory of the genetic toolkit for the development of model species (Chapter 2). Finally, we analyze the regulation and function of these genes in the complex hierarchies that govern animal development (Chapter 3). This crucial background knowledge of the major transitions in animal evolution and the genetic logic of animal design sets the stage for the analysis of mechanisms of morphological evolution.

The second part of the book examines the genetic mechanisms underlying the evolution of animals at different morphological levels. We take a case study approach by focusing on the best-understood examples of the evolution of the genetic toolkit, the diversity of body plans and body parts, and novel structures. In the final chapter, we discuss why and how changes in gene regulation have played a primary role in the evolution of diversity across the morphological spectrum—from small-scale differences within or between species, to the large-scale differences that distinguish higher taxa.

We have provided selected references for further reading at the end of each chapter. By no means should these citations (or this book) be taken as the primary or exclusive references on a topic. For both brevity and to circumvent questions of priority in ideas or evidence, we have avoided attributions to specific authors in the text.

One of the inspirations for our approach was Mark Ptashne’s classic A Genetic Switch, in which many of the basic physiological and molecular principles of gene regulation were illuminated by focusing on the bacteriophage λ. In the preface, Ptashne stated that “one of the charms of molecular biology is that the answers it provides to fundamental questions for the most part can be easily visualized.” Few fields in biology can rival the aesthetic appeal of the new comparative embryology. Indeed, the visualization of members of the genetic toolkit in action during the development of different species has already become a surrogate for analyzing final forms. For those who find conceptual beauty in the logic and molecular anatomy of genetic switches, the genetic switches controlling animal anatomy may be even more appealing. Not only do they control the striking patterns of gene expression within developing embryos, but as we shall see, they are also key to understanding how the wonderful, but presently dwindling, diversity of animal forms has evolved.

**CHANGES IN THE SECOND EDITION**

The revision and expansion of From DNA to Diversity for this second edition is driven by advances on many fronts. Increased understanding of developmental mechanisms, systematic exploration and comparisons of animal genomes, and inquiries into new models of morphological evolution have provided a wealth of case studies from which we have selected new material. Much of the new coverage in this edition is found in the second part of the book, which has been expanded to five chapters (Chapters 4–8) from four in the first edition. Information and references have been updated throughout the book. Again, we stress that these citations are selective and that neither they nor this book should be taken as the primary or exclusive reference on a topic.
The book’s overall organization remains the same, with the first three chapters devoted to the history of animals, developmental genetics, and genetic regulatory mechanisms. The second part of the book (Chapters 4–8) examines the evolution of animals at different morphological levels. The explosion in genome sequence data has provided an enormous increase in the quantity and quality of information concerning the evolution of the genetic toolkit for animal development. Many animal genomes, including our own, have been sequenced since the publication of the first edition. Some of the major insights from genome studies have been added to Chapter 4.

The growth of evolutionary developmental biology has provided new insights into the diversification of specific body plans and the origins of animal novelties. Chapters 5 and 6 have been revised and expanded to incorporate new findings ranging from mechanisms of segmentation in spiders, to the evolution of the cephalopod body plan, and the origin of the turtle shell.

There has also been an increasing focus on models of variation within species and of divergence of traits. Some of the simplest models of phenotypic variation and evolution involve the color patterns of mammals, birds, and insects. In several cases, the identity of genetic differences responsible for variation between populations is now known. We have added a new chapter (Chapter 7) that focuses on models of variation and divergence among closely related species.

ACKNOWLEDGMENTS

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The central focus of this book is to identify the genetic mechanisms underlying the evolution of animal design, particularly with regard to the patterning of animal body plans and body parts. To approach this mystery, new discoveries and ideas from developmental genetics must be integrated into the larger framework of the evolutionary history of animal life. This history is reconstructed from many fields of study—in particular, paleontology, systematics, and comparative biology. In this chapter, we present a brief overview of animal evolution from these three perspectives. This discussion provides a historical foundation for the consideration of the mechanistic questions that are addressed in subsequent chapters.

First, we discuss the origin of animals and the radiation of the major animal phyla based on evidence gleaned from the fossil record. Most living phyla have ancient origins, and the fundamental differences between them evolved long ago. Two milestones in early animal history that are of special interest are the evolution of bilaterally symmetrical animals and the explosive radiation of these forms in the Cambrian period more than 500 million years ago.

Second, we examine the phylogenetic relationships among animals. Understanding the direction of evolutionary change in morphological, developmental, or genetic traits and the ability to make inferences about animal ancestors requires knowledge of the structure of the animal evolutionary tree. While traditionally based upon morphological comparisons, new phylogenies based on DNA and protein sequences have revealed unexpected relationships among anatomically disparate animals, refuting long-held notions about which phyla are more closely related.

Third, we consider the comparative anatomy of selected phyla with the aim of identifying some of the major trends in the evolutionary diversification of individual phyla. In particular, we focus on the modular organization of the body plans and body parts of larger animals—the vertebrates, arthropods, and annelids. Much of the large-scale morphological diversity within these phyla (for example, between different classes) involves differences in the

... an understanding of regulation must lie at the center of any rapprochement between molecular and evolutionary biology; for a synthesis of the two biologies will surely take place, if it occurs at all, on the common field of development.

—Stephen Jay Gould
Ontogeny and Phylogeny (1977)
number and pattern of modular elements (segments, appendages, and so on). The recognition of the modular organization of these animals is an important conceptual link to understanding the genetic logic controlling their development and the mechanisms underlying the evolution of diversity.

ANIMAL ORIGINS AND THE FOSSIL RECORD

The fossil record is our primary window into the history of life. It provides many kinds of information that cannot be inferred from living animals. Fossils give us pictures of extinct forms that may be ancestors of modern animals, provide minimal estimates of the time of origin or divergence of particular groups, reveal episodes of extinctions and radiations, and, in favorable circumstances, offer detailed accounts of the evolution of important structures.

The search for the origins of modern animals begins with an assessment of the Cambrian fossil record. It has been known since before Darwin’s time that animal diversity increased dramatically during this period, which spans an age from roughly 545 to 490 million years ago (Ma). Molluscs, arthropods, annelids, chordates, echinoderms, and representatives of most other modern phyla make their first appearance in Cambrian fossil deposits (Fig. 1.1). The emergence of large, complex animal forms and their radiation over a 10 to 25 million year interval in the Early–Middle Cambrian is often referred to as the “Cambrian Explosion.”

The appearance of these animals in the Cambrian fossil record gives us only a minimum estimate of their time of origin. The crucial question about the Cambrian Explosion is whether it marks the origin of animals or the origin of modern phyla. Did most phyla first arise in this short period, or did they predate their preservation in the Cambrian fossil record? Although the Precambrian animal fossil record is relatively scarce, several kinds of fossil evidence indicate that the origins of most modern phyla predate the Cambrian. First, the fossil record of some modern groups clearly begins before this period. For example, body fossils of both cnidarians and sponges predate the Cambrian (Fig. 1.2). Both of these groups are diploblastic animals, composed of two tissue layers. The cnidarians have a radically symmetrical body design that distinguishes them from sponges and from a much larger number of modern phyla that are triploblastic—a that is, composed of three tissue layers—and have bilaterally symmetrical body designs (the Bilateria). Second, Precambrian deposits contain evidence in the form of trace fossils, the record of the meanderings and burrowings of animals in sediments, which indicate the existence of some bilaterian forms (Figs 1.2 & 1.3d) well before the Cambrian Explosion. A third piece of potential evidence for earlier animal origins is the Ediacaran fauna (575–544 Ma), named for the Australian locale in which they were first discovered.

The biological interpretation of Ediacaran fossils and their relationships, if any, to modern animals remains controversial. Several distinct body plans have been identified, including radially symmetrical types and a number of frond-like and tube-like forms (Fig. 1.3). None of these bear any clear-cut similarity to modern animals, so they have been difficult to place on the tree of animal evolution. Some of the Ediacaran fossils could represent diploblastic forms related to cnidarians or sponges. Others could be primitive bilaterians that possess some, but not all, features of modern bilaterians.

The difficulties in placing Ediacarans in the scheme of animal evolution have led to the proposal that they represent an extinct experiment in multicellular life. On the other hand, perhaps their lack of resemblance to modern groups is exactly what should be expected.
Chapter 1: A Brief History of Animals

of primitive animals. It is possible that the Ediacaran fauna include both extinct types of diploblastic animals and primitive ancestors of modern bilaterians. The fossil record indicates that some Ediacaran forms persisted into the Cambrian, but then died out as bilaterians, sponges, cnidarians, and ctenophores flourished.

Given the uncertainty of the relationship of the Ediacarans to modern phyla and the paucity of body fossils prior to the Cambrian, it is difficult to pinpoint the origins of modern animals based on the fossil evidence. Consequently, biologists have turned to other methods to try to identify when major animal groups diverged. Using the evolution of protein and ribosomal RNA sequences between species to calibrate **molecular clocks**, estimates of the time of divergence of most animal phyla have been made that range from approximately 650 Ma to more than 1000 Ma. While these estimates remain controversial, even the most conservative estimate suggests a period of more than 100 million years before the beginning of the Cambrian in which most bilaterian phyla had arisen but led a paleontologically cryptic existence.

![Figure 1.1](image)

**Figure 1.1**

**Cambrian animal fossils**

Representatives of many modern phyla are found in Cambrian deposits and are made up of repeating units. (a) *Aysheaia pedunculata*, an onychophoran; (b) *Burgessochaeta setigera*, a polychaete annelid; (c) *Pikaia gracilens*, a chordate; (d) *Olenoides serratus*, a trilobitomorph arthropod; (e) *Waptia fieldensis*, a crustacean-type arthropod.

It is widely believed that primitive bilaterians may have been very small and their size limited by atmospheric and oceanic oxygen levels. This fact would help to explain their slim fossil record before the Cambrian (Fig. 1.2). In the last few years, evidence has also been gathered that suggests a possible mass extinction at the boundary between the Proterozoic and Cambrian. Whatever the cause of such an event, it may have hastened the extinction of Ediacaran forms and opened up the ecological opportunity for bilaterians to radiate. Environmental and ecological changes may have removed constraints on bilaterians, permitting the evolution of larger animals. In addition, competitive interactions among bilaterians may have
Figure 1.3

Pre-Cambrian animal fossils and traces

(a) Ediacaria, a radially symmetrical form from deposits in Australia. (b) Calcified fossils in limestone from Namibia. (c) Pteridinium, a frond-like ediacaran fossil form built of repeating units. (d) Trace fossils made in sediments by bilaterian animals.

facilitated the evolution of skeletonized taxa, more sophisticated predatory and defense behaviors, and the variety of anatomical innovations that unfolded in the Cambrian.

THE ANIMAL TREE

There are about 35 living animal phyla. To understand the origin and evolution of any feature found in one or more of these groups, it is necessary to have a picture of the phylogenetic relationships among animals. Ideally, the fossil record would present a complete, ordered, unambiguous picture of the branching pattern of the animal tree. Unfortunately, it does not. As the divergence of most bilaterian phyla appears to have predated the emergence of recognizable members of modern phyla in the fossil record, we must make our inferences from later, more derived forms.

Constructing an accurate picture of metazoan relationships has been challenging, and many alternative schemes of animal phylogeny have been proposed and scrutinized over recent decades and continue to be evaluated. Most approaches have relied on anatomical and embryological comparisons. In general, phylogenies are determined according to shared characters that are presumed to be derived and therefore reflect a close relationship. For example, all animal phyla are thought to be more closely related to each other than to any other nonanimal phylum, because of similarities in animal multicellularity, cell structure and morphology, and cell signaling. Members of the most closely related protist group, the choanoflagellates, share a similar cell architecture with sponges but are not multicellular. What is most difficult to determine is whether apparent similarities between animals (for example, segmentation in arthropods and annelids) are due to common ancestry, are superficial, or evolved independently. Also, different tree topologies can emerge when different characters are used or when the same characters are weighted differently.

One way to circumvent the reliance on morphological comparisons is to use molecular genetic characters to construct animal phylogenies. As taxa diverge, the sequences of DNA, RNA, and protein molecules diverge as well; the relative degree of divergence can therefore be used to infer phylogenetic relationships. In addition, the presence or absence of particular genes, or the linkage of a group of genes on chromosomes, can be used to construct phylogenetic trees. New methods based on molecular sequences have been combined with morphology-based approaches to both prune and strengthen the animal tree.

We now recognize shared morphological, developmental, and genetic traits that suggest that the Bilateria can be organized into three great clades (a set of species descended from a common ancestor) (Fig. 1.4):

- The deuterostomes, including chordates, echinoderms, ascidians, and hemichordates. The deuterostomes are named for a shared feature of early embryonic development in which the mouth forms from a site separate from the blastopore, an opening in the early embryo.

- Two groups of protostomes, in which the mouth develops from the blastopore. The protostomes are divided into the lophotrochozoans, including annelids, molluscs, and brachiopods, many of which share a trochophore larval stage in their life cycle, and a clade consisting of the arthropods, onychophora, and priapulids.
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Figure 1.4
Metazoan phylogeny
The current picture of metazoan phylogeny showing representatives of three major bilaterians clades—the deuterostomes, the Lophotrochozoa, and the arthropod + onychophora + priapulid clade.
Within these great clades, the branching order has been less well resolved, such that it is unclear which phyla are more closely related. It is worth noting that the recent assignment of arthropods and annelids to two different protostome clades and the assignment of pseudocoelomate phyla among different clades are major changes from previous portraits of the animal tree. The phylogenetic placement of the nematodes, including the model organism *Caenorhabditis elegans*, remains controversial, because their rapid molecular clock complicates analysis. Some phylogenies place the nematodes close to the arthropod + onychophora + priapulid clade and others more basally near the common ancestor of all bilaterian phyla.

The anatomical and developmental features of the Bilateria are very distinct from those of the basal metazoans (cnidaria, ctenophores, and porifera). The evolutionary links between basal metazoans and the bilaterians are difficult to perceive. Indeed, as we will see in Chapter 4, major differences exist between the genetic toolkit of these two groups, and the differences are much more substantial than those between most bilaterians. Because of the long divergence time since the radiation of these groups, the phylogenetic relationships between cnidarians, sponges, and ctenophores and the last common ancestor of the Bilateria are uncertain. Many extinct animal lineages, as yet unknown from the fossil record, may have branched off of the metazoan tree between the last common ancestor of all animals and of the Bilateria (Fig. 1.4).

The gaps in the fossil record; the great differences in anatomy, development, and genome content between radially symmetrical animals and bilaterians; and the cryptic early history of bilaterians, make inferences about the morphological transformations involved in the origin of animal body plans very speculative. Paleontologists have introduced the concept of *disparity* to refer to differences among body plans and use the term *diversity* to refer to the number of species within a group. The genetic and developmental bases of the morphological diversification of a *particular* body plan within a phylum are far more accessible than is the origin of different body plans. Therefore, we will focus primarily on evolutionary trends within a few select phyla, such as the arthropods and chordates, making the implicit assumption that the same sort of genetic mechanisms involved in the evolution of large-scale morphological diversity within phyla also gave rise to fundamental differences in body plans.

**GENERAL FEATURES OF ANIMAL DESIGN AND DIVERSITY**

One of the most outstanding features of animal design, particularly of larger bilaterians, is their construction from repeating structures (or modules). The segments of arthropods and annelids and the vertebrae (and associated processes) of vertebrates are the basic units of body plan organization in these phyla (Fig. 1.5a–c). Similarly, many body parts such as the insect wing (Fig. 1.5d) and the *tetrapod* hand (Fig. 1.5e) are composed of repeated structures.

An important trend in the morphological evolution of animals has been the *individualization* of modular elements. For example, among the arthropods, we observe a large number of different segment types in crustaceans and insects. This diversity far exceeds that found in the onychophora, a phylum closely related to the arthropods. Thus the evolution of the onychophoran/arthropod clade has been marked by increased diversity of segment types from the more uniform patterns found in earlier forms. Similarly, in some mammals, teeth are differentiated into molars, premolars, canines, and incisors, whereas in the ancestral condition exhibited by most reptiles, the teeth are of uniform shape. Because the diversification of the
Chapter 1: A Brief History of Animals

The body plans of many major phyla, including the annelids (a), arthropods (b), and chordates (c), are composed of many repeating parts. Some of these parts are similar or identical in appearance to other parts; others are individuated. Sets of serially homologous structures are shaded a unique color. Body parts, such as a butterfly wing (d), or a fossil tetrapod limb from the amphibian fossil Acanthostega (e), are also composed of repeating structures or patterns, some of which are differentiated from others. For example, Acanthostega has eight digits, but like its modern descendants, only five distinct types of digits can be distinguished.

Source: Parts a–c from Weatherbee SD, Carroll SB. Selector genes and limb identity in arthropods and vertebrates. Cell 1999; 97: 283–286; part e from Michael Coates.
number, morphology, and function of these repeated units characterizes many of the large-scale differences that distinguish related taxa, understanding how repeated structures form and become individualized is a prerequisite for understanding the developmental basis of large-scale morphological evolution.

The modular organization of animal bodies and body parts has long been recognized by comparative biologists. William Bateson, in his classic treatise *Materials for the Study of Variation* (1894), identified several kinds of organization found among animals. More importantly, he was the first to bring a Darwinian perspective to the question of how different body patterns may have evolved. Bateson focused particularly on the repetition of parts, cataloguing a large number of rare, but naturally occurring, variants that differed from the norms within various species with regard to either the number or individualization of characters. He suggested that these variations within species could provide insight into the evolution of the large-scale morphological discontinuities between species. For example, variations in the number of body segments within onychophora and centipede species, and of vertebrae in humans and pythons, suggested to Bateson that such discontinuities arose at some frequency in populations and therefore represented plausible steps in the morphological diversification of species.

The question of whether evolution may progress in large, discrete steps remains controversial (we will address this issue in Chapter 8). Nevertheless, these sorts of variants and the organizational concepts espoused by Bateson have been enormously helpful in understanding the genetics and developmental logic underlying the modularity of animal design. In fact, they led to the discovery of genes that play key roles in morphological evolution, albeit not in the fashion Bateson first imagined.

Four fundamental kinds of large-scale, evolutionary differences in morphology are most prevalent in modularly organized animals and are the most significant in terms of adaptation:

1. *Changes in the number of repeated parts* Bateson referred to this type of change as **meristic variation** when describing differences within species. Differences in segment number and vertebral number are some of the most obvious characteristics that distinguish classes of arthropods and various classes and orders of vertebrates, respectively (Fig. 1.6).

2. *Diversification of serially homologous parts* A series of reiterated parts are termed **serially homologous**. The individualization of repeated parts in an animal reflects the diversification of serially homologous structures. For example, arthropod appendages are serially homologous structures. In the course of arthropod evolution, ancestrally similar appendages have evolved into antennae, various mouthparts, walking legs, and genital structures. In vertebrates, serially homologous vertebrae have evolved into distinct cervical, thoracic, lumbar, and sacral vertebral types.

3. *The diversification of homologous parts* One of the most prevalent trends in animal evolution is the morphological diversification of **homologous** parts between lineages. The same structures in different lineages are termed “homologous” when they share a common history, even if they no longer serve the same function. For example, all tetrapod forelimbs are homologous (Fig. 1.7). Despite their differing appearances and functions, bird wings, bat wings, and human forelimbs have all conserved the basic architecture of the tetrapod forelimbs.
Figure 1.6

Meristic differences among arthropods and among vertebrates

Among arthropods such as this trilobite (a), crustacean (b), centipede (c), and insect (d), the number of body segments differs, as does the diversity of segment morphology. Among vertebrates, the number of vertebrae and associated processes differs considerably between a fish (e), frog (f), python (g), and chimpanzee (h).
Figure 1.7

The diversification of homologous parts

All vertebrate forelimbs are homologous structures whose anatomy has undergone considerable diversification in the evolution and adaptation of these various vertebrate lineages. Not to scale.

4. The evolution of novelties New characters or “novelties” may evolve from a preexisting structure or arise de novo and become adapted to a new purpose. The evolution of feathers, fur, teeth, antlers, and butterfly wing eyespots are examples of such morphological novelties.

Considering that modularly organized animals are among the most diverse groups (in terms of both the number and morphology of species), could there be a correlation between body design and evolutionary diversity? One possible explanation for this relationship is that modular organization allows one part of the animal to change without necessarily affecting other parts. The evolution of genetic mechanisms that control the individualization of parts would allow for the uncoupling of developmental processes in one part of the body from the developmental processes in another part of the body. In this fashion, for example, vertebrate forelimbs can evolve into wings while hindlimbs remain walking legs. Dissociation of the forelimb and hindlimb developmental programs allows further modifications to occur selectively in either structure, such as the development of feathers in the forelimb of birds and scales in the hindlimb.

EVOLUTION AND DEVELOPMENT: DNA AND DIVERSITY

To understand the major trends in animal diversity and the various kinds of morphological evolution, we must first understand how animal form is generated. Morphology is the product of development, the process through which a single fertilized egg cell gives rise to an entire organism. The physical basis of animal diversity has been viewed since Darwin’s time as the outcome of development. Until very recently, however, the developmental principles underlying animal design remained unknown. Although experimental embryologists of the late 1800s and the first half of the 1900s had identified many fascinating phenomena concerning the organization of embryos and the formation of particular structures, the mechanisms responsible for these properties were beyond their reach.

With better understanding of the nature of genes and the process of gene regulation, development has been increasingly viewed as a process orchestrated by the products of genes. Thus the puzzles of embryology, such as how cells come to know their position and identity within a developing animal, have become rephrased in genetic terms. Given that the DNA of (most) all cells in an animal is identical, how do different cells acquire the unique morphologies and functional properties required in the diverse organs and tissues of the body? We now understand that this process occurs through the selective expression of distinct subsets of the many thousands of genes in any animal’s genome in different cells. How genes are turned on and off in different cells over the course of animal development is an exquisitely orchestrated regulatory program whose features are only now coming into detailed view.

If morphological diversity is all about development, and development results from genetic regulatory programs, then is the evolution of diversity directly related to the evolution of genetic regulatory programs? Simply put, yes. But to understand how diversity evolves, we must first understand the genetic regulatory mechanisms that operate in development. In other words, what is the genetic toolkit of development and how does it operate to build animals? In the next two chapters, we will examine some of the general features of the genetic and regulatory logic of animal development.
SELECTED READINGS

The Early Fossil Record


Molecular Clocks and Animal Evolution


Animal Phylogeny


Animal Design and Evolution: Modularity and Homology


The foremost challenge for embryology has been to identify the genes and proteins that control the development of animals from an egg into an adult. Early embryologists discovered that localized regions of embryos and tissues possess properties that have long-range effects on the formation and patterning of the primary body axes and appendages. Based on these discoveries, they postulated the existence of substances responsible for these activities. However, the search for such molecules proved fruitless until the relatively recent advent of genetic and molecular biological technologies. The most successful approach to understanding normal development has involved the isolation of single gene mutations that have discrete and often large-scale effects on body pattern.

In this chapter, we take an inventory of the essential genetic toolkit for animal development. We concentrate on genes first discovered in insects, where systematic screens for developmental genes were pioneered. Importantly, however, it turns out that related genes are present in many other animals. We describe how members of the genetic toolkit were identified and what kinds of gene products they encode. In addition, we illustrate the general correlation between these genes’ patterns of expression with the development of the morphological features they affect. Finally, we briefly survey their distribution and function in other animals.

Only a small fraction of all genes in any given animal constitute the toolkit that is devoted to the formation and patterning of the body plan and body parts. Two classes of gene products with the most global effects on development are of special interest: families of proteins called transcription factors that regulate the expression of many other genes during development, and members of signaling pathways that mediate short- and long-range interactions between cells. The expression of specific transcription factors and signaling proteins marks the location of many classically defined regions within the embryo. These proteins control the formation, identity, and patterning of most major features of animal design and diversity.

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W. Bateson
Material for the Study of Variation (1894)

. . . if the mystery that surrounds embryology is ever to come within our comprehension, we must . . . have recourse to other means than description of the passing show.

T.H. Morgan
Experimental Embryology (1927)
BEFORE THE TOOLKIT—ORGANIZERS, FIELDS, AND MORPHOGENS

Long before any genes or proteins affecting animal development were characterized, embryologists sought to identify the basic principles governing animal design. In their search, they focused on the large-scale organization of the primary body axes, the differentiation of various germ layers (ectoderm, mesoderm, and endoderm), and the polarity of structures such as appendages and insect segments. By manipulating embryos and embryonic tissues, primarily by transplantation and ablation, researchers discovered many important properties of developing embryos and tissues. Much of the fascination of embryology stems from the remarkable activities of discrete regions within developing embryos in organizing the formation of body axes and body parts. Furthermore, these classical concepts of embryonic organization present a very useful framework for considering how that organization can change during evolution. We will briefly review some of these experiments and ideas before addressing their genetic and molecular manifestations.

The first demonstration of organizers—regions of embryos or tissues that have long-range effects on the fate of surrounding tissues—was achieved by Mangold and Spemann in 1924. They transplanted the lip of the blastopore, the invagination where mesoderm and endoderm move inside the amphibian embryo, of a newt gastrula into another newt embryo and found that the transplanted tissue could induce a second complete body axis (Fig. 2.1a). The additional embryo induced was partly derived from the transplanted graft and partly derived from the host. The equivalent of the “Spemann organizer” in amphibians has been found in chick and mouse embryos, and it is now recognized to be a structure characteristic of all chordate embryos.

Other organizers with long-range effects on surrounding tissues have been identified in the developing vertebrate limb bud. Transplantation of a discrete patch of posterior tissue to an ectopic anterior site induces the formation of limb structures (digits, tendons, muscles) with mirror-image polarity to the normal anteroposterior order (Fig. 2.1b). By contrast, transplantation or removal of anterior tissue has no effect on limb development, suggesting that this posterior region of the limb bud, dubbed the zone of polarizing activity (ZPA), organizes anteroposterior (that is, the thumb-to-pinkie axis) polarity and limb formation.

Another organizer operates from the most distal tip of the limb bud, the apical ectodermal ridge (AER). Removal of this region truncates the limb and deletes distal elements (digits), whereas transplantation of the AER to an early limb bud can induce outgrowth of a duplicate limb (Fig. 2.1b).

One explanation for the long-range polarizing and inductive effects of the Spemann organizer, ZPA, and AER is that these tissues are sources of inducer molecules, or morphogens—that is, substances whose concentrations vary within a tissue and to which surrounding cells and tissues respond in a concentration-dependent manner. The response to a morphogen depends, then, on the distance of the responding tissue from the source. For example, if the ZPA is a source of a morphogen, then diffusion of this substance can establish a gradient of inducer concentration. Induction of different digit types depends on the morphogen concentration, with low levels of morphogen inducing anterior digits (thumb) and high levels inducing posterior digits (pinkie) (Fig. 2.1b).

Organizers have been demonstrated and morphogens postulated in insects as well as vertebrates. Ligature and cytoplasmic transplantation experiments first suggested that the anteroposterior axis of certain insect embryos is influenced by two organizing centers, one at
Chapter 2: The Genetic Toolkit for Development

Figure 2.1
Organizers in animal embryos

Transplantation and ablation experiments have been used to investigate the long-range organizing activities of embryonic tissues. (a) The Spemann organizer. The dorsal blastopore lip of an early amphibian embryo can induce a second embryonic axis and embryo when transplanted to the ventral region of a recipient embryo. (b) Limb organizers. The apical ectodermal ridge (AER) is required for formation of distal limb elements. Removal leads to loss of structures; transplantation to specific ectopic sites induces extra elements. The zone of polarizing activity (ZPA) organizes the anteroposterior pattern; transplantation to an ectopic site induces extra digits with reverse polarity. (c) Insect egg organizer. Ligation of the insect *Euscelis* embryo (marked by the gray line) early in development deletes the thorax and abdomen; later ligations leave more segments intact. However, transplantation of the posterior pole cytoplasm (marked by the black dot) into the anterior of a ligated embryo induces the formation of a complete embryo. This result demonstrates that the posterior cytoplasm has organizer activity. (d) Within insect segments, epithelial polarity is organized by signaling sources. Ablation of a segment boundary (indicated by the interruption of the black line) reorganizes segment polarity (indicated by the orientation of small black hairs).

each pole of the egg (Fig. 2.1c), that behave as sources of morphogens. Similarly, the polarity of cells within insect segments appears to be organized by signals that produce a graded pattern (Fig. 2.1d).

One difficulty with this picture of morphogen-producing organizers arises when we attempt to explain the boundaries of their range of influence. All of the cells in a growing embryo are in contact with other cells, so how is it that some parts respond and others do not? One explanation involves the concept of the morphogenetic field. Early embryologists demonstrated that some parts of developing animals, such as the forelimb field, could be transplanted to another site and still differentiate properly—that is, into a forelimb. In addition, if undetermined cells were introduced into the field, they could become incorporated into the limb. These transplantable, self-regulating fields are discrete physical units or modules of embryonic development. They form bounded domains within which specific programs of morphogenesis occur. The term “primary field” applies to the entire embryo before the axes are determined; the limbs, eyes, and other organs are termed “secondary fields,” or organ primordia.

Secondary fields may be further subdivided into “tertiary fields,” defined by physical or developmental boundaries. Compartments are one special type of subdivision. First demonstrated within the wing imaginal disc of the fruit fly Drosophila melanogaster, compartments are composed of populations of cells that do not intermix with cells outside the compartment.

Further progress in understanding the nature of organizers, morphogens, and fields stalled after their discovery and description in the first half of the 1900s. The impasse was ultimately broken by the discovery of genes whose products governed the activity of organizers, behaved as morphogens, and controlled the formation and identity of embryonic fields. These genes make up the “toolkit” for animal development.

THE GENETIC TOOLKIT

Animal genomes contain thousands of genes. Many of these genes encode proteins that function in essential processes in all cells in the body (for example, metabolism, biosynthesis of macromolecules) and are often referred to as “housekeeping genes.” Other genes encode proteins that carry out specialized functions in particular cells or tissues within the body (for example, oxygen transport, immune defense) or, to extend the housekeeping metaphor, in specific “rooms” in the “house.” But here we are interested in a different set of genes, those whose products govern the construction of the house—the toolkit that determines the overall body plan and the number, identity, and pattern of body parts.

Toolkit genes have generally first been identified based on the catastrophes or monstrosities that arise when they are mutated. Two sources of toolkit gene mutations exist. The first source comprises rare, spontaneous mutations that arise in laboratory populations of model animals (for example, fruit flies, mice). The second source consists of mutations induced at random by treatment with mutagens (such as chemicals or radiation) that greatly increase the frequency of damaged genes throughout the genome. Elegant refinements of the latter approach, particularly in Drosophila melanogaster, have enabled systematic searches for members of the genetic toolkit for animal development.

Intensive screens for genes that affect the formation of the insect embryonic and adult body patterns and analysis of the structure, function, and expression of the proteins they encode have revealed several critical features of the genetic toolkit for development:
1. The toolkit is composed of a small fraction of all genes. Only a small subset of the entire complement of genes in the genome affects development in discrete ways.

2. Most toolkit genes encode either transcription factors or components of signaling pathways. Therefore, toolkit genes generally act, directly or indirectly, to control the expression of other genes.

3. The spatial and temporal expression of toolkit genes is often closely correlated with the regions of the animal in which the genes function.

4. Toolkit genes can be classified according to the phenotypes caused by their mutation. Similar mutant phenotypes often reflect genes that function in a single developmental pathway. Distinct pathways exist for the generation of body axes, for example, and for the formation and identity of fields.

5. Many toolkit genes are widely conserved among different animal phyla.

Because the discovery of the insect toolkit has offered a direct path to identifying developmental genes in other animals, we will begin our inventory of the genetic toolkit for animal development by considering *Drosophila melanogaster*.

**The Drosophila toolkit**

Classifying genes according to their developmental function

Many mutations have been isolated that alter the embryonic and/or adult body pattern of *Drosophila*. It has proved very useful to group the genes affected by these mutations into several categories based on the nature of mutant phenotypes. Most toolkit genes can be classified according to their function in controlling the identity of fields (for example, different segments and appendages), the formation of fields (for example, organs and appendages), the formation of cell types (for example, muscle and neural cells), and the specification of the primary body axes.

We begin by considering the genes that control the identity of segments and appendages. This choice is made partly for historical reasons and partly to follow a hierarchical approach. The genes controlling field identity were among the very first toolkit members discovered, and their identification inspired much of the genetic and molecular biological innovations that catalyzed the discovery of the rest of the toolkit. In addition, they are among the most globally acting developmental genes that affect animal form. Next, we discuss genes that control the patterning of fields at progressively finer scales, from the formation of entire fields, to compartments within fields, and then to differentiated cell types.

Homeotic genes and segmental identity

Among the most fascinating kinds of abnormalities to be described in animals are those in which one normal body part is replaced with another. Bateson catalogued several oddities of this nature, coining the term homeotic to describe such transformations. Among the most common homeotic variants noted by Bateson were arthropods in which one type of appendage formed in the position of another, such as a leg in place of an antenna (Fig. 2.2a), and vertebrates in which one type of vertebra or rib replaced another, such as a thoracic vertebra in place of a cervical vertebra (Fig. 2.2b).
Intriguing as Bateson’s specimens were, most were one-of-a-kind museum pieces in which only one member of a bilateral pair of structures was affected. To carry out a thorough investigation of the phenomenon of homeosis and its genetics, researchers required mutants that would breed true in subsequent generations. In 1915, Calvin Bridges isolated a spontaneous mutation in *Drosophila*, dubbed *bithorax*, in which part of the haltere (the posterior flight appendage in flies) was transformed into wing tissue. The haltere and wing are serially homologous appendages, so the *bithorax* mutation causes the partial transformation of the identity of a structure on the third thoracic segment (the haltere) into its serial homolog found on the second thoracic segment (the wing). A more complete transformation of the entire haltere into a wing can occur if additional mutations are combined with *bithorax*, producing a four-winged fly (Fig. 2.3).

In the following decades, several more homeotic mutants were identified in *Drosophila*, and in other insects as well. All of these homeotic mutations transform the identities of segments and their associated structures into those of other segments. For example, certain *Antennapedia* mutations cause the transformation of antennae into legs (Fig. 2.3), which are...
also serial homologs. The direction of the homeotic transformations depends on whether a mutation causes a loss of homeotic gene function where the gene normally acts, or a gain of homeotic gene function in places where the homeotic gene does not normally act. For example, \textit{Ultrabithorax} (\textit{Ubx}) acts in the haltere to promote haltere development and repress wing development. Loss-of-function mutations in \textit{Ubx} transform the haltere into a wing. Dominant mutations that cause \textit{Ubx} to gain function in the wing transform that structure into a haltere. Similarly, the antenna-to-leg transformations of \textit{Antennapedia} mutants reflect a dominant gain of \textit{Antennapedia} gene function in the antenna.

The fascination with homeotic mutants stems from two issues. First, it is startling that a single gene mutation could change entire developmental pathways so dramatically in a complex animal. Second, it is curious that the structure formed in the mutant is a well-developed likeness of another body part.

More detailed understanding of homeotic gene function was made possible by some particularly ingenious methods for analyzing the effects of mutations on the behavior of a group of cells in otherwise normal (or “wild-type”) tissues. That is, rather than being limited to examining the effect of homeotic mutations on whole animals, the behavior of clones of mutant cells could be observed within otherwise normal animals (Fig. 2.4). This technique...
was used to determine that the effects of homeotic mutations generally remain limited to cells with mutant genotypes; such behavior is termed **cell autonomous**. Thus a patch of cells in the haltere that lacks *Ubx* function forms wing tissue, even when it is surrounded by normal haltere cells (Fig. 2.4). This finding suggested that homeotic genes act within cells to select their developmental fate. Homeotic genes, and other genes with analogous functions in controlling cell fate, are therefore known as **selector genes**.

Although homeotic genes were first identified through spontaneous mutations affecting adult flies, they are required throughout most of *Drosophila* development to determine segmental identity. Systematic screening for homeotic genes led to the identification of eight linked genes, collectively referred to as **Hox genes**, that affect the specification of particular segment identities in the developing *Drosophila* embryo, larva, and adult. In addition to *Ultrabithorax* (*Ubx*) and *Antennapedia* (*Antp*), they include *labial* (*lab*), *proboscipedia* (*pb*), *Deformed* (*Dfd*), *Sex combs reduced* (*Scr*), *abdominal-A* (*abd-A*), and *Abdominal-B* (*Abd-B*). Generally, the complete loss of any *Hox* gene function causes transformations of segmental identity and is lethal in early development. The spontaneous homeotic mutants found in viable adults are caused by partial loss of gene function or are dominant such that in heterozygotes normal gene function is provided by the wild-type allele.

One of the most intriguing features of these *Hox* genes is that they are linked in two **gene complexes** in *Drosophila*, the Bithorax and Antennapedia Complexes; each complex contains several distinct homeotic genes. Furthermore, the order of the genes on the chromosome and within the two complexes corresponds to the rostral (head) to caudal (rear) order of the segments that they influence, a relationship described as **colinearity** (Fig. 2.5).
Figure 2.5

**The Hox genes of Drosophila**

Eight Hox genes regulate the identity of regions within the adult and embryo. The color coding represents the segments and structures that are affected by mutations in the various Hox genes.

**Figure 2.6**

Methods for visualizing gene expression in developing animals

The two most common means of visualizing where a gene is transcribed and its protein product is synthesized are (left) **in situ** hybridization of complementary RNA probe to mRNA and (right) immunolocalization of protein expression. The procedures for each method are indicated. Gene expression patterns are visualized as the product of enzymatic reactions (left) or with fluorescently labeled compounds (right).
The relationship between the structure of Hox gene complexes and the phenotypes of Hox mutants was illuminated by the molecular characterization of both the Bithorax Complex and the Antennapedia Complex. Cloning of the Hox genes provided the means to uncover when and where each of the eight genes is expressed during development. The ability to visualize Hox and other gene expression patterns during development was crucial to understanding the correlation between gene function and phenotypes. Localization of Hox genes’ RNA transcripts by in situ hybridization or of Hox proteins via immunological methods (Fig. 2.6) revealed that all Hox genes are expressed in spatially restricted, sometimes overlapping domains within the embryo. These genes are also expressed in subsets of the developing larval imaginal discs, which proliferate during larval development and differentiate during the pupal stages to give rise to the adult fly.

The patterns of Hox gene expression generally correlate with the regions of the animal affected by homeotic mutations. For example, the Ubx gene is expressed within the posterior thoracic and most anterior abdominal segments of the embryo (Fig. 2.7a). The development of these segments is altered in Ubx mutants. In larvae, Ubx is expressed in the developing haltere, but not in the developing wing (Fig. 2.7b–e). This expression correlates with the requirement for Ubx to promote haltere development and to suppress wing identity.

The boundaries of Hox gene expression in the Drosophila embryo are not segmental, but usually begin in the posterior part of one segment and extend to (or beyond) the anterior portion of the next-most-posterior segment, a unit dubbed a parasegment. In the various imaginal tissues of the developing adult, homeotic genes are often expressed in segmental domains. For example, flies have three pairs of legs, with one pair extending from each of the three thoracic segments. Each pair of adult legs has a distinctive morphology. Indeed, genetic analysis has shown that the morphology of the first legs is largely influenced by the Scr gene, the second legs by the Antp gene, and the third legs by the Ubx gene. These respective genetic requirements correlate with the respective patterns of homeotic gene expression in the developing imaginal legs.

It is crucial to understand the distinction between Hox gene function in determining the identity of a field, as opposed to a requirement for Hox gene function in the formation of the field. The antennae, mouthparts, and walking limbs of flies all develop from serially homologous limb fields. In the absence of homeotic genes, each limb field develops, but with antennal identity. Therefore, Hox genes specify the particular identity, but are not required for the formation of the limb fields. The expression and function of Hox genes are not limited to body segments and their appendages. These genes act as region-specific selectors in all three germ layers (ectoderm, mesoderm, and endoderm) and in diverse structures and tissues.

The homeobox

The large effects of Hox genes on the developmental fates of entire segments and structures made the nature of the proteins encoded by these genes of special interest. The close genetic linkage and similar function of Drosophila Hox genes suggested that they might have evolved through the tandem duplication of one or more ancestral Hox genes. This idea led to the discovery that the DNA sequences of the Hox genes of the Bithorax and Antennapedia Complexes were similar enough to hybridize to each other. This similarity was traced to a 180 base-pair (bp) stretch of DNA, dubbed the homeobox, that encodes a 60 amino acid protein domain (the homeodomain); the sequence of the homeodomain is very similar among the
homeotic proteins (Fig. 2.8). The structure of the homeodomain resembles the DNA-binding domain of many prokaryotic regulatory proteins, suggesting that homeotic gene products exert their effects by controlling gene expression during development and that the homeodomain binds to DNA in a sequence-specific manner.

The homeobox gene family is large and diverse. In fact, the homeodomain motif is found in approximately 20 other distinct families of homeobox-containing genes, all of which encode DNA-binding proteins.
Another class of selector genes acts within specific developing fields to regulate the formation and/or the patterning of entire structures. Such genes have been identified in *Drosophila* through spontaneous or induced mutations that selectively abolish or reduce the development of the eye, wing, limbs, and heart. Several of these genes also have the remarkable ability to induce the formation of ectopic organs when they are expressed at different sites in the animal.

Perhaps the best-known *Drosophila* field-specific selector gene is the *eyeless* (*ey*) gene. Flies that lack *ey* function can reach adulthood, but never develop a compound eye (Fig. 2.9a,b). Molecular characterization of the *ey* gene revealed that it encodes a member of a particular homeobox gene family (*Pax6*), suggesting that the Ey protein acts as a DNA-binding transcription factor to regulate the expression of other genes.

The *ey* gene is expressed in the developing eye field in the embryo, and in the larval eye imaginal disc, before the formation of the units (ommatidia) that make up the compound fly eye (Fig. 2.10b). Most remarkable, however, is the ability of the *ey* gene when expressed elsewhere in the developing fly, such as in the imaginal wing or leg discs, to induce the formation of eye tissue composed of properly organized, pigmented ommatidia (Fig. 2.9c,d). Its ability to reprogram other developing tissues to form eyes suggests that *ey* is a major regulatory gene in the genetic program of eye development.

The *Distal-less* (*Dll*) gene displays similar properties with respect to the formation of *Drosophila* limbs. Named for the effect of its mutations on the formation of the proximodistal axis of the limbs, *Dll* affects the development of all limbs, including the walking legs, mouthparts, antenna, and genitalia. Complete loss of *Dll* function truncates all limbs (that is, the limbs lack distal elements). The *Dll* gene is yet another type of homeobox-containing gene, suggesting that *Dll* also exerts its effects by regulating the expression of other genes. It is expressed in the limb primordia in the embryo, and in the distal portion of all imaginal limb fields.
Expressing \( Dll \) in places where it is not normally active can induce the outgrowth of ectopic limbs.

Development of the flight appendages of \textit{Drosophila} depends on the function of a pair of genes, \textit{vestigial} (\( vg \)) and \textit{scalloped} (\( sd \)), whose products act together in a molecular complex. Fruit flies with \( vg \) and \( sd \) mutations lack wings and halteres altogether. The Vg and Sd proteins are expressed in the wing and haltere primordia in the embryo and in fields of cells within the imaginal discs that will give rise to the flight appendages (see Fig. 2.10c). As is the case with the other field-specific selector genes, expression of \( vg \) with \( sd \) in developing eyes, legs, antenna, or genitalia can induce the formation of wing tissue. The Vg and Sd proteins form a

(Fig. 2.10c,d). Expressing \textit{Dll} in places where it is not normally active can induce the outgrowth of ectopic limbs.

Figure 2.9

\textbf{The eyeless selector gene controls eye development}

(a) Normal fly head with eye. (b) The ey mutant fly lacks the eye. (c) Expression of the \textit{ey} gene induces the formation of pigmented eye tissue at new sites, including (d) on the wing.

Source: Photographs courtesy of Georg Halder.
Figure 2.10
Field-specific selector genes

(a) Development of parts of the Drosophila adult depend upon the function of the ey (eyes), vg (flight appendages), and Dll (limbs) selector genes. (b–d) These genes are expressed in both the embryonic primordia (left) and larval imaginal discs (right), which will give rise to these structures.

Source: Photomicrographs courtesy of Georg Halder and Grace Panganiban.
complex that binds to DNA, indicating that their selector function is mediated by regulation of gene expression.

The formation of the *Drosophila* heart depends on still another selector gene, dubbed *tinman* (*tin*). Mutants lacking *tin* function lack a heart. The *tin* gene is expressed in the developing mesoderm and in all cells that will form the cardiac tissue of the fly. It is a member of a distinct homeobox family, and thus also a DNA-binding protein that acts by controlling gene expression.

**Compartment selector genes**

Several genes have been identified in *Drosophila* that act within certain developing fields to subdivide them into separate cell populations, or compartments. The *engrailed* (*en*) gene acts in the posterior part of all segments of the embryo; it is expressed continuously such that the posterior portions of all structures that develop from these segments also express *en* (Fig. 2.11a). The function of the *engrailed* gene is best understood in the embryo and in the developing wing, where it acts to determine posterior identity. Mutations in this gene cause posterior cells to develop as anterior cells but with reversed segmental polarity, resulting in mirror-image duplications of anterior tissue. The *engrailed* gene encodes member of a distinct class of homeodomain-containing transcription factors.

A second compartmental selector gene, *apterous* (*ap*), subdivides the developing wing imaginal disc into dorsal and ventral compartments (Fig. 2.11b–e). Complete loss of *apterous* function blocks wing development, whereas loss of *apterous* function within a subset of dorsal cells transforms their identity to ventral fate. The Apterous protein belongs to yet another class of homeodomain-containing transcription factors.

**Cell-type-specific selector genes**

Another class of selector genes operates within developing fields to control the differentiation of particular cell types. The formation of neuroblasts and other neural precursor cells in *Drosophila* requires the action of members of the Achaete-Scute Complex (AS-C), a gene complex that contains four genes. Loss of AS-C gene function in the embryo prevents formation of the nervous system; loss or reduction of individual AS-C gene functions in particular body regions in the imaginal tissues of the developing adult fly causes loss of particular sensory bristles.

All four AS-C genes encode structurally related transcription factors. The genes are expressed in dynamic and complex patterns that foreshadow the formation of central and peripheral nervous system elements in the larva and adult. The development of neural precursors is initiated within clusters of cells that express AS-C genes, from which a single precursor segregates, divides, and gives rise to neurons and associated cells (Fig. 2.12). A similar process involving a distantly related group of transcription factors specifies muscle development in *Drosophila*. The *twist*, *nautilus*, and *Dmef-2* genes control the development and differentiation of muscle cells.

**Formation of the body axes**

**Systematic searches for developmental genes in Drosophila**

Many of the selector genes described in the previous section were first identified on the basis of the adult phenotypes of spontaneous mutants in *Drosophila*. Most of those mutations,
however, did not completely disrupt the gene’s function during development. Complete loss of function of many selector genes is lethal at earlier stages of development. Therefore, to find genes that control other aspects of embryo organization and patterning, genetic screens had to be designed that could identify recessive lethal mutations.
Two types of systematic screens have harvested the lion’s share of the *Drosophila* genetic toolkit. The first searched for all loci that were required in the fertilized egg, or *zygote*, for proper patterning of the larva. The second type of search sought to identify those genes whose products function in the egg for proper patterning, before the zygotic genome becomes active. Genes whose products are provided by the female to the egg are called *maternal effect* genes. Mutant phenotypes of strict maternal effect genes depend only on the genotype of the female parent (Fig. 2.13).

The two types of systematic mutagenesis screens revealed that mutations in only a small fraction of all genes in the genome have very specific effects on the organization and patterning of the embryo and larva. In addition to their maternal or zygotic actions, these mutants and the corresponding genetic loci can be classified according to the embryonic axis affected (anteroposterior or dorsoventral), and the type of patterning defect observed. Molecular characterization of these genes identified many of the first known representatives of widely shared transcription factor families and signaling pathways. Indeed, the molecular analysis of many of these genes in *Drosophila* led to the development of tools to isolate them.
from other animals. Thus the systematic inventory of the *Drosophila* genome is representative of what is generally known about the types of molecules with large-scale effects on animal patterning.

**The anteroposterior axis**

A few dozen *Drosophila* genes are required for proper anteroposterior patterning of the embryo and larva. These genes are grouped into five classes based on their realm of influence on embryonic pattern. Each class represents a progressively finer subdivision of the developing embryo.

The first class consists of the maternal effect genes, such as the *bicoid* gene (which affects the anterior region of the embryo) and the *nanos* and *caudal* genes (which affect the posterior region) (Fig. 2.14).

The second class contains the zygotically active *gap genes*, which include the *bunchback*, *Krüppel*, *giant*, *knirps*, *tailless*, and *buckebein* genes, each of which regulates the formation of a contiguous set of segments. Mutations in gap genes lead to gaps in segmentation (Fig. 2.15a,b).

The third class comprises the *pair-rule genes*, such as *fushi tarazu*, *even-skipped*, *bairy*, and *paired*, which act at a double-segment periodicity. Pair-rule mutants display defects in part of each pair of segments (Fig. 2.15c).
The fourth class consists of the segment polarity genes, which affect patterning within each segment. Mutants in this class lack the normal polarity of pattern elements within segments and display polarity reversals and segmentation defects (Fig. 2.15d). The products of two segment polarity genes, wingless and hedgehog, are largely responsible for the polarity-organizing activities identified within insect segments by classical transplantation and ablation techniques.

The fifth class includes the homeotic selector genes, which we discussed earlier in this chapter.

Collectively, these five classes of mutants indicate that the segmental body plan of the Drosophila larva is progressively specified by genes acting over the realm of the whole embryo, in subregions of the embryo, in every other segment, and, ultimately, in each individual segment.

**The dorsoventral axis**

The dorsoventral axis of the Drosophila embryo is also regionally subdivided. Several maternal effect genes are required to initiate the establishment of dorsoventral polarity. Embryos from
females that lack the activity of any of these genes, such as dorsal, are “dorsalized”—that is, ventral structures do not form in these animals. The zygotically active genes decapentaplegic (dpp), zerknüllt (zen), short gastrulation (sog), twist (twi), and snail (sna) all play major roles in the subdivision of the dorsoventral axis. In addition, neurogenic genes, such as Delta (Dl) and Notch (N), are required in distinct dorsoventral subregions for the formation of the ectoderm.
Expression of toolkit genes

The phenotypes of mutant fruit flies tell only part of the story regarding what developmental genes do. To fully understand the link between these genes and patterning, we must know the timing and location of the genes’ expression patterns and the molecular nature of the genes’ protein products. The identification of gene classes that, when mutated, cause similar effects in development raises the possibility that a given class of genes may affect the same developmental process or genetic pathway, or that the genes may encode products with similar functions.

Analysis of the expression of toolkit genes has revealed a very informative correlation between the locations at which genes are expressed in development and the pattern of defects caused by mutations. For each of the five classes of anteroposterior axis-patterning genes, a clear correspondence exists between the regions of the embryo in which the gene is transcribed (or the protein product is localized) and the regions affected by mutations in that gene. For example, the bicoid and nanos proteins are expressed in graded patterns emanating from the anterior and posterior poles of the Drosophila embryo, respectively (Fig. 2.16). These proteins are largely responsible for the organizing activities that classical experiments identified as residing at the two poles of the insect egg.

The gap genes are expressed in blocks of cells that correspond to the future positions of the segments affected by gap gene mutants (Fig. 2.17a). The pair-rule genes are expressed in one transverse stripe per every two segments, for a total of seven stripes that span 14 future body segments; the stripes correspond to the periodicity of defects in mutant embryos (Fig. 2.17b). The segment polarity genes are expressed in each segment, in 14 or 15 transverse stripes (Fig. 2.17c). The various dorsoventral patterning genes are expressed in different
domains along the dorsoventral axis that correspond to regions of the embryo that give rise to elements such as the mesoderm, neuroectoderm (the part of the ectoderm from which the ventral nervous system develops), and amnioserosa (a dorsal sheet of extraembryonic cells) (Fig. 2.18).

**Toolkit gene products: transcription factors and signaling pathway components**

The proteins encoded by selector and axial patterning genes most often belong to one of two categories: transcription factors or components of signaling pathways. Ultimately, these proteins exert their effect through the control of gene expression. Thus developmental
processes such as embryonic axis formation and segmentation are organized by regulating gene expression in discrete regions and cell populations of the embryo.

The transcription factors found in the *Drosophila* toolkit include representatives of most of the known families of sequence-specific DNA-binding proteins. These families are distinguished by the type of secondary structures in the folded protein that are involved in protein subunit interactions and contact with DNA.

Most transcription factors possess either a helix-turn-helix, zinc finger, leucine zipper, or helix-loop-helix (HLH) motif (Fig. 2.19; Table 2.1). The homeodomain superfamily belongs to the helix-turn-helix class of factors, for example (Fig. 2.19a). Three proteins with divergent homeodomain sequences—Bicoid, Fushi tarazu, and Zen—have very different roles in organizing the anteroposterior axis, segmentation, and dorsoventral axis patterning, respectively. All three proteins are also encoded within the Antennapedia Complex, surrounded by *Hox* genes. Most other homeodomain proteins are encoded by genes dispersed throughout the genome.
Figure 2.19

Structural motifs of major transcription factor families

(a) The second and third α-helices of the homeodomain form a helix-turn-helix structure. (b) The zinc finger motif involves a coordination complex of Zn²⁺ with critically positioned cysteine (C) and histidine (H) residues. The "finger" contacts DNA. (c) The leucine zipper structure is formed by association of two subunits with regularly spaced leucine residues. (d) The helix-loop-helix structure is similar to the leucine zipper except that a protein loop interrupts the helices and the association of the subunits.
### TABLE 2.1  **Transcription factors in the Drosophila genetic toolkit for development, and mouse homologs**

<table>
<thead>
<tr>
<th>Domain</th>
<th>Drosophila gene</th>
<th>Developmental function</th>
<th>Mouse homolog(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Homeodomain</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Homeodomain</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zinc finger</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th></th>
<th>Drosophila gene</th>
<th>Developmental function</th>
<th>Mouse homolog(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>labial</td>
<td>Homeotic</td>
<td>Hox a1, b1, c1</td>
</tr>
<tr>
<td></td>
<td>proboscipedia</td>
<td>Homeotic</td>
<td>Hox a2, b2</td>
</tr>
<tr>
<td></td>
<td>deformed</td>
<td>Homeotic</td>
<td>Hox a4, b4, c4, d4</td>
</tr>
<tr>
<td></td>
<td>Sex combs reduced</td>
<td>Homeotic</td>
<td>Hox a5, b5, c5</td>
</tr>
<tr>
<td></td>
<td>Antennapedia</td>
<td>Homeotic</td>
<td>Most similar to Hox6-8</td>
</tr>
<tr>
<td></td>
<td>Ultrabithorax</td>
<td>Homeotic</td>
<td>Most similar to Hox6-8</td>
</tr>
<tr>
<td></td>
<td>abdominal-A</td>
<td>Homeotic</td>
<td>Most similar to Hox6-8</td>
</tr>
<tr>
<td></td>
<td>Abdominal-B</td>
<td>Homeotic</td>
<td>Most similar to Hox9-13</td>
</tr>
<tr>
<td></td>
<td>bicoic</td>
<td>Maternal anteroposterior axis organizer</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>caudal</td>
<td>Maternal/zygotic anteroposterior axis organizer</td>
<td>Cdx genes</td>
</tr>
<tr>
<td></td>
<td>fushi-tarazu</td>
<td>Pair-rule segmentation</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>even skipped</td>
<td>Pair-rule segmentation</td>
<td>evx genes</td>
</tr>
<tr>
<td></td>
<td>paired</td>
<td>Pair-rule segmentation</td>
<td>Pax3</td>
</tr>
<tr>
<td></td>
<td>zerknultt</td>
<td>Dorsoventral axis patterning</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>engrailed</td>
<td>Segment polarity/posterior compartment selector</td>
<td>En genes</td>
</tr>
<tr>
<td></td>
<td>apterous</td>
<td>Dorsal wing compartment selector</td>
<td>Lmx genes</td>
</tr>
<tr>
<td></td>
<td>eyeless</td>
<td>Eye field selector</td>
<td>Pax6</td>
</tr>
<tr>
<td></td>
<td>Distal-less</td>
<td>Limb field selector</td>
<td>Dlx genes</td>
</tr>
<tr>
<td></td>
<td>tinman</td>
<td>Mesoderm, heart selector</td>
<td>Nkx2.x genes</td>
</tr>
<tr>
<td></td>
<td>extradenticle</td>
<td>Hox co-factor</td>
<td>Pbx genes</td>
</tr>
<tr>
<td></td>
<td>hunchback</td>
<td>Gap segmentation</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Krüppel</td>
<td>Gap segmentation</td>
<td>KLF genes</td>
</tr>
<tr>
<td></td>
<td>tailless</td>
<td>Gap segmentation</td>
<td>tlx</td>
</tr>
<tr>
<td></td>
<td>huckebein</td>
<td>Gap segmentation</td>
<td>Sp1/Egr genes (Krox20)</td>
</tr>
<tr>
<td></td>
<td>knirps</td>
<td>Gap segmentation (steroid receptor)</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>odd-skipped</td>
<td>Pair-rule segmentation</td>
<td>Osr genes</td>
</tr>
<tr>
<td></td>
<td>snail</td>
<td>Dorsoventral axis patterning</td>
<td>Snail, slug</td>
</tr>
</tbody>
</table>
Several of the gap genes encode zinc finger proteins, whereas genes such as the dorsoventral patterning gene \textit{twist} and the pair-rule segmentation gene \textit{hairy} encode basic HLH proteins. The identification of these DNA-binding motifs in proteins has allowed biologists to deduce the biochemical function of a gene product by inspecting its encoded sequence, rather than resorting to exhaustive biochemical analysis. Furthermore, because these motifs are involved in contact with DNA, they are often constrained with respect to evolutionary changes in their sequence. As a result, these highly conserved motifs are useful for isolating gene homologs in other taxa.

The second major category of toolkit genes encode proteins involved in the process of cell signaling, either as ligands, receptors for ligands, or components involved in the intracellular transduction of signals. At least seven major signaling pathways operate in the \textit{Drosophila} embryo: the \textit{Hedgehog}, \textit{Notch}, \textit{Wingless}, \textit{Dpp}/transforming growth factor-\(\beta\) (TGF-\(\beta\)), \textit{Toll}, epidermal growth factor (EGF), and fibroblast growth factor (FGF) signaling pathways (Table 2.2).

All of these pathways have similar elements—namely, each has at least one signaling ligand, at least one receptor spanning the cell membrane, and at least one DNA-binding transcription factor that responds to signaling inputs by binding to target genes to turn them on or off (Fig. 2.20). Although the signaling logic may be similar among pathways, the biochemistry is not. Many structural types of ligands and receptors exist, and each pathway regulates the activity of different transcription factors. The response to ligand binding is mediated by a variety of mechanisms that often involve post-translation modifications (for example, protein phosphorylation, proteolysis, binding to a cofactor) that regulate the activity of, or the translocation of, transcription factors to the cell nucleus (Fig. 2.20). The similar mutant phenotypes of some developmental genes reflect their involvement in the same signaling pathway. Any loss of a signaling ligand, receptor, or member of the signal transduction machinery can cause the pathway to fail.

<table>
<thead>
<tr>
<th>Domain</th>
<th>Drosophila gene</th>
<th>Developmental function</th>
<th>Mouse homolog(s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Helix-loop-helix</td>
<td>Achaete-Scute Complex genes</td>
<td>Proneural</td>
<td>\textit{MASH} genes</td>
</tr>
<tr>
<td>nautilus</td>
<td>Myogenic</td>
<td>\textit{myoD}</td>
<td></td>
</tr>
<tr>
<td>Dmef</td>
<td>Myogenic</td>
<td>\textit{mef}</td>
<td></td>
</tr>
<tr>
<td>hairy</td>
<td>Pair-rule segmentation</td>
<td>\textit{hairy}</td>
<td></td>
</tr>
<tr>
<td>twist</td>
<td>Dorsoventral axis patterning</td>
<td>\textit{twist}</td>
<td></td>
</tr>
<tr>
<td>Other</td>
<td>dorsal</td>
<td>Dorsoventral axis organizer</td>
<td>\textit{NF-(\kappa)/rel} genes</td>
</tr>
<tr>
<td>giant</td>
<td>Gap segmentation</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>vestigial</td>
<td>Wing selector</td>
<td>\textit{tondu}</td>
<td></td>
</tr>
<tr>
<td>scolopced</td>
<td>Wing selector</td>
<td>\textit{TEF-1}</td>
<td></td>
</tr>
</tbody>
</table>

Table 2.1 | Continued

**TABLE 2.2** Major signaling pathways in the *Drosophila* and mouse genetic toolkits

<table>
<thead>
<tr>
<th>Pathway</th>
<th>Components</th>
<th><em>Drosophila</em> toolkit</th>
<th>Mouse toolkit</th>
</tr>
</thead>
<tbody>
<tr>
<td>TGF-β</td>
<td>Ligands</td>
<td>decapentaplegic, 60A, screw</td>
<td>BMP genes, TGF-β, Activin</td>
</tr>
<tr>
<td></td>
<td>Receptors</td>
<td>thick veins, Punt, saxophone</td>
<td>BRK genes, ALK genes</td>
</tr>
<tr>
<td></td>
<td>Transcription factors</td>
<td>Mad, Medea</td>
<td>Smad genes</td>
</tr>
<tr>
<td>Wingless</td>
<td>Ligands</td>
<td>wingless, Dwnt genes</td>
<td>Wnt genes</td>
</tr>
<tr>
<td></td>
<td>Receptors</td>
<td>frizzled genes, arrow</td>
<td>frizzled genes, LRP-5/6</td>
</tr>
<tr>
<td></td>
<td>Transcription factors</td>
<td>DTCF/pangolin</td>
<td>TCF genes, LEF1</td>
</tr>
<tr>
<td>Notch</td>
<td>Ligands</td>
<td>Delta, Serrate</td>
<td>Dil (Delta-like) genes, Jagged/serrate genes</td>
</tr>
<tr>
<td></td>
<td>Receptors</td>
<td>Notch</td>
<td>Notch genes</td>
</tr>
<tr>
<td></td>
<td>Transcription factors</td>
<td>Suppressor of Hairless</td>
<td>RBP-Jk/CBF1</td>
</tr>
<tr>
<td>Hedgehog</td>
<td>Ligands</td>
<td>hedgehog</td>
<td>Sonic hedgehog, Indian hedgehog, Desert hedgehog</td>
</tr>
<tr>
<td></td>
<td>Receptors</td>
<td>Patched, smoothened</td>
<td>Patched genes, smoothened</td>
</tr>
<tr>
<td></td>
<td>Transcription factors</td>
<td>cubitus interruptus</td>
<td>Gli genes</td>
</tr>
<tr>
<td>Toll</td>
<td>Ligands</td>
<td>spatzle</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Receptors</td>
<td>Toll</td>
<td>TLR genes</td>
</tr>
<tr>
<td></td>
<td>Transcription factors</td>
<td>dorsal</td>
<td>NF-κB/rel genes</td>
</tr>
<tr>
<td>EGF</td>
<td>Ligands</td>
<td>Spitz, Argos, gurken</td>
<td>TGFrα, EGF</td>
</tr>
<tr>
<td></td>
<td>Receptors</td>
<td>torpedo (EGF-R)</td>
<td>EGF-R</td>
</tr>
<tr>
<td></td>
<td>Transcription factors</td>
<td>pointed</td>
<td>ets genes</td>
</tr>
<tr>
<td>FGF</td>
<td>Ligands</td>
<td>branchless</td>
<td>FGF genes</td>
</tr>
<tr>
<td></td>
<td>Receptors</td>
<td>breathless, heartless</td>
<td>FGF-R genes</td>
</tr>
<tr>
<td></td>
<td>Transcription factors</td>
<td>Not known</td>
<td></td>
</tr>
</tbody>
</table>

Figure 2.20
A generic signaling pathway

Most signaling pathways operate through similar logic but have different proteins and signal transduction mechanisms. Signaling begins when membrane-bound receptors bind a ligand, leading to the release or activation of associated intracellular proteins. Receptor activation often leads to the modification of inactive transcription factors that are translocated to the cell nucleus, bind to cis-regulatory DNA sequences or to DNA-binding proteins, and regulate the level of target gene transcription.
Pleiotropy of toolkit genes

Because of the essential roles played by toolkit genes in the initial pattern of events in *Drosophila* development, flies with mutations in toolkit genes typically fail to survive the early stages of the life cycle. Most of these genes also play critical roles later in development, however. Toolkit genes with multiple functions during development are termed **pleiotropic**. For instance, individual genes may act in different developmental events in the same tissue over the course of development, in different tissues at the same stage of development, or in different tissues over the entire course of development. The multiple roles played by individual genes can be unmasked by mutations that affect only a subset of a gene’s functions, by conditional mutations (such as temperature-sensitive mutations), or by genetic techniques for investigating the requirements for gene function in subsets of cells at various stages of development.

Signaling pathway components are especially pleiotropic. For example, the *Wingless* pathway plays diverse roles in many tissues throughout *Drosophila* development. It is required early in the ectoderm to organize segment polarity; later in the embryonic ectoderm to direct the formation of the leg and wing imaginal discs; days later in the larval wing field to organize dorsoventral polarity, wing outgrowth, and sensory organ patterning; and finally to organize the polarity of the eye, leg, and other tissues. All components of the *Wingless* pathway are required in each of these settings, even though the ultimate regulatory and morphogenetic “output” of the pathway is different in each case. In Chapter 3, we will see how transcription factors and signaling pathways achieve their tissue-specific effects.

**SHARING OF THE GENETIC TOOLKIT AMONG ANIMALS**

**Hox genes**

One of the most exciting and unexpected discoveries that occurred soon after the cloning of the *Hox* genes was the detection of genes with related sequences in all sorts of animals. Using the homeobox to search for similar sequences in other genomes by hybridization, researchers isolated *Hox*-related genes from a broad sample of other animals. The similarity between the sequences of the homeodomains of genes isolated from frogs, mice, and humans and the original *Drosophila* *Hox* sequences was surprisingly extensive given the vast evolutionary distances between these animals. As many as 59 of the 60 amino acid residues were shared between the most similar homeodomains (Fig. 2.21).

An even greater surprise emerged with the physical mapping of vertebrate *Hox* genes. The map revealed that these *Hox* genes occurred in four large, linked complexes and that the order of the *Hox* genes within these complexes paralleled the order of their most related counterparts in the insect *Hox* complexes (Fig. 2.22). The vertebrate complexes define 13 groups of *Hox* genes, compared with the eight genes in *Drosophila*, although not every *Hox* gene is represented in each vertebrate complex (Fig. 2.22). Furthermore, the relative order of expression of vertebrate *Hox* genes along the anteroposterior (rostrocaudal) axis of vertebrate embryos correlates with gene position in each complex (Fig. 2.22).

With the invention of techniques for knocking out gene function in mice, it became possible to analyze the functions of the 39 *Hox* genes in the four mouse *Hox* complexes. This analysis has been complicated by **genetic redundancy**—that is, the expression and function of
Figure 2.21
The similarities of Drosophila and vertebrate Hox protein sequences

The sequence of the Drosophila Dfd homeodomain and C-terminal flanking region and the sequences of several members of the vertebrate Hox 4 genes are shown. Note the great sequence similarity between the Drosophila and vertebrate proteins, and among the vertebrate Hox proteins.

Figure 2.22
Hox gene complexes and expression in vertebrates

(a) In the mouse, four complexes of Hox genes, comprising 39 genes in all, occur on four different chromosomes. Not every gene is represented in each complex, however. (b) The Hox genes are expressed in distinct rostrocaudal domains of the mouse embryo.

two or more similar Hox genes in overlapping domains. In some cases, loss of function of a specific Hox gene causes the homeotic transformation of the identity of particular repeated structures, such as vertebrae, and, in other cases, the loss of particular organs (Fig. 2.23). Conversely, the expression of Hox genes in more anterior sites often causes the reciprocal transformations. Similar results have been obtained in birds, amphibians, and fish, which indicates that in vertebrates, as well as Drosophila, Hox genes act as region-specific selector genes.

Hox genes also affect the development of unsegmented animals. In the nematode Caenorhabditis elegans, for example, Hox genes regulate the differentiation of cell types and certain structures along the main body axis. As Hox genes have been found on all branches of the metazoan tree and play such important roles in body patterning, we will devote considerable attention to their evolution and function in later chapters.

**Field- and cell-type-specific selector genes**

The discovery of homologs of Drosophila Hox genes in vertebrates and other animal phyla inspired the search for homologs of other Drosophila selector and developmental genes.

---

**Figure 2.23**

*Hox* genes regulate vertebrate axial morphology

The morphologies of different regions of the vertebral column are regulated by Hox genes. (a) In the mouse, normally six lumbar vertebrae arise just anterior to the sacral vertebrae. (b) In mice lacking the function of the posteriorly acting Hoxd11 gene, and possessing one functional copy of the Hoxd11 gene, seven lumbar vertebrae form and one sacral vertebra is lost. (c) In mice lacking both Hoxa11 and Hoxd11 function, eight lumbar vertebrae form and two sacral vertebrae are lost. The anterior limit of Hoxd11 expression is at the first sacral vertebrae. Loss of these Hox gene functions transforms the sacral vertebrae into lumbar vertebrae.

Source: Photographs courtesy of Dr Anne Boulet, HHMI, University of Utah.
Through cross-hybridization between insect genes and vertebrate genes, families of vertebrate genes were isolated that included homologs of the *Distal-less* (*Dlx*), *tinman* (*Nkx2.5*), and AS-C (MASH) genes. Vertebrate homologs of *eyeless* (*Pax6*), *scalloped* (*TEF*-1), *apterous* (*Lmx*), *engrailed* (*En1* and *En2*), *nautilus* (*myoD*), and *Dmef* (*mef*), genes are also known (Table 2.1).

Thus homologs of most identified field- and cell-type-specific selector genes have been found in vertebrates. Of even greater interest, however, are the structures and cell types in which these vertebrate homologs function. For example, the vertebrate *eyeless* homolog, *Pax6*, is involved in the development of the vertebrate eye. Mutations that reduce the activity of the mouse *Pax6* gene, called *small eye* (*Sey*), result in loss of eye tissue, including loss of the retina, lens, and cornea in homozygous mutants that lack all *Sey* function (Fig. 2.24). Mutations in the human *Pax6* gene, *Aniridia*, similarly affect eye development. Furthermore, when a version of the *Sey* gene is introduced into and expressed in flies, it behaves just like the *eyeless* gene in terms of its ability to induce new eye tissue. The *Pax6* gene has been found to be associated with eye development across the metazoan tree, despite the differences in the architectures and optic principles of animal eyes.

Similar results have been found for vertebrate homologs of the *Distal-less*, *tinman*, and *achaete-scute* genes. In vertebrates and other phyla, *Distal-less*-related genes are expressed in an enormous variety of appendages with very distinct morphologies and functions.
Tinman/Nkx2.5 homologs are also expressed in, and required for, the development of the vertebrate heart. Similarly, AS-C homologs are expressed in neural precursors in vertebrates.

**Signaling pathways: classical organizers, and morphogens**

The components of all the major signaling pathways in *Drosophila* also have vertebrate counterparts (Table 2.2). These pathways operate in many tissues throughout vertebrate development. Importantly, some widely shared signaling proteins play important roles in the classical organizers defined in vertebrate embryos. For example, searches for molecules with activities associated with the Spemann organizer in the *Xenopus* embryo revealed that several of the secreted signaling proteins have potent inducing activities. One protein, dubbed Chordin because of its activity in inducing dorsal derivatives (the notochord), is a homolog of the *Drosophila* Short gastrulation protein. Both the vertebrate and *Drosophila* proteins interact with members of the TGF-β signaling protein family, and both are involved in dorsoventral axis formation.

Signaling proteins have also been found that account for the activities of the ZPA and AER in the vertebrate limb bud. For example, Sonic hedgehog (Shh), a homolog of the *Drosophila* Hedgehog signaling protein, is expressed in the posterior mesenchyme of the limb bud, precisely where the ZPA is localized (Fig. 2.25). Consistent with Shh carrying out the organizing
activity of the ZPA, expression of Shh in the anterior of the limb induces the same ectopic mirror-image duplications of digits as does transplantation of the ZPA. Similarly, members of the fibroblast growth factor signaling protein family are expressed in the distal tip of the limb bud in the AER and promote its activity (Fig. 2.25).

THE TOOLKIT AND ANIMAL DESIGN

The visualization of fields, compartments, and organizers as domains of selector gene expression and as sources of signaling proteins with demonstrable long-range effects on the patterning of embryonic fields has provided concrete molecular evidence of the fundamental roles that these units of organization play in embryonic development. The identification of genes that affect the formation and function of these units of animal body organization represents a first step toward understanding the developmental genetic logic underlying animal design. To understand how an animal is built, we need to understand which mechanisms establish these spatial domains in a growing embryo and how gene products function together to shape animal patterns.

The conservation of the genetic toolkit provokes many developmental and evolutionary questions. How do such different structures as the insect compound eye and the vertebrate lens-type eye develop when their formation is controlled by such similar, even functionally interchangeable, genes? And, what does the conservation of genes involved in building anatomically different structures with similar biological functions tell us about animal ancestors? We’ll confront the mechanistic issues in Chapter 3 and tackle the historical questions in Chapter 4.

SELECTED READINGS

Animal Development—General Texts


Organizers, Fields, and Morphogens


Homeotic Genes in Insects and the Homeobox

Field-specific Selector Genes


Compartment Selector Genes


Cell-type-specific Selector Genes


Genetics of Embryonic Axis Formation in Drosophila

Nüsslein-Volhard C, Wieschaus E. Mutations affecting segment number and polarity in 

St Johnston R, Nusslein-Volhard C. The origin of pattern and polarity in the *Drosophila* embryo. 

**Sharing of the Toolkit Among Animals**

Duboule D, Dollé P. The structural and functional organization of the murine HOX gene family 


Graham A, Papalopulu N, Krumlauf R. The murine and *Drosophila* homeobox gene complexes 


Quiring R, Walldorf U, Kloter U, Gehring WJ. Homology of the eyeless gene of *Drosophila* to 

Riddle C, Johnson R, Laufer E, Tabin C. Sonic hedgehog mediates the polarizing activity of the 
CHAPTER 3

Building Animals

The fundamental mystery of development is how a single fertilized egg cell multiplies and differentiates into a complex animal composed of a large number and wide variety of specialized organs, tissues, and cell types. Identification of the genes with the most pronounced effects on development is merely a first step to an understanding of the process. The second challenge is to understand how the activity of these genes unfolds in time and space to control the formation and patterning of animal body plans and body parts. This process is primarily a matter of gene regulation.

Every feature of animal development depends on both serial and parallel gene functions that act within and between regulatory hierarchies. Because evolutionary changes in animal development can arise from changes in the operation of regulatory hierarchies, understanding the genetic logic and molecular circuitry of major regulatory programs is fundamental to understanding the evolution of morphology.

In this chapter, we examine the architecture of developmental regulatory hierarchies at two levels. First, we analyze the genetic logic of the hierarchies that control the formation and patterning of the primary embryonic axes, appendages, organs, and other major features of selected model animals. Second, we analyze the ways in which combinations of regulatory inputs within these hierarchies are integrated at a molecular level by regulatory elements of developmental genes to control their stage- and tissue-specific expression and functions.

GENE REGULATION IN METAZOANS

The evolution of animal bodies composed of different cell types required mechanisms for turning gene expression on and off in specific cells at particular times during development. The temporal and spatial dimensions of animal development, the thousands of genes in animal genomes, the tendency for the same gene product

I should like to work like the archaeologist who pieces together the fragments of a lovely thing which are alone left to him. As he proceeds, fragment by fragment, he is guided by the conviction that these fragments are part of a larger whole which, however, he does not yet know.

—Hans Spemann, Embryonic Development and Induction (1938)

Several genetic steps seem to be interposed between the reception of a signal, extrinsic to the genome, and its translation first into genetic and later into developmental terms. A hierarchy of genes may be involved in this process...

—Antonio García-Bellido (1975)
DNA to Diversity: Molecular Genetics and the Evolution of Animal Design

Both the protein machinery involved in transcription to produce messenger RNAs (mRNAs) and the structure of genes at the DNA level are more complicated in metazoans than in prokaryotes. Activation of tissue-specific gene expression requires the assembly of protein complexes involving at least three components (Fig. 3.1):

- RNA polymerase II and its associated general transcription factors
- Cell-specific, tissue-specific, field-specific, or signal transducer-type activators
- Coactivators that make contacts between activators and the general transcription machinery and influence the local state of chromatin

Importantly, gene regulation is not all about activation. The spatial and temporal specificity of gene expression depends a great deal on specific repressors that act to suppress transcription, often through corepressors that affect the local state of chromatin. Many of the genes in the toolkit for animal development are transcriptional activators or repressors.

The general transcription machinery is assembled on DNA sequences called promoters, which are found near the site of the start of transcription. Activators and repressors bind to
other DNA sequences to affect the transcriptional activity of the gene. Sometimes, these sites may be located in the vicinity of the promoter regions. Often, however, the regulatory proteins that mediate the transcriptional activity of individual developmentally regulated genes bind to specific sites in discrete DNA regions, called **cis-regulatory elements** (or cis-elements), which can influence the level of transcription by several orders of magnitude, from sites that may be many thousands of base-pairs away from the promoter. The modularity of these elements is illustrated by their ability to regulate transcription of essentially any gene in cis. Such cis-elements are usually identified and their function studied by analyzing their activity in regulating a heterologous **reporter gene** (Fig. 3.2). For example, they may regulate transcription through the looping of distant sequences that brings bound activators and coactivators in contact with the transcription initiation complex at the promoter; they may also act by affecting the state of chromatin (Fig. 3.1).

The expression of individual genes at particular stages and locations in the developing animal is often controlled by cis-elements that are distinct from those controlling gene expression at other times and places. Thus metazoan genes not only consist of the coding sequences for a particular protein, but also often possess a modular array of cis-elements that act as genetic switches to control gene expression in a variety of different contexts. The modularity of metazoan cis-regulatory DNA is critical both to the specificity of gene interactions during development and to evolutionary changes in gene expression during the evolution of new morphologies.

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**THE ARCHITECTURE OF GENETIC REGULATORY HIERARCHIES**

**General features and approach**

Many genes can affect the same structure, pattern, or process. In many cases, it is impossible to determine the order of gene action solely by examining mutant phenotypes. Nor is it usually clear whether one gene plays a primary role in a process and other genes play subordinate roles. For example, for any two genes \(A\) and \(B\) (which based on similar mutant phenotypes appear to control the same process), several regulatory relationships are possible. \(A\) could affect the expression or function of \(B\), \(B\) could affect \(A\), \(A\) and \(B\) could affect each other, or \(A\) and \(B\) may have no effect on each other. With additional genes, one can quickly see that the number of possible regulatory relationships among a set of genes increases exponentially with the size of the set. Given that genetic screens have often identified multiple loci that have similar developmental effects, it is clear that to decipher the temporal order of action and hierarchical relationships between genes, more information is necessary.

Molecular techniques have revealed that the patterns in which major developmental genes are expressed within various body regions, organs, and cells of the developing animal usually correspond to the structures of the larvae or adult whose formation, pattern, or differentiation are affected by mutations in these genes. This understanding has inspired a whole new approach to embryology and the genetic analysis of developmental regulatory mechanisms. Rather than focus on final physical forms, which are often arrested prematurely and disfigured in lethal mutants, the patterns in which key developmental genes are expressed at different developmental stages can be used as surrogates for the ultimate form. Visualization of gene activity in developing animals provides a much more direct and dynamic (that is, both
Animal genes often contain multiple independent elements that control gene expression in different places and/or different times during embryogenesis (for example, A, B, C). The cis-regulatory elements are generally identified through their ability, when placed in cis to a heterologous reporter gene and inserted back into a host genome, to control the pattern, timing, and/or level of gene expression. The most frequently employed reporter genes encode either (1) enzymes that can be detected in situ with chromogenic substrates or specific antibodies or (2) proteins that fluoresce in vivo.
Chapter 3: Building Animals

The study of model organisms such as *Drosophila* has revealed a few general features of the architecture of genetic regulatory hierarchies underlying developmental programs. First, development is a continuum in which every pattern of gene expression has a preceding causal basis—namely, a previous pattern of gene activities. Second, regulatory information often flows through “nodal points”—key genes that integrate multiple spatial inputs and whose products often control a major feature of the future pattern. For example, the location of organ primordia often requires inputs from anteroposterior and dorsoventral coordinate systems that are integrated by selector genes, which directly control the formation of these primordia. By focusing on the genetic control of such nodal genes, we can simplify and resolve the regulatory logic of networks that might appear quite complicated at the formal genetic level. Third, many genes, particularly components of signaling pathways and transcription factors, are deployed at several stages of development in distinct spatial patterns and are involved in a variety of regulatory hierarchies.
Regulatory logic—pathways, circuits, batteries, feedback loops, networks, and the connectivity of genes

The analysis of regulatory interactions between genes has given rise to a host of commonly used terms, often applied with somewhat broad meanings, to describe the higher-order relationships and connections between genes. Because the liberal use of these terms can obscure the concepts they are intended to represent, we take a moment here to define their use in this book.

Let’s start with a pathway. We use this term to describe components that are obligately linked in the transmission of information. Signaling pathways, which may be linear or branched, are composed of components that depend on one or more upstream or downstream components to exert their effects (Fig. 3.3). Multiple structurally related ligands, receptors, or transducers might be able to transmit information in a given pathway.

We define a circuit to be larger than a pathway, encompassing additional components that are not obligately linked (Fig. 3.3). For example, while components of signaling pathways...
depend on one another to conduct the signal to the cell nucleus, the nature of the pathway output is usually context-dependent. When gene regulation is the output, the target genes of a given individual pathway usually differ between developmental stages and spatial locations in the developing embryo. We use the term “circuit” to describe a regulatory pathway that includes particular target genes. Thus two circuits that employ the same pathway are different if they regulate different target genes.

A group of connected circuits constitutes a regulatory network. These circuits may be connected in series or they may consist of parallel circuits that are connected by one or more links (Fig. 3.3). Some genes in the network may act earlier than others, or control the activity of many genes, organizing the network into a hierarchy. Hierarchies may be vertically organized, comprising many tiers of genes that control lower tiers of genes (Fig. 3.3). Alternatively, they may be organized more horizontally, with single genes directly controlling the expression of a large group or battery of target genes (Fig. 3.3).

Model regulatory hierarchies and the key genetic switches that operate them

The regulatory mechanisms that underlie a large number of developmental processes in the several model species belonging to a few different phyla have been analyzed in recent years. In this book, we concentrate primarily on insect and vertebrate examples because they illustrate general mechanisms of the control of toolkit gene expression and function during development. Importantly, knowledge derived from these model systems forms the foundation of the comparative approaches to the evolution of morphology described in subsequent chapters.

Our discussion focuses on the regulatory hierarchies that control the sequential generation of spatial coordinate systems that unfolds during embryogenesis—from the generation of the major body axes, to the formation of primary and secondary fields, to the patterning of individual fields. We examine at the molecular level how organizers function within embryos and fields, how gradients of morphogens are interpreted within fields, how selector genes regulate the formation and identity of individual fields, and how cis-regulatory elements integrate combinations of regulatory inputs to control complex spatial patterns of toolkit gene expression.

THE INSECT BODY PLAN

The genetic regulatory hierarchies that establish the major features of the Drosophila melanogaster body plan are of interest for two reasons: (i) because they are among the best understood in any animal and illustrate potentially general principles, and (ii) because they provide a basis for comparison between Drosophila and other insects and arthropods. The structure of these regulatory hierarchies is critical to mechanisms of arthropod body plan evolution. The major features of the insect body plan that have both developmental and evolutionary significance include the following:

- Segmentation
- The organization of segments into distinct regions forming the head, thorax, and abdomen
The identities of paired, jointed appendages on different head and thoracic segments

The limbless abdomen

The two pairs of dorsal flight appendages

Fruit flies are holometabolous insects, meaning that they undergo a complete metamorphosis between their larval and adult forms during the pupal stage. The larval body plan differs from the adult body plan (for a view of the fly life cycle, see Fig. 3.4). Here we examine regulatory mechanisms involved in both larval and adult patterning.

From egg to segments: the anteroposterior coordinate system

In just one day following fertilization, the Drosophila embryo develops from a single nucleus in a huge yolk-filled egg about 0.5 mm long into a highly organized segmented, motile, feeding larva with a complex nervous system and the future adult tissues growing as imaginal structures within it. Genetic screens identified five tiers of regulatory genes involved in organizing body pattern along the primary anteroposterior (A/P) axis of the developing embryo. As discussed in Chapter 2, the maternal effect, gap, pair-rule, segment polarity, and homeotic genes have large and distinct effects on the patterning of the A/P axis.
Genetic and molecular analyses of the regulatory interactions within and among these five tiers of genes have elucidated both the general logic of and many specific molecular interactions that operate within the A/P patterning hierarchy. These molecular mechanisms, in turn, illustrate several important general concepts about the function and transcriptional regulation of pattern-regulating genes during animal development.

Figure 3.5 shows the basic outline of the A/P axis regulatory hierarchy. The generation of the periodic, segmental organization of the larva from an initially aperiodic egg involves...
regulatory interactions between gene products translated from maternal mRNAs deposited in the egg, transcriptional activation of zygotic genes by certain maternal activators, and combinatorial action of segmentation gene products to refine the expression patterns of many zygotic segmentation genes into iterated domains. This hierarchy has four key features:

1. Generation of gradients of maternal transcription factor proteins
2. Transcriptional activation of and cross-regulation by the gap genes
3. Transcriptional regulation of individual pair-rule gene stripes by combinations of maternal and gap proteins
4. Regulation of segment polarity gene expression by pair-rule and segment polarity proteins

**Generation of maternal transcription factor gradients**

The Bicoid (Bcd) and Caudal (Cad) proteins are homeodomain-type transcription factors, and the Hunchback (Hb) protein is a zinc-finger-type transcription factor. The mRNAs encoding these proteins are deposited in the egg and translated during early embryonic development. Because the early embryo is a syncytium, lacking any cell membranes that would impede the diffusion of protein molecules, these transcription factors can move freely throughout the cytoplasm. The concentration of each protein is graded along the A/P axis of the embryo (Fig. 3.6). These maternal gradients are important because several downstream segmentation genes are regulated by Bcd, Cad, and Hb (depending on their concentrations). Early gene regulation in the fly embryo provides a general model for the concentration-dependent control of gene expression by gradients of regulatory proteins.

In *Drosophila*, these three maternal gradients are formed by different mechanisms. The Bicoid gradient results from the localization of maternal mRNA at the anterior pole of the egg. Translation of this mRNA and diffusion of the Bicoid protein toward the posterior creates a concentration gradient. The Hunchback gradient is generated by selective inhibition of the translation of the ubiquitous *hb* mRNA in posterior regions (Fig. 3.6b). This inhibition is regulated by the product of the *nanos* gene, which binds to the *hb* mRNA. The Nanos (Nos) protein is in a posterior-to-anterior concentration gradient, generated by the localization of its mRNA to the posterior end of the egg. The caudal mRNA, like hunchback, is distributed evenly throughout the egg, although selective inhibition of its translation by the Bicoid protein generates a posterior-to-anterior Caudal protein gradient (Fig. 3.6b).

**Transcriptional activation of and cross-regulation by gap genes**

The gap genes are the first zygotic genes to be expressed in discrete regions along the A/P axis. The Bcd, Hb, and Cad proteins are involved in the initial regulation of these genes. One key regulatory interaction is the activation of the second phase of *hunchback* expression by the Bicoid protein in the anterior half of the embryo. This activation occurs through a direct interaction of the Bicoid protein with a cis-regulatory element of the *hb* gene. DNA sequences 5′ to the *hb* promoter contain several binding sites for the Bicoid protein that are necessary for the Bcd-dependent activation of the *hb* gap domain (Fig. 3.6c). Bcd binds to these sites cooperatively—that is, the binding of one Bcd protein molecule to one site facilitates the binding of other Bcd molecules to nearby sites. More than one occupied Bcd site is necessary
to generate the sharp boundary of \( h_b \) transcription, suggesting that this cis-regulatory element requires a threshold concentration of the Bcd protein to occupy multiple binding sites before it can activate \( h_b \) expression. In this way, the \( h_b \) gene “reads” or “interprets” the Bcd gradient.

Different gap genes are deployed in distinct domains in the embryo, depending on the levels of the Bcd, Cad, and Hb proteins. Each gene possesses cis-regulatory elements that contain different arrangements of binding sites with different affinities for these transcription factors. Consequently, gap genes are activated at different positions along the A/P axis. Their expression domains are further refined through mutually repressive interactions between gap proteins (all of which are transcription factors) and the cis-regulatory elements of gap genes expressed in adjacent domains.
Initiation of periodic pair-rule gene expression

The aperiodic maternal and gap protein expression patterns establish the periodic, seven-striped patterns of certain pair-rule genes, called “primary” pair-rule genes. The way in which the pattern of pair-rule stripes is created by aperiodically distributed regulators had been one of the major mysteries of *Drosophila* segmentation. The solution to this puzzle highlights one of the most important concepts in the spatial regulation of gene expression in developing animals—namely, the independent control of distinct cis-regulatory elements in a single gene.

The key discovery was that the seven stripes that make up the expression patterns of the primary pair-rule genes *hairy* and *even-skipped* are controlled independently. That is, different cis-regulatory elements control the expression of different stripes. The seven-striped pattern of *eve* expression, for example, represents the sum of seven sets of regulatory inputs into seven separate cis-elements of the gene (Fig. 3.7). Detailed analysis of individual stripe cis-regulatory elements revealed that the position of a “simple” stripe was controlled by no fewer than four regulatory proteins. For example, for the second *eve* stripe, Bcd and Hb act as broadly distributed activators, and the boundaries of the stripe are sharpened via repression by the Giant and Krüppel gap proteins (Fig. 3.7). This roughly 700-bp element contains multiple binding sites for each activator and repressor. The cis-regulatory element acts as a genetic switch, integrating a host of regulatory protein activities to regulate *eve* expression in one three- to four-cell-wide stripe in the embryo.

Other *eve* and *hairy* stripe elements are regulated by different combinations of maternal activators and gap proteins found at other positions in the embryo. The periodic expression of these primary pair-rule proteins, in turn, regulates the periodic expression of downstream “secondary” pair-rule genes.

Regulation of segment polarity genes by pair-rule proteins

The 14-stripe patterns of several segment polarity genes (corresponding to 14 segments) are initiated by the pair-rule regulatory proteins, then subsequently refined and maintained by the segment polarity gene products themselves. Pair-rule proteins regulate alternating (7 out of 14) segment polarity stripes. For example, Ftz acts through a cis-regulatory element in the *engrailed* gene to help activate the even-numbered *en* stripes. The Odd-skipped (Odd) pair-rule protein is expressed slightly out of phase with the Ftz protein and represses expression of the *en* gene. In this manner, narrow *en* stripes are activated in cells that express Ftz but do not express Odd.

After the pair-rule proteins position the initial expression domains of segment polarity genes in the growing (now cellularized) embryo, interactions among the segment polarity proteins and genes refine and maintain these patterns. These regulatory circuits are important because they maintain boundaries between compartments and segments; they also regulate the expression of other genes that control the morphogenesis of polarity. The pathways that operate within these circuits include some of the most widely deployed pathways in animal development. Much has been learned about their general mechanisms through the study of segment polarity in *Drosophila*.

Two major signaling pathways regulate the maintenance and elaboration of polarity within *Drosophila* segments: the Wingless and Hedgehog pathways. The Hh protein is produced in all posterior cells in each segment, which also express *engrailed*. Hh signals anterior cells
Figure 3.7  
Regulation of a pair-rule stripe: combinatorial control of an independent cis-regulatory element

The regulation of the even-skipped cis-regulatory element controls the formation of the second stripe in the early embryo. (a) The stripe 2 element controls just one of seven stripes of eve expression. (b) The stripe forms within the domain of the Bicoid and Hunchback proteins and at the edge of the Giant and Krüppel gap protein domains. The former are activators, and the latter are repressors, of eve stripe 2 expression. (c) The eve stripe 2 element spans from about 1.7 to 1.0 kilobases upstream of the eve transcription unit. (d) Within this element, several binding sites for each regulator exist. The net output of the combination of activators and repressors is expression of the narrow eve stripe.

through the Ptc receptor and several transducers (Fig. 3.8), to activate *wingless* expression in adjacent anterior cells. Wg, in turn, signals back to posterior cells through the Wingless receptor complex (D-fz and Arrow) and various signal transducers (see Table 2.2), to maintain *engrailed* expression, *Hh* expression, and the continuity of the regulatory circuit (Fig. 3.8). Signaling between cell populations to initiate and maintain gene expression patterns is a general feature of cellular fields. Indeed, these signaling molecules play a wide variety of roles in animal development.

**General lessons from the segmentation genes**

The A/P segmentation hierarchy illustrates five general concepts concerning the spatial regulation of gene expression in animal development:

1. The concentration-dependent response of genes to graded inputs is illustrated by the regulation of gap target genes by the Bicoid protein. Both threshold and graded responses to inducers are major themes of gene regulation in cellular fields as well.

2. The action of both activators and repressors determines all of the gap and many of the pair-rule and segment polarity gene expression patterns. The refinement of gene expression patterns from those covering most of the embryo (maternal proteins), to 15- to 20-cell-wide regions (gap proteins), to 3- to 4-cell-wide stripes (pair-rule proteins), and ultimately to 1- to 2-cell-wide stripes (segment polarity proteins), depends on activators that define potential areas of gene activation and repressors that restrict
the areas in which target genes are expressed. The spatial repression of gene expression to carve ever-finer patterns out of larger domains is a major regulatory theme in cellular fields.

3. Many genes are regulated by two or more activators or repressors. **Combinatorial regulation** imposes greater specificity and allows for a greater diversity of spatial patterns.

4. **Multiple independent cis-regulatory elements** regulate the expression of many segment genes. Individual elements control individual domains of gene expression (for example, each pair-rule stripe). Furthermore, many segmentation genes are also expressed in the developing nervous system in unique patterns that are controlled by other discrete cis-regulatory elements. The utilization of multiple independent elements controlling different spatial domains of gene expression is a general theme of developmental gene regulation.

5. The sequential activation of gap, pair-rule, and segment polarity genes, with each tier of genes being dependent on the preceding tier, constitutes a **regulatory hierarchy**. The domino-like activation of genes in sequence assures the proper temporal deployment of developmental genes.

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**The dorsoventral axis coordinate system**

Like the A/P axis, the dorsoventral (D/V) axis of the *Drosophila* embryo is subdivided into a series of domains that give rise to different tissues. Ventral-most cells will form mesoderm, more ventrolateral cells will give rise to the neuroectoderm, lateral regions generate the dorsal epidermis, and dorsal cells produce an extraembryonic structure, the amnioserosa (Fig. 3.9a). Although subdivision of the D/V axis is accomplished by an entirely different set of genes from those that regulate the A/P axis, both transformations occur through similar transcriptional regulatory mechanisms. Combinations of activators and repressors, including some that work in a concentration-dependent manner, act on discrete cis-regulatory elements to carve out the spatial boundaries of downstream pattern-regulating genes.

A maternal transcription factor gradient also organizes the D/V axis. This gradient is established by regulating the nuclear localization of the Dorsal (Dl) protein. The highest concentrations of the nuclear Dorsal protein are found in ventral cells, lower levels occur in ventrolateral and lateral regions, and no nuclear Dorsal protein is found in dorsal-most regions (Fig. 3.9c). The protein is required in the ventral region to induce ventral tissues.

More than 40 zygotic genes are regulated by different threshold responses to Dorsal protein concentration along the D/V axis, a few of which have been examined in great detail (Fig. 3.9b). The response of genes to the Dorsal gradient depends on the number and affinity of binding sites for the Dorsal protein within the cis-regulatory elements of target genes. Low-affinity sites are occupied only at high concentrations of Dl nuclear protein, which are found in ventral-most cells. For example, the *twist* and *snail* response elements contain low-affinity sites and are activated in ventral-most cells. By contrast, high-affinity sites can be occupied at lower concentrations of Dorsal. The *rhomboid* gene cis-element, for example, contains high-affinity sites and is activated by lower concentrations of Dorsal in lateral regions. This protein also acts as a repressor to prevent the expression of genes such as *zen* and *dpp* in ventral and lateral regions of the embryo.
Figure 3.9
The dorsal gradient and the dorsoventral genetic regulatory hierarchy
(a) Different zygotic genes are expressed in different spatial domains along the D/V axis of the embryo. Some of these domains correspond to populations of cells that will give rise to distinct regions of the embryo (such as the mesoderm and neurogenic ectoderm). (b) The battery of genes regulated by Dorsal protein. (c) Dorsal regulation of target genes is dependent on threshold concentrations of protein. The concentration of nuclear Dorsal protein is highest in ventral (V) cells and lowest in dorsal (D) cells. Certain target genes are activated at different concentrations of Dorsal (thresholds 1, 2, and 3).

The sharpness of the expression boundaries of the zygotic D/V patterning genes is refined by further regulatory interactions. The snail (sna) gene enhancer, which is bound by Dl, also requires the product of the twist (twi) gene to be activated. Synergistic interactions between Dl and Twi ensure a sharp on/off border of sna expression, which coincides with the boundary between and regulates the differentiation of the mesoderm from the neuroectoderm (Fig. 3.9c). The rhomboid (rbo) element is repressed by the Snail protein, which eliminates rbo expression from the mesoderm and restricts it to the neuroectoderm (Fig. 3.9c). A cascade of subsequent interactions, involving primarily the activity of the Dpp signaling molecule, further subdivides the dorsal region of the embryo.

**The Hox ground plan**

In *Drosophila* and many other animals, the spatial and temporal expression patterns of individual Hox genes are much more complicated than those of most segmentation or D/V axis-patterning genes. The number of cis-elements and trans-acting regulators for each gene is therefore considerably greater for the Hox genes. The regulation of Hox gene expression has the following major features:

- Activation in a broad domain, usually consisting of two or more parasegments
- Modulation of the levels of expression within this domain
- Expression in all three germ layers, often in patterns that are slightly out of register between each germ layer
- Dynamic changes in the level and domains of expression throughout the subsequent course of development

Numerous transcription factors act on *Drosophila* Hox genes; likewise, many cis-acting regulatory elements control the various features of Hox expression. In the embryonic ectoderm, for example, most Hox genes are regulated by as many as six classes of regulators. All three classes of zygotic segmentation genes are involved in Hox regulation. Gap proteins regulate the broad initial domains of Hox transcription, selected pair-rule proteins are involved in setting the initial spatial register of certain Hox gene domains, and segment polarity proteins, such as the Engrailed protein, help provide intrasegmental modulation of Hox gene expression levels. A fourth level of control involves other Hox proteins—the posterior boundaries and levels of expression of some Hox genes are regulated by more posteriorly acting Hox proteins. A fifth level of control involves autoregulatory feedback mechanisms that operate to maintain Hox expression domains. Finally, a large group of chromatin-associated proteins, termed the Polycomb group and Trithorax group of proteins, act to maintain the repressed or activated transcriptional state, respectively, of many Hox genes (Fig. 3.10).

The Ultrabithorax gene provides a good example of the constellation of cis-regulatory elements that control Hox gene expression. Different cis-elements of Ubx control expression in particular parasegmental domains, in a manner analogous to individual pair-rule stripe elements (Fig. 3.11). These elements contain binding sites for several activators and repressors. The pattern of Ubx expression controlled by these elements is the net output from numerous positive and negative inputs. Other Ubx elements control the level of expression within parts of parasegments, in appendage fields later in development, and in other germ layers such as the mesoderm (Fig. 3.11).
**Figure 3.10**  
_Hox_ genes are regulated by many inputs  
Several classes of gene products regulate the initiation, modulation, and maintenance of _Hox_ gene expression in the developing embryo and larva of _Drosophila_. Positive regulatory inputs are denoted by arrows; negative regulatory inputs are shown by a truncated line.

**Figure 3.11**  
The cis-regulatory elements of the _Ubx_ gene  
(a) Three elements that regulate _Ubx_ expression in the embryo are shown. Each element controls expression in selected subsets of parasegments.  
(b) The BRE element contains binding sites for several segmentation proteins (Ftz, Hb, En, Tll) and a dorsoventral patterning protein (Twi) within a 500-bp span.  
(c) Expression driven by the BRE element in parasegments 6, 8, 10, and 12 is the net output of the various positive and negative regulatory inputs shown in this figure.
Secondary fields: integrating the anteroposterior and dorsoventral coordinate systems

The anteroposterior and dorsoventral axis regulatory genes establish a two-dimensional coordinate system over the entire embryo and within each segment. The Hox ground plan is superimposed on this coordinate system and provides the regulatory information for differentiating the events that take place within different segments.

Once the two coordinate systems and Hox ground plan are deployed, the development of secondary fields begins in the Drosophila embryo. Neural precursors and the primordia for structures such as the heart, salivary glands, and imaginal discs (which will later give rise to the adult appendages and body wall) form at discrete A/P and D/V coordinates within particular segments. The positions of these structures in the embryo are specified in a manner analogous to the way a geographical position is defined on the globe through the combination of latitude and longitude.

Specification of organ precursor position, number, and identity requires integration of regulatory inputs from both coordinate systems, the Hox genes, and sometimes other regulators. As described in Chapter 2, the development of various precursors and organ fields is controlled by specific selector genes. The activation of these genes is often the initial step in the specification of these structures. The regulation of these field- and cell-type-specific selector genes reflects the molecular output of the integration of the various spatial regulatory systems. The analysis of their regulation reveals a great deal about the genetic logic underlying early organogenesis. There are several well-studied examples of the integration of axial inputs in the specification of embryonic organs and secondary fields.

The limb fields

The ventral limb fields (legs, mouthparts, and others) and dorsal flight appendage fields (wing, haltere) arise from small populations of approximately 20 cells that are initially specified during mid-embryogenesis. The first sign of the specification of these primordia is the activation of regulatory genes within them. The expression of the Distal-less gene marks the development of the ventral limb primordia. Dll is expressed initially in small clusters of cells in the ventrolateral ectoderm in many of the head segments and in each thoracic segment. The ventrolateral and anteroposterior positions of these clusters of cells are specified by signals that organize patterning along the dorsoventral and anteroposterior axes of each segment.

Both positive and negative regulatory inputs are used to position the clusters. Along the A/P axis, the Wingless signaling protein is a positive regulator of Dll expression. Dll is not induced everywhere that the Wg signal is present, however. Instead, Dll expression is restricted to a ventrolateral position by negative regulation in dorsal cells by the Dpp signal and in ventral cells by a signal acting through the EGF receptor pathway (Fig. 3.12).

All three signals—and hence the proper coordinates for Dll expression—are present in each trunk segment, yet Dll is not expressed in the abdominal segments. This situation occurs because Hox regulation of Dll is superimposed on the segmentally reiterated coordinate system to prevent activation in the abdomen. The Ubx and abd-A proteins bind directly to and repress a cis-regulatory element of the Dll gene that controls Dll expression in the embryonic thoracic segments (Fig. 3.12).
The *Dll* gene possesses many *cis*-regulatory elements that function in limb primordia in parts of the embryonic head and/or thorax. Some of these elements are regulated by other *Hox* genes, others appear to be redundant. The existence of several elements allows for the independent control of the timing, level, and pattern of *Dll* expression in different segments and at different stages of development.

**The wing primordia**

The formation of the wing primordia is also regulated by intrasegmental signals and repressed in specific segments by *Hox* proteins. Cells that form the wing primordia migrate out of the limb field and are specified later and at a more dorsal position than are the limb primordia. Unlike...
the limb primordia, the wing primordia form in a region characterized by high levels of Dpp signaling (Fig. 3.12). Primordia formation is selectively repressed in the first thoracic and all abdominal segments by the Scr, Ubx, and Abd-A proteins (Fig. 3.12). We will have more to say about the role of Hox genes in the evolution of insect limb and wing number in Chapter 5.

The salivary gland

The salivary gland primordia form within the ventral epidermis of parasegment 2 in the developing embryo. The positioning of this organ and its restriction to parasegment 2 follow a similar logic, as we have described for the appendage primordia. Expression of the transcription factor Forkhead (Fkh) is an early event in salivary gland development. The regulation of the fkh gene parallels the regulation of salivary gland organogenesis. Although the fkh gene and salivary gland development are activated by the Hox protein Scr in parasegment 2, they are limited to the ventral epidermis by negative regulation, from the Dpp signal dorsally, and the EGF receptor pathway more ventrally (Fig. 3.12).

Neural and muscle precursors

The patterns of embryonic neural and muscle precursors in the ectoderm and mesoderm, respectively, are also generated by combinatorial inputs of A/P and D/V signals and modified by Hox regulatory inputs. Rows of proneural clusters that will give rise to segmentally iterated patterns of neuroblasts in the early embryo are marked by the expression of Achaete-Scute Complex genes, whose regulation is a nodal point for the patterning of the central and peripheral nervous system. In the AS-C, cis-regulatory elements integrate inputs from segmentation proteins and dorsoventral-axis patterning proteins that establish proneural clusters. Later in embryogenesis, segment-specific patterns of peripheral neural precursors arise from combinatorial regulation of AS-C genes by segmentation, dorsoventral, and Hox regulatory inputs.

The AS-C contains many cis-regulatory elements that control discrete subpatterns of AS-C gene expression. Muscle precursors, marked by the expression of the nautilus gene, are positioned by combinatorial signals within the mesoderm as well as by Hox inputs that regulate segment-specific muscle patterns.

Patterning within secondary fields: organizing signals and selector genes

The intrasegmental coordinate systems in the embryo and the Hox ground plan control the initial position, size, and number of developing Drosophila fields. As these fields grow and begin to differentiate, regulatory hierarchies within these fields establish new coordinate systems and organizers that control the formation and morphogenesis of the final structure.

Perhaps the best-studied genetic regulatory program of all secondary fields is that of the Drosophila wing. This wing develops from a portion of the larval imaginal disc that will also give rise to the body wall of the second thoracic segment. The wing disc grows from roughly 30–40 cells in the first instar larva to approximately 50,000 cells in the third instar larva before morphogenesis transforms it into the adult wing and thoracic body wall. Coordinate systems and selector genes operate within the wing field to control the position, number, and differentiation of the various pattern elements.
The regulatory logic and several of the molecular mechanisms involved in the hierarchies that control wing formation and patterning have been described in sufficient detail that some general concepts regarding the patterning of secondary fields have emerged. These concepts, which apply to other insect appendages as well as to vertebrate limbs, involve regulatory mechanisms that are similar to those operating to position and regulate the formation of pattern elements in other cellular fields. Because the number, position, size, and morphology of pattern elements within secondary fields are important aspects of morphological diversity, understanding these regulatory mechanisms is crucial to developing a picture of the evolution of gene expression and morphology within a field.

The development of secondary fields from primordia established in the embryo involves three major processes:

1. The generation of new coordinate systems during the growth of the field
2. The placement and specification of pattern elements within the field
3. The differentiation of field identity from other, serially homologous fields

The genetic regulatory mechanisms governing these processes involve the integration of combinatorial inputs by cis-regulatory elements, similar to those described earlier for the generation of body axes and the initial specification of the fields. One set of regulators includes short- and long-range signaling proteins (morphogens) that determine the area within a field in which a given gene is activated or repressed. The sources of these signals are usually oriented with respect to one patterning axis of the field (anteroposterior, dorsoventral, or proximodistal). A second regulator controls the field- or cell-type-specific response to these signals, such as a selector gene. The identity of serially homologous fields is determined by another layer of regulation superimposed on the signaling inputs and field-specific selector genes by Hox genes or analogous regulators that are expressed in only one serial homolog. The integration of signaling, field-specific selector, and Hox inputs through cis-regulatory elements of target genes controls gene expression patterns in a particular field.

In the Drosophila wing and haltere (the wing’s serial homolog), the identity and function of the major signaling (Dpp, Hh, Wg, N), wing-specific selector (Vg, Sd), and Hox (Ubx) inputs in the wing patterning hierarchies have been studied extensively. Three coordinate systems operate in the wing field:

1. The first coordinate system is involved with patterning along the A/P axis.
2. The second coordinate system acts along the D/V axis.
3. The third coordinate system, the proximodistal (P/D) axis, affects the integration of these inputs and the specification of the entire wing field.

Next, we examine the genetic regulatory hierarchies that govern the formation and operating of these coordinate systems and see how they control gene expression and wing pattern.

**The anteroposterior coordinate system**

The A/P coordinate system encompasses the sequential organizing activities of the Engrailed, Hedgehog, and Dpp proteins. Patterning along the A/P axis of the wing is controlled by a hierarchy involving compartment selector genes and multiple signaling pathways.
Posterior cells are segregated from anterior cells by a lineage restriction imposed by the *engrailed* gene, which is expressed in all posterior cells but not in anterior cells. This lineage restriction creates a smooth boundary between the anterior and posterior compartments, across which important inductive events take place. Due to the regulatory activity of *engrailed*, cells in the posterior (but not the anterior) compartment express the short-range signaling molecule Hedgehog (Fig. 3.13a). Cells in the anterior compartment express the Patched protein (see Table 2.2), a receptor for the Hh signal. Transduction of Hh signaling induces a stripe of cells on the anterior side of the A/P boundary to express the Dpp signaling protein (Fig. 3.13a,b). Dpp, in turn, acts as a morphogen and diffuses across both the anterior and posterior compartments from its source along the A/P compartment boundary, thereby regulating gene expression in a concentration-dependent manner.

**The dorsoventral coordinate system**

The D/V coordinate system includes the sequential organizing activities of Apterous and the Notch and Wingless pathways.

The organization of the D/V axis is regulated in a fashion somewhat analogous to the regulation of the A/P axis organization. Dorsal cells are segregated from ventral cells by a lineage restriction imposed by the *apterous* compartment selector gene (see Table 2.1), which is expressed in all dorsal cells (Fig. 3.13a). The Apterous protein regulates the expression of Serrate and Fringe (Fig. 3.13b & see Table 2.2), two proteins that interact with the ubiquitously expressed Notch receptor protein. Along both sides of the D/V boundary, Notch-mediated signaling between cells induces expression of the Wg signaling protein in a stripe that straddles the D/V boundary. Wg acts as a morphogen, diffusing from its source to regulate gene expression in both the dorsal and ventral compartments.

**Signal integration by the vestigial field-specific selector gene**

One important downstream target of the Dpp, N, and Wg organizing signals emanating from the compartment boundaries is the *vg* field-specific selector gene. The *vg* gene is expressed in all of the cells in the wing disc that will form the wing. The regulation of the overall pattern of its expression is the sum of inputs from two separate *cis*-regulatory elements that respond to signals from each coordinate system (Fig. 3.13c). The “boundary” *cis*-element is directly activated along the D/V boundary through the Notch pathway. The “quadrant” *cis*-element is expressed in the complementary pattern in the rest of the wing field through activation by the Dpp pathway and other regulators (Fig. 3.13c). Expression of this field-specific selector gene depends directly on the signaling sources of the wing field and indirectly on the compartmental selectors that establish the expression of these signals.

**Combinatorial regulation of wing patterning by signaling proteins and the Vg/Sd selector proteins**

Within the wing field, a large number of genes function to pattern various features of the wing. For example, the AS-C genes are activated along the anterior extent of the D/V boundary to generate the sensory organs of the leading edge of the adult wing. The *blistered* or *Drosophila Serum Response Factor* (D-SRF) gene is expressed in all of the cells that lie between the developing wing veins. Many genes are involved in the positioning and differentiation of the
Figure 3.13
The genetic regulatory hierarchy in the Drosophila wing
(a) Two systems organize the pattern along the dorsoventral (left) and anteroposterior (right) axes of the wing. (b) Compartmental selector genes establish signaling sources along each compartment boundary. Transduction of these signals leads to the activation of downstream target genes. (c) One important target that is directly regulated by these signals is the vestigial wing selector gene. The vg gene is expressed in all wing cells through the sum of the activities of two cis-regulatory elements—one active along the compartment boundaries, and one active in the remaining four quadrants of the wing field.
veins that provide the structural support for and the fluid transport system of the wing. The expression of each gene required for the development of wing pattern elements depends on combinatorial inputs from signaling pathways and the wing selector proteins Vg and Scalloped (the DNA binding partner of Vg). Specifically, the signaling proteins regulate where within the wing field a given gene is activated or repressed, and the activities of Vg/Sd make the response to these signals specific to the wing field.

**The response to organizing signals**

Various ligands for the major signaling pathways (Serrate, Dpp, Wnt, Hedgehog, and so on) are deployed in dynamic patterns in the wing field. Some signals appear to have a greater range of influence than others do. The Hedgehog protein, for example, induces Dpp expression over a few cell diameters, whereas the Dpp signal produced within these cells influences cells as many as 20 or more cell diameters away.

At the level of gene regulation, the response to signaling inputs may be graded or threshold. In addition, some genes are activated at high concentrations of signal, whereas others are activated at a wider range of concentrations. The differential response of genes can create nested patterns of gene expression along a given axis of the wing field. For example, the *spalt* gene is activated at high Dpp levels, the *omb* gene at lower Dpp levels, and the *vg* quadrant element at still lower levels; consequently, the three genes are expressed in nested, partially overlapping domains centered on the A/P axis (Fig. 3.14). The different responses of target genes to levels of the Dpp signal fit the classic description of a morphogen.

**Figure 3.14**

*Activation of target genes by the Dpp morphogen gradient*

The Dpp protein is expressed along the anterior/posterior compartment boundary of the wing, and spreads from this source to establish a concentration gradient. The *spalt* (purple), optomotor-blind (*omb*; yellow), and *vg* (red) target genes are activated in nested patterns centered along this boundary. The differential response of target genes to a common signal is critical to pattern formation in cellular fields.

Source: Courtesy of Jaeseob Kim.
The action of the Vg/Sd selector genes

Because the various signaling pathways deployed in the wing field are active elsewhere in the body, regulatory mechanisms must exist to impart specificity to their activities in any given tissue. This role is fulfilled by the field-specific selector genes. In the wing, the Vg and Sd proteins are required to permit appendage formation and to effect a wing-specific response of target genes to particular signals. The molecular basis for the Vg/Sd selector function is as follows: Sd acts as a sequence-specific DNA binding protein that binds to *cis*-regulatory elements that control patterns of target gene expression in the wing. Sd forms a complex with Vg; in this complex, the Vg protein acts as a transcriptional activator. Importantly, however, the binding of Vg/Sd is not sufficient to activate target genes. A second cue is required from one or more signaling pathways or another regulatory input. This combinatorial regulation by the selector and signaling input restricts gene activity to particular cells within the wing field (Fig. 3.15).

Many genes that are expressed in discrete patterns in the wing field are also expressed in entirely different patterns in other fields. The wing-specific aspects of these genes' expression are controlled by independent wing-specific *cis*-regulatory elements that respond to and contain binding sites for Vg/Sd and at least one other input. Other *cis*-regulatory elements

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**Figure 3.15**

Integration of signaling and selector protein inputs by *cis*-regulatory elements

Target genes are activated in the wing field by a combination of inputs from one or more signal transducers and the Vg/Sd selector proteins. Localized activation of the Hedgehog (left), Notch (middle), or Dpp (right) signaling pathways induces target gene expression (red) where the signaling input and selector input (Vg/Sd; blue stippling) overlap. Each target gene possesses wing-specific *cis*-regulatory elements that integrate inputs from the appropriate signal transducer (Cl, Mad/Medea, Su(H); see Table 2.2) and Vg/Sd.
control gene expression elsewhere in the body. The modularity of the various cis-regulatory elements allows for the independent regulation of genes in the wing field and in other regions of the body.

The modification of regulatory hierarchies in secondary fields by Hox genes

The regulatory events in the growing wing field described previously transpire in the absence of Hox gene activity. Similarly, the patterning of the antennal field does not involve any Hox gene. On the other hand, the development of the haltere (the serial homolog of the wing) and the serial homologs of the antenna (which include all limb-type appendages) does involve Hox genes. The Hox proteins act to differentiate these structures by modifying developmental programs between serially homologous fields.

Before we discuss how Hox genes modify these complex morphologies and regulatory hierarchies, it is useful to briefly summarize our understanding of Hox protein functions. All Hox genes encode homeodomain proteins with similar DNA-binding specificities. One way that Hox proteins exert specific regulatory control is by interacting with protein cofactors. The Extradenticle (Exd) protein, another homeodomain-containing protein, is a key cofactor that interacts with several Hox proteins. Certain Hox-regulated cis-regulatory elements contain both Hox and adjacent Exd protein binding sites; occupancy of both sites and interaction between Hox and Exd molecules are required for gene regulation.

Hox proteins can act as both transcriptional activators and repressors, with their precise roles depending on a number of as yet unknown variables. In addition to cofactor interactions, post-translational modifications (such as phosphorylation) and the context of Hox binding sites can influence Hox activity. The core recognition sequence for Hox proteins is only about 6 bp long, so the genome contains many low- and high-affinity sites for these proteins.

The striking feature of Hox genes is that their expression in a new position can be sufficient to transform one structure into another. For example, expression of Antp in the antenna, where it is normally absent, causes the development of a leg. Similarly, expression of Ubx in the wing transforms that structure into a haltere. How can changes in a single gene so radically alter development? Do Hox proteins act globally to modify the expression of genes at the top of regulatory hierarchies? Or do they act throughout hierarchies, modifying some regulatory interactions but not others?

The picture that has emerged is one depicting Hox genes as "micromanagers" that act at many levels of regulatory hierarchies on selected components. The development of the haltere, for example, depends on the same major regulators (apterous, engrailed, hedgehog, vestigial, and so on) as its serial homolog, the wing. Yet, the morphology of the haltere is dramatically different in terms of its smaller size, balloon shape, lack of veins or large bristles, and details of cell architecture. The Ubx protein modifies the wing-patterning hierarchy to shape the development of the haltere by acting on a selected subset of genes that influence features of wing pattern formation (Fig. 3.16). For example, Ubx suppresses the production of the Wg protein in the posterior compartment of the haltere disc. This protein also represses AS-C genes, resulting in the suppression of bristle formation at the edge of the haltere. Along the A/P axis of the haltere, certain Dpp-regulated genes are specifically repressed by Ubx, while others are expressed in patterns similar to those in the wing.
The Ultrabithorax-regulated hierarchy in the Drosophila haltere

The haltere is a serial homolog of the wing. The Ubx protein regulates the morphological differentiation of the haltere by selectively modifying the wing regulatory hierarchy at many levels. **(top)** Genes or regulatory elements that are Ubx-regulated in the haltere are shown in red. **(bottom)** The expression patterns of genes in the wing and haltere imaginal discs. (top row, left to right) engrailed, apterous, and dpp expression are similar in the two fields. (bottom row, left to right) spalt, vg quadrant enhancer, and achaete expression differ between the wing and haltere discs due to the actions of Ubx.

How is the selective regulation of genes controlled by Ubx in the haltere? Once again, the key is \textit{cis}-regulatory elements. Recall that wing-specific patterns of gene expression are controlled by discrete elements. In genes that are directly controlled by Ubx in the haltere, wing-specific \textit{cis}-regulatory elements also contain functional sites for the Ubx protein, which generally acts to repress wing-patterning genes in the developing haltere field. \textit{The modularity of cis-regulatory elements enables the selective and differential regulation of gene expression by Hox proteins between serially homologous fields.}

\section*{THE VERTEBRATE BODY PLAN}

The development of the vertebrate body plan has long been a focus of experimental embryology. Many fundamental concepts such as organizers, fields, and morphogens were derived first from observations of vertebrate embryos. The major features of adult vertebrate morphology, including segmented vertebral columns, paired appendages, and skulls, have undergone considerable evolutionary diversification. Therefore, we will focus on the developmental genetics of these major features here, and consider their evolutionary origins and modification in subsequent chapters.

Most vertebrate embryology has focused on a few amphibian (\textit{Xenopus}, newts), avian (chick, quail), mammalian (mouse), and fish (zebrafish, medaka) species. The early development of these embryos and the mechanisms that orient the primary axes differ considerably, particularly between species with yolky eggs (for example, amphibians, fish) and species that rely upon extraembryonic structures. Nevertheless, all early vertebrate development converges on an embryonic stage in which the head is distinct, a neural tube extends along the dorsal midline above the \textit{notochord}, and part of the mesoderm is subdivided. At this stage, the major anteroposterior axis (often called the “rostrocaudal” axis) is well defined, the general rostrocaudal domains of major head, trunk, and tail regulatory genes are initially determined, and the secondary fields (for example, limb buds) that will give rise to appendages and other structures are set to emerge. In the following discussion, we will see that the developmental regulatory logic and mechanisms controlling the patterning of vertebrate body axes and cellular fields revisit some already familiar themes mentioned earlier in this chapter.

\textbf{The rostrocaudal axis: somite formation and segmentation}

The mesoderm of vertebrates gives rise to many organs and to the tissues that form major features of vertebrate body architecture. The mesoderm is subdivided into several regions that give rise to different tissues (Fig. 3.17a,b). The \textit{lateral plate mesoderm} is the source of elements of the circulatory system, the lining of body cavities, and all of the mesodermal components of the limbs (except the muscles). The \textit{paraxial mesoderm} is subdivided into metameric subunits called \textit{somites}. The somites provide the framework for the metameric organization of somite-derived tissues, including the axial skeleton (vertebrae and ribs), the dermis and muscles of the back, and the skeletal muscles of the body wall and limbs. Because somite-derived structures such as the vertebrae and ribs have characteristic shapes in different regions of the rostrocaudal axis, the generation and individuation of the somites and their derivatives are fundamental to the evolution and diversification of the vertebrate body plan.
Segmentation of the paraxial mesoderm and somite formation involve a sequential, temporally ordered process that proceeds from the anterior to the posterior of the embryo. New somites are formed from the rostral end of the paraxial mesoderm at regular intervals. In the mouse and the chick, a defined, albeit different number of somites are produced, at roughly 90-minute intervals, whereas in zebrafish a new somite forms every 20 minutes.

Two regulatory systems have been identified that underlie the temporal and spatial regulation of somite formation. One system behaves as a segmentation clock, involving the oscillating expression patterns of genes in the developing somites. The second system translates the clock into a periodic arrangement of segment (somite) boundaries.
The best characterized set of genes whose expression oscillates during somite formation include relatives of the *Drosophila hairy* and *Enhancer of Split* gene family (HES/HER genes), a modulator of the Notch receptor protein *Lunatic fringe*, and the Notch ligand deltaC. These genes are expressed in a dynamic wave that sweeps across the presomatic mesoderm from posterior to anterior once during the formation of each somite (Fig. 3.17c). Oscillating gene expression and somite formation in all vertebrates appears to be regulated by the Notch pathway, although the roles of individual genes vary in different species. The synchronized expression of these genes suggests that they lie downstream of a common cycling activator. The translation of oscillating transcription factor expression into segment boundaries is regulated by the Fgf8 signaling molecule. Fgf8 is expressed in a graded manner in the presomitic mesoderm (PSM) and the level of expression is important for regulating the position of somite boundaries. The integration of these two systems together with antero-posterior patterning information, including *Hox* genes, creates the unique identities along the vertebrate segmented body plan.

**The vertebrate Hox ground plan**

The somites of the vertebrate embryo give rise to the major axial structures, including the vertebrae, ribs, and skeletal muscles, as well as the dermis. The vertebrae of the spinal column and their associated processes are of five distinct types: cervical, thoracic, lumbar, sacral, and caudal (Fig. 3.18a). The transitions between these types occur at specific somite positions in each species. Similarly, the forelimb and hindlimb buds, which develop from the unsegmented lateral plate mesoderm, arise at specific axial positions. Therefore, the morphologies of somite derivatives at different axial levels must be genetically regulated.

Patterning along the rostrocaudal axis in all vertebrates is regulated by the *Hox* genes. Vertebrates have four or more complexes of *Hox* genes; the mouse, for example, has 39 *Hox* genes. In general, the anterior boundaries of expression of the *Hox* genes correlate with the respective locations of the genes within a complex, a principle termed *colinearity*. Thus the *Hox1* genes (which are related to the *labial* gene) are expressed in the most anterior positions, and the *Hox9-13* genes (which are related to the *Abd-B* gene) are expressed in more posterior positions (see Fig. 2.22b). The anterior boundaries of *Hox* gene expression are generally sharper than their posterior boundaries and lie in spatial register between the somitic and lateral plate mesoderm. However, they are not in register with *Hox* gene expression in the neural tube. The tissue-specific boundaries of *Hox* gene expression indicate that independent genetic regulatory mechanisms control *Hox* expression in the neural tube.

Patterning of part of the neural tube, the posterior region of the head, and the hindbrain also involves segmentation along the A/P axis. The process of hindbrain segmentation is evolutionarily conserved in vertebrates in terms of the number of subdivisions or rhombomeres formed (seven to eight), the neuroanatomical organization of each region, and the pattern of expression of various genes, including transcription factors and signaling proteins that establish rhombomere segmentation, identity, and cell behavior. The best-known regulatory genes involved in this process are the *Hox1* through *Hox4* genes, various members of which are expressed in and affect the development of discrete rhombomeres (Fig. 3.18a,b).

The same themes pertain to *Hox* regulation in the vertebrate hindbrain (and elsewhere) as were highlighted for the *Drosophila Hox* genes.
Figure 3.18
Regulation of *Hox* gene expression in the vertebrate hindbrain

(a) The vertebrate hindbrain forms anterior to the spinal cord. (b) The hindbrain is overtly segmented into rhombomeres (r1–r7). (c) Summary of regulatory gene expression in r1–r7. The arrows denote regulatory interactions between two transcription factors (Kreisler and Krox20) and two *Hox* genes of interest. (d) A schematic of the genetic region around the *Hoxa*2 and *Hoxa*3 genes. Specific cis-regulatory elements (R3 and R5) control the response of *Hoxa*2 to Krox20 in r3/r5. Note the presence of several other elements controlling *Hox* expression in other tissues. This diversity of cis-elements is typical of *Hox* genes.

Numerous independent cis-acting regulatory elements control features of individual Hox gene expressions in different rostrocaudal domains and in different germ layers.

Both positive and negative regulatory interactions are necessary to define the initial domains of gene expression.

Autoregulatory and cross-regulatory interactions between Hox genes are necessary to maintain domains of gene expression.

In the developing mouse hindbrain, a few direct transcriptional regulators of Hox gene expression patterns have been identified, including the products of the Krox20 and Kreisler genes, and products of Hox genes themselves. In addition, cis-regulatory elements have been identified that control rhombomere-specific patterns of Hox gene expression (Fig. 3.18d). For example, expression of the Hoxa2 and Hoxb2 paralogs in rhombomeres 3(r3) and 5(r5) are controlled by distinct elements, independent of those that control expression in the mesoderm and other tissues. The Krox20 protein is specifically expressed in these two rhombomeres and directly regulates the r3/r5 enhancers (Fig. 3.18c). Similarly, expression of the Hoxa3 and Hoxb3 genes in r5/r6 is controlled by discrete elements. The Kreisler protein is specifically expressed in these developing rhombomeres, and it binds to and directly regulates Hoxa3 and Hoxb3 expression through these elements (Fig. 3.18c).

Expression of the Hox genes is coupled to the segmentation clock, providing coordination of somitogenesis and the specification of regional identity. But apart from these rhombomere patterns, not much is known about the transcription factors that position Hox domains along the rostrocaudal axis. In addition, from the cis-regulatory elements of the Hox complexes, we can infer that the regulatory network likely involves many interactions. When the intergenic regions of a few pairs of Hox genes were analyzed in greater detail, a substantial number of regulatory elements were found interspersed among the Hox genes. Separate cis-regulatory elements that control gene expression in the lateral mesoderm, neural crest, spinal cord, tail bud, limb buds, and somites have been identified, for instance (Fig. 3.18d). The temporal, spatial, and tissue-specific aspects of each pattern must be regulated by a variety of transcription factors acting through these elements. The different anterior expression boundaries of adjacent genes within complexes could reflect the differential response of cis-regulatory elements to the same regulator or the specific response of each element to different regulators. Interestingly, some of the cis-acting regulatory elements are shared between pairs of adjacent genes, and this sharing may be a mechanism that preserves the clustering of Hox genes.

**Vertebrate limb development**

Among the most important structures in vertebrate evolutionary history are the paired pelvic and pectoral appendages. The evolution of the tetrapod limb from the paired fins of fish and the subsequent modification of limb morphologies used in flying, running and jumping, burrowing, and the return to water involved major changes in the anatomy of paired appendages. Coupled with the longstanding research on the experimental embryology of limb buds, immense interest is focused on the developmental genetics of vertebrate limb formation and patterning. For our purposes, there are six major aspects of limb formation and patterning, each of which represents a potential point of divergence between taxa in the functional evolution of limb morphology:
- The positioning of the limb bud along the rostrocaudal axis
- The initial outgrowth of the limb bud
- Establishment and signaling from the limb organizers
- The formation of the limb proximodistal axis and deployment of Hox gene expression in the limb bud
- The specification of major skeletal and cartilage elements
- The regulation of forelimb and hindlimb identity

These developmental processes unfold as a cascade of events that progressively establish the limb field, its two organizers, and many pattern elements (Fig. 3.19).
The developmental regulatory logic involved in these processes of limb formation and patterning are analogous to those discussed for insects. The position of the limb field depends on signals whose distribution is regulated along the main body axis. The outgrowth of the limb field involves regulatory hierarchies that establish the anteroposterior and dorsoventral axes of the limb field, and integration of these inputs regulates patterning along the proximodistal axis. Superimposed on the potentially equivalent hierarchies in the forelimb and hindlimb fields are selector genes that specifically modify the development of these limbs.

**Positioning of the limb buds along the rostrocaudal axis**

Limb development begins with the proliferation of mesenchymal cells from within the limb field of the lateral plate mesoderm. As these cells accumulate under the epidermis, a bulge appears—the growing limb bud. This proliferation appears to be under the control of local signals emanating from nearby mesoderm. One strong candidate for the source of this proliferative signal is fibroblast growth factor 10 (FGF10), which is expressed near the site of limb bud initiation and can induce the formation of ectopic limb buds when expressed at novel positions along the rostrocaudal axis. Although *Fgf10* is initially expressed throughout the lateral plate mesoderm, it becomes restricted to the presumptive limb areas. Expression in these regions is maintained by a combination of specific Wnt signaling molecules (members of the signaling protein family to which the insect Wg protein belongs) and T-box (Tbx) transcription factors (Fig. 3.20a).

Limbs of different vertebrates arise at different positions, with respect to somite number, along the rostrocaudal axis. Nevertheless, the forelimb buds always arise at the most anterior position of *Hoxc6* expression at the transition from the cervical vertebra to the thoracic vertebra. This consistent location suggests that the positioning of the limb bud is determined by both axial coordinates (*Hox* genes) and local expression of specific *Wnt* genes.

**Outgrowth of the limb bud: dorsoventral and anteroposterior regulatory hierarchies**

The early limb bud consists of mesodermal cells and an overlying ectodermal epithelium. As it forms, the mesodermal cells induce the overlying ectoderm to express *Fgf8* (Fig. 3.20b) and these cells will eventually form the apical ectodermal ridge (AER), a prominent thickening of cells at the edge of the limb bud where its dorsal and ventral halves meet. The AER organizer is a major signaling center that produces signals such as FGF8 (and other related FGFs), which are necessary for the continued proliferation and differentiation of the growing limb bud. Early loss of the AER, or its signaling functions, prevents outgrowth of the buds and limb formation.

Induction and maintenance of the AER as well as dorsoventral patterning of the limb depends upon the interplay between several signaling pathways in the ventral limb ectoderm (Fig. 3.21). FGF10 signals from the limb mesoderm to the overlying ectoderm and induces *Fgf8* expression in the early ventral limb ectoderm, or pre-AER cells. This induction is mediated by the BMP and Wnt signaling pathways (Fig. 3.21b).

These same two signaling pathways also regulate the expression of *En-1*. *En-1* regulates dorsal patterning of the limb, which occurs later during limb bud outgrowth. *En-1* represses expression of the Wnt-7a protein, which is then expressed only in the dorsal ectoderm and
Figure 3.20
Regulation of vertebrate limb bud formation

Dorsal view of a vertebrate forelimb-forming region showing genes involved in (a,b) limb initiation and (c,d) A/P patterning. Rostral is to the top and distal is to the right. (a) The forelimb bud is induced by Wnt2b expression in the chick lateral plate mesoderm. In response to Wnt2b signaling, Tbx5 expression is activated, which maintains Fgf10 expression in the presumptive forelimb region. (b) Fgf10 signals to the overlying ectoderm and, through Wnt3a, induces Fgf8 expression. These Fgf8-expressing cells will give rise to the AER and maintain proliferation of the bud. (c) In the absence of Hedgehog signaling, GLI3 exists in a repressor state (GLI3R). Mutual antagonism of GLI3R and HAND2 prepatterns the limb mesenchyme prior to SHH signaling. (d) Shh is induced in the posterior of the limb bud by HAND2 and restricts GLI3R activity to the anterior mesenchyme. The two signaling sources (ZPA, AER) are maintained by reciprocal interactions involving Shh, Fgfs and the BMP antagonist Gremlin (GRE).

regulates dorsoventral polarity. The actions of Wnt-7a appear to be mediated by the Lmx1b protein, a homolog of the *Drosophila apterous* gene that is expressed in all dorsal mesodermal cells of the limb (Fig. 3.21b).

Patterning along the A/P axis of the limb bud depends on the function of the zone of polarizing activity (ZPA), another organizer with potent signaling activity. The ZPA is the source of the Sonic hedgehog (Shh) protein in the limb field. The ZPA and Shh influence the polarity of pattern elements along the A/P axis. In particular, digits with reversed polarity can be induced when either the ZPA is transplanted or Shh expression is induced in the anterior part of the limb.

The restriction of the ZPA and Shh expression to the posterior portion of the limb bud suggests that the limb bud contains anteroposterior positional information that predates Shh expression and function. This preexisting information may consist of rostrocaudal regulatory patterns derived from the lateral plate mesoderm. Two proteins, the basic helix-loop-helix transcription factor HAND2 (dHAND) and Gli3, a homolog of the *Drosophila* *Cubitus interruptus* protein and a transcriptional effector of Hedgehog signaling play critical roles in anteroposterior patterning of the limb. Gli3 is expressed in the anterior limb mesoderm and HAND2 in the posterior limb mesoderm, and they each act to restrict each other’s expression (Fig. 3.20c). HAND2 regulates the activation of Shh expression in the posterior distal mesoderm. Therefore, Shh expression in the ZPA appears to be regulated both by signals from the AER and inputs from the lateral plate mesodermal cells that will become incorporated into the limb bud.
The organizers in the limb field do not act independently of one another. Formation and maintenance of the ZPA depends on the presence of a functional AER, and, in turn, maintenance of the AER depends on the existence of ZPA function. Thus a regulatory circuit connects FGF signaling to the induction of Shh expression, and Shh signaling to FGF expression in the AER (Fig. 3.20d). These inductive interactions appear to operate through intermediate signals, although all of the proteins are not known as of yet.

**Integration of organizing signals to form the proximodistal axis and deployment of Hox genes in the limb field**

After the formation of the ZPA and AER and the proliferation of mesodermal cells, proximal mesenchymal cells begin to form cartilaginous condensations that prefigure the limb skeletal pattern. These condensations arise in a proximal to distal order, so that the humerus (the bone of the upper arm) forms first, the radius and ulna next, and the carpals (wrist bones) and digits last. The development of these individual pattern elements suggest that limb development goes through discrete temporal phases and that pattern formation is controlled by the localized expression of regulatory genes, cued by the signals emanating from the organizers, the AER and the ZPA.

The Hox genes play important roles in limb patterning, albeit in a different fashion than in appendage patterning in arthropods. Whereas individual Hox genes or pairs of Hox genes are expressed in arthropod limb fields, a much larger number of Hox genes are deployed in nested domains of vertebrate limb fields (Fig. 3.22). Detailed studies of normal and ectopic Hox gene expression and analysis of limb development in mice lacking one or more Hox gene functions have revealed a complex and dynamic spatiotemporal pattern of Hox gene patterns and gene interactions in determining the formation of limb pattern elements.

Three temporal phases of Hox gene expression appear to correlate with the temporal sequence of proximodistal pattern element formation, particularly in regard to the Hox9-13 genes of the HoxA and HoxD complexes (Fig. 3.22a–c). The first phase of Hox expression is not polarized, appears to be Shh-independent, and is associated with the development of the most proximal limb elements (upper arm/leg) (Fig. 3.22a). Subsequent phases of Hox expression arise in nested patterns whose polarity depends on AER and ZPA functions. The second phase of Hox expression occurs in the next most proximal elements (forearm or lower leg) (Fig. 3.22b). The third phase of Hox expression includes most distal elements (wrist and hand, ankle and foot) (Fig. 3.22c).

These spatial patterns of Hox gene expression reveal that many related Hox proteins are expressed in overlapping as well as adjacent domains. No simple one-to-one correspondence exists between a particular Hox gene and the growth and patterning of any particular limb pattern element. Rather, the formation and identity of particular elements appear to reflect combinations of Hox protein functions, some of which are clearly acting redundantly. For example, loss of either Hoxa-13 or Hoxd-13 function has limited effects, whereas loss of both genes dramatically affects the development of the distal limb. Similarly, Hoxd11 or Hoxa11 mutations have minor effects on the formation of the radius and ulna, whereas these long bones are almost completely lost in the double mutant.

The analysis of cis-acting regulatory elements that control Hox gene expression in the limb bud has highlighted the complexity of Hox expression. Both gene-specific and global regulatory elements appear to control the expression of individual Hox genes and groups of Hox
Figure 3.22

Hox gene expression and function in vertebrate limb development

(a–c) Three temporal phases of Hox gene expression in the limb bud correlate with the elaboration of three distinct elements of the limb. (a) Phase I involves a more limited set of Hox genes expressed across the entire bud in the region that will give rise to the upper arm or thigh (stylopod). Mutations in these genes primarily affect this structure. (b) Phase II expression is initiated in response to Sonic hedgehog signaling and is polarized with respect to the A/P axis of the limb bud. Disruption of multiple Hox genes that have pronounced phase II patterns affects formation of the bones corresponding to the lower arm or calf (zeugopod). (c) Phase III expression occurs in the hand and foot (autopod). Disruption of certain Hox genes primarily affects autopod formation and patterning. (d) The cis-regulatory elements of the Hoxd9-13 genes. Both global and local elements regulate Hox expression in the limb bud. Multiple Hox genes are influenced by remote elements active in the zeugopod and autopod. Local elements, such as RXI (red), affect expression of individual genes in restricted domains.

genes in the limb bud, respectively. For example, phase II expression of the \textit{Hoxd12} gene is regulated by both a nearby element and a remote global element, whereas phase III expression is controlled by a remote global element (Fig. 3.22d). Thus the complex patterns of \textit{Hox} gene expression are built up from many potentially independent inputs, which creates the potential for morphological diversification through changes in the relative timing and spatial regulation of individual elements.

**Regulatory networks controlling the differentiation of major limb pattern elements**

The skeletal defects caused by \textit{Hox} mutations indicate that one function of \textit{Hox} proteins may be to regulate the position and timing of cartilage development and subsequent growth and differentiation in the limb field. Evidence suggests that multiple signaling molecules (Bmps, Wnts, Fgfs, another member of the Hedgehog signal family, Indian hedgehog-Ihh), Sox transcription factors, and \textit{Hox} genes regulate the progression of bone differentiation and thus major features of limb pattern formation. Certain \textit{Hox} proteins appear to act early in the differentiation pathway to prevent its progression, either by promoting the proliferation of bone forming cells or by preventing their differentiation.

The positions of the joints between limb skeletal elements are also spatially regulated. One regulator of this process, the GDF5 signaling protein (a member of the large TGF-\beta superfamily), is specifically expressed in regions where bone development does not occur. GDF5 expression presages the location of joints throughout the limb field (as well as elsewhere in the body). Clearly, the regulation of the spatial expression of these various cartilage and bone promoting and inhibiting factors is key to the skeletal pattern formed.

**The regulation of forelimb and hindlimb identity by selector genes**

The serially homologous tetrapod forelimb and hindlimb are believed to have evolved from the paired pectoral and pelvic fins of fish, respectively. Although the developing forelimbs and hindlimbs both utilize organizing signals that are necessary for appendage outgrowth, no single transcription factor required for limb outgrowth has been identified as yet. The \textit{Hox} genes, for example, do not play the role of selector genes for limb field formation or identity. No \textit{Hox} mutations result in homeotic transformations between limb types.

A few other transcription factors have, however, been identified as selectors of forelimb and hindlimb identity in vertebrate limbs. The paired homeodomain-containing protein \textit{Pitx-1} and the T-box-containing protein \textit{Tbx4} are expressed specifically in the hindlimb mesenchyme, and the T-box gene \textit{Tbx5} is expressed specifically in the forelimb mesenchyme (Fig. 3.23). In addition to the early requirement of these genes for limb outgrowth, the differentiation between forelimb and hindlimb identity is regulated, at least in part, by these selector genes. Although each is required for proper identity, none of the three genes is sufficient by itself to confer limb-specific identity. Hence, additional regulators of limb identity may act upstream of these genes, or perhaps limb identity results from a number of field-specific inputs.

These and other potential field-specific regulators presumably function to control the differential expression of genes between the two fields (Fig. 3.23). Several \textit{Hox} genes, including \textit{Hoxc10} and \textit{Hoxc11}, are differentially expressed between limb types and are potential
targets of the limb-specific selectors. Given the important morphological and functional diversity of vertebrate limb morphologies, it will be important to characterize the levels in the limb regulatory hierarchy at which changes have arisen between taxa.

**REVIEW: THE GENERAL LOGIC AND MECHANISMS CONTROLLING GENE EXPRESSION IN CELLULAR FIELDS**

This chapter has presented an overview of the genetic regulatory logic and mechanisms that operate to control development in a few model species. It concentrated on the genes of the general animal toolkit for development as well as the mechanisms that control large-scale patterning of the main body axes and secondary fields. Although we explored only a modest number of regulatory hierarchies in representatives of just two phyla, the similarities in the regulatory logic and mechanisms allowed us to identify some general themes concerning the regulation and function of toolkit genes and the architecture of the regulatory hierarchies that progressively specify pattern formation in cellular fields. The identification of these general themes is critical for understanding trends in animal evolution.

Because the regulatory mechanisms of development are themselves the product of evolution, their architecture is a reflection of the evolutionary processes that assembled them. Armed with our knowledge of regulatory architectures in one species, we can begin to make comparisons with other taxa with different morphologies to identify at what level and through what genetic mechanisms developmental and morphological diversity arises.
From this survey of regulatory hierarchies and mechanisms, we underscore three general themes with regard to the architecture of regulatory hierarchies, the molecular mechanisms controlling gene expression, and the functions of toolkit genes in controlling gene expression and patterning in cellular fields:

1. *The development of the growing embryo and its body parts occurs in a spatially and temporally ordered progression through the sequential generation of coordinate systems.* The position of any adult pattern element is determined through a series of hierarchies that subdivide the embryo and its morphogenetic fields into progressively finer elements. From the coordinate systems that subdivide the major embryonic axes, secondary fields are specified that contain their own autonomous coordinate systems. Domains within these fields can be further subdivided and specified through the establishment of local domains of gene expression. Ultimately, the temporal and spatial segregation of these processes allows for the modular organization of bilaterians and the individualization of body parts and pattern elements, such that morphology evolves independently from the rest of a field or the body plan.

2. *The modularity of cis-regulatory elements acting on individual genes allows for the independent spatial and temporal control of discrete features of gene expression and function.* Many gene expression patterns are actually the sum of the functions of many independent cis-regulatory elements. The modularity of cis-regulation is crucial to the control of the specificity of gene interactions and function during development. The independent spatial and/or temporal regulation of gene expression permits individual genes to have different but specific functions in different contexts. Thus, while operating through identical signal transduction pathways, a signaling protein can be expressed in entirely unrelated populations of cells in two different tissues (by virtue of its own cis-elements) and can regulate the expression of completely different target genes (through different cis-regulatory elements of its target genes).

   In this light, it is not adequate or accurate to describe a given toolkit gene function solely in terms of the protein it encodes, because the function of that protein almost always depends on the context in which it is expressed. Instead, toolkit genes should be viewed as consisting of both a functional protein and a suite of regulatory elements that control its deployment. Each element represents a separate genetic function.

   The modular organization of gene-specific control elements has two profound evolutionary implications. First, this modularity is the product of evolution and therefore directly reflects the mechanisms that allow new and independent patterns of gene expression to evolve. Second, the modular control of extant genes allows for changes in one aspect of gene expression and function without affecting any other functions. In other words, modularity facilitates the dissociation of gene functions as well as the evolution of new interactions and potential morphologies.

3. *Spatial patterns are the product of combinatorial regulation* In the examples of regulatory hierarchies and individual genes expression patterns analyzed in this chapter, we saw that new patterns were created by the combined inputs of preceding patterns. The integration of D/V and A/P inputs to position primordia, the positioning of a pair-rule stripe, the positioning of organizers, and the field-specific pattern of a regulatory gene, for example, are all derived from the integration of multiple inputs (some positive, some negative) by cis-regulatory elements.
Combinatorial control is key to both the specificity and the diversity of gene expression. In terms of specificity, it provides a means to localize gene expression to discrete cell populations using inputs (such as signaling pathways) that are not cell-type-specific or tissue-specific. In terms of diversity, combinatorial mechanisms provide a means to generate a virtually limitless variety of spatial patterns through the overlapping inputs of positive and negative regulators and autoregulatory feedback mechanisms. As the number of discrete domains of gene activity increases in a field, the potential combinations of regulators and patterning outputs increase exponentially.

The evolutionary significance of combinatorial regulation is obvious. New gene expression patterns can evolve as new combinations of regulatory inputs are integrated. In turn, the new gene expression patterns create the potential for further change, and the cycle continues.

From these general themes, we can start to anticipate how regulatory hierarchies and gene expression patterns might be cobbled together in the course of evolution and how they might change. To begin to understand what has occurred, we must take a broad inventory of the genetic toolkit across the animal kingdom and examine the regulatory mechanisms for building animals in a phylogenetic and comparative context to see how morphological diversity evolves.

SELECTED READINGS

Gene Regulation in Metazoans


From Egg to Segments: The Anteroposterior Coordinate System


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### The Dorsoventral Axis Coordinate System


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### The Hox Ground Plan


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### Secondary Fields: Integrating the Anteroposterior and Dorsoventral Coordinate Systems


**The Wing Field**


**Selector Proteins and Target Gene Regulation**


### Hox Genes’ Modification of Regulatory Hierarchies in Secondary Fields


### Vertebrate Somite Formation and Segmentation


The Vertebrate Hox Ground Plan


Vertebrate Limb Initiation and Early Development


**Patterning Along the Vertebrate Limb Axes**


CHAPTER 4

Evolution of the Toolkit

The presence of many similar developmental regulatory genes among long-diverged and vastly different animals such as the mouse and the fruit fly raises an apparent paradox. If bilaterians share a related set of developmental genes, how has morphological diversity evolved? There must be genetic differences between animal lineages that direct the development of different animal morphologies. One potential source of genetic differences among animals is the coding content of their genomes, including the number and biochemical functions of toolkit genes. Another possible source is the way that toolkit genes are used during development, including both the timing and pattern of their expression and their interactions with other developmental genes in regulatory circuits and networks.

This chapter focuses on the assembly and expansion of the genetic toolkit for development during animal evolution and the role of gene evolution in the origins of morphological complexity. The number of genes contained within an evolving genome is dynamic, and gene duplication increases the information content and potential complexity of developmental programs. Representatives of large gene families are shared among bilaterians, and these families continue to evolve new members in different animal lineages. The expansion of gene families during animal evolution can be traced by comparing the genes found in extant (living) organisms and by mapping gene duplication events relative to animal phylogeny. In particular, organisms that are basal (most deeply branching) members of a clade indicate the state of the toolkit before the evolution and radiation of more recent groups.

Analysis of gene families reveals two periods of major genomic change during animal evolution that correlate with the emergence of more complex animal forms. One interval occurred at the transition to triploblastic bilaterians early in the animal tree; the other is found at the base of the vertebrate lineage. By contrast, the content of the genetic toolkit for development appears roughly equivalent among other morphologically disparate bilaterian phyla.

Species and groups of species which are called aberrant, and which may fancifully be called living fossils, will aid us in forming a picture of the ancient forms of life. Embryology will often reveal to us the structure, in some degree obscured, of the prototypes of each great class.

—Charles Darwin, The Origin of Species (1859)
Thus expansion of the toolkit correlates with increased animal complexity, but not with diversity.

Evolutionary changes in the proteins that make up the toolkit are more often not the primary source of genetic differences that underlie the morphological diversity of animal forms. Indeed, some homologous genes from long-diverged animal phyla share similar biochemical functions. More significantly, some toolkit genes have similar developmental functions, and these similarities may be used to make inferences about the possible anatomy of animal ancestors, even without fossil evidence. The shared developmental functions among disparate animal phyla suggest that these toolkit genes controlled the development of various anatomical features in the last common ancestor of bilaterians.

**THE HISTORY OF GENE FAMILIES**

*Conservation of developmental regulatory genes: inferences about animal ancestors from comparative genomics*

Animal genomes contain many thousands of genes, many of which are required for basic processes common to cellular life. These “housekeeping” genes are shared by most living organisms and predate the evolution of multicellular animal life. Because housekeeping genes are fundamental to cell structure, viability, and function, they are not the most likely candidates for the genes critical to the evolution of complex animal body plans. Instead, this chapter focuses on the subset of genes that controls patterning and differentiation—a namely, the components of the genetic toolkit for development. Previous chapters have described the developmental functions and interactions of many toolkit genes in the model organisms such as *Drosophila* and the mouse. But how and when did this toolkit of genes evolve? And how can we hope to understand the early evolution of toolkit genes, given that the ancient animals that carried those genes are now extinct?

Much of our understanding of the evolution of the genetic toolkit for development is based on deductive logic, using information from the genomes of extant organisms (including mammals such as humans and mice, the insect *Drosophila*, the nematode worm *Caenorhabditis elegans*, the yeast *Saccharomyces cerevisiae*, and some plants) to extrapolate to the past. Similarity between gene sequences found in two (or more) different organisms is most easily explained by common history. In other words, a gene that is conserved among a group of animals was present in the last common ancestor of that group. During the subsequent independent evolution of each animal lineage from the last common ancestor, the gene sequence diverged to create related (but not identical) genes, one in each species. Genes with similar sequences that are found within a single animal genome are also related, as they are products of the duplication and divergence of ancestral genes.

The knowledge of the complete genome sequence of a growing list of animals allows the comparison of extant genomes and the reconstruction of ancestral genomes. The systematic comparison of whole genomes provides a history not only of gene sequences but also of repetitive sequences, gene organization, gene order (and rearrangements), gene duplication, and *cis*-regulatory regions, as well as organismal phylogeny. Furthermore, as more genomes are sequenced, it becomes possible to determine ancestral and derived conditions with respect to the number and identity of genes, and to make well-informed inferences about the
Chapter 4: Evolution of the Toolkit

The conservation of toolkit genes in fruit flies and mice discussed in earlier chapters anticipated the findings of comparative genomics, and extends to humans, *Fugu rubripes* (pufferfish), *Ciona intestinalis* (urochordate, basal to the chordate lineage), *C. elegans*, and other bilaterian species (Box 4.1). All of these organisms inherited the basic set of toolkit genes from the last common ancestor of all bilaterian phyla (Fig. 4.1).

The characterization of toolkit genes in some organisms is particularly informative. An animal lineage that branches near the base of a clade represents a basal member or an outgroup which can help to establish the ancestral condition for that clade. For example, the cnidarians are an outgroup to the bilaterian clade, the cephalochordate *Amphioxus* is a close outgroup to the vertebrates, and the onychophora are an outgroup to the arthropods. The resemblance of these animals to their extinct ancestors has led them to be called “living fossils.” The complement of developmental genes shared between these outgroups and representative model organisms reflects the state of the toolkit before the radiation of bilaterians, vertebrates, and arthropods, respectively. As Darwin foreshadowed (see the opening quote for this chapter), they “aid us in forming a picture of the ancient forms of life.”

**Gene duplication**

One major process involved in the assembly of the bilaterian toolkit for development is the duplication and divergence of genes and the creation of gene families. For example, more than 40% of the genes in the nematode *C. elegans* have sequence similarity to other *C. elegans* genes and thus arose at some point from gene duplication events. In fact, tens to hundreds of genes are duplicated in animal genomes every million years, a significant contribution to genome evolution. Duplicated genes, which are often linked in tandem, may arise from slipping errors during DNA replication, errors in the repair of double-strand DNA breaks, or unequal cross-over events during recombination. Some developmental genes are found as closely related, linked gene pairs, indicating that they derive from tandem duplication events. In *Drosophila*, for example, the gene pairs *engrailed* and *Invertoed*, *spalt* and *spalt-related*, and *gooseberry* and *gooseberry-neuro* are all tightly linked. Over time, tandemly duplicated genes may become physically separated through chromosomal rearrangements and translocations.

Such tandemly duplicated genes create a unique opportunity for further expansion of a gene family. Unequal crossing over between mispaired tandem copies leads to one chromosome with a duplication and one chromosome with the corresponding deletion (Fig. 4.2). The chromosome that contains the duplicated region also carries a new chimeric gene, located between the parental copies. If more than two tandemly arrayed members of a gene family exist, unequal crossing over can lead to gene duplication as well as the creation of a new chimeric gene, when the cross-over is out of register by more than one gene. This mechanism of expansion of related tandemly arrayed genes may explain the evolution of the *Hox* complex, for example. New *Hox* genes have evolved as chimeras (or duplicates) of older *Hox* genes, and the *Hox* genes at each end of the complex may represent the “oldest” members of the complex.

Large-scale duplications of chromosomal segments or even of entire genomes (tetraploidy) have also occurred during animal evolution. Such duplication events generate large syntenic blocks, which are duplicated arrays of genes that are found in the same order, distributed throughout the genome. This mechanism of gene duplication rapidly increases
Box 4.1 Identifying and Analyzing Toolkit Genes in Different Animals

The conservation of developmental regulatory genes allows members of gene families to be identified in different animals based on sequence similarity. Historically, molecular biology techniques such as degenerate polymerase chain reaction (PCR) and library screening have facilitated the isolation of homologous genes from both model and nonmodel organisms. Both of these techniques rely on the ability to detect sequence similarity by nucleic acid hybridization between two homologous genes.

Degenerate PCR takes advantage of mixtures of short oligonucleotides that match all possible codon combinations for two conserved peptide sequences within a gene; these mixtures are used to amplify the intervening region from a target pool of nucleic acid (typically, genomic DNA or cDNA). Thus genes that are characterized by conserved protein motifs are good candidates for degenerate PCR, which can rapidly and selectively isolate a region of a gene from any animal’s genome.

Library screening uses a known gene sequence as a probe to isolate other genes with similar DNA sequence. Libraries are pools of randomly isolated, but unsorted, pieces of genomic DNA or cDNA that can be screened multiple times to identify genes of interest. Such screens are particularly useful for isolating genes from nonmodel organisms.

The identification of toolkit genes in model organisms has become an exercise in searching computer databases. The advent of genome sequencing projects provides large databases for sequence comparison. The number, sequence, and chromosomal map position of members of conserved gene families can be rapidly cataloged based on sequence similarity. Internet-based computer resources, some of which are listed below, are the best way to search and analyze genome sequences from model organisms.

The identification of conserved genes from different organisms allows gene sequences to be analyzed for evolutionary relatedness. Many computer programs and tools are available for aligning and comparing the conserved sequences of members of gene families. Large molecular biology Internet servers in the USA and Europe include the following:

- The National Center for Biotechnology Information (NCBI) at the National Institute of Health (http://www.ncbi.nlm.nih.gov)
- The European Bioinformatics Institute, part of the European Molecular Biology Laboratory (http://www.ebi.ac.uk/index.html)
- The ExPASy Molecular Biology Server at the Swiss Institute of Bioinformatics (http://www.expasy.ch)

These sites integrate sequence databases (GenBank, SWISS-PROT), protein family databases (PFAM, Prosite), genome project websites, and other genome analysis tools. Powerful computer programs that are used to generate molecular phylogenies (“gene trees”) include PHYLIP (http://evolution.genetics.washington.edu/phylip.html) and PAUP* (http://www.sinauer.com/Titles/frswofford.htm). These programs build molecular phylogenies of related gene sequences using several computer models of molecular evolution.
Figure 4.1

Metazoan phylogeny

The phylogenetic relationships between many Metazoan phyla have been resolved using molecular data. Three primary bilaterian clades exist: the deuterostomes (including echinoderms, hemichordates, urochordates, cephalochordates, and vertebrates), the arthropod + onychophora + priapulid clade, and the lophotrochozoans (including annelids, molluscs, most flatworms, and lophophorates). The last common ancestor of all bilaterian phyla is indicated in the figure. Basal branches off of the prebilaterian stem lineage lead to the Cnidaria (jellyfish, anemones, coral) and the Porifera (sponges); the position of the nematodes is uncertain.
the total number of genes within a genome. The presence of two, three, or four copies of many
developmental genes and syntenic regions of linked genes in vertebrate genomes provides
evidence of large-scale duplications or tetraploidization events in the vertebrate lineage
(Fig. 4.3). The initial tetraploidy of genes and chromosomes may gradually vanish, as linked
genes become separated or are lost due to chromosomal rearrangements and deletions.

Because the duplication and divergence of genes parallels the branching of animal lineages,
specific terminology has been developed to describe the historical relationships between
gene family members found both within and between different animal genomes. All of the
genes in a given gene family share sequence similarity and hence are termed homologs, as
they share common ancestry. Two important distinctions are made among homolog genes,
based on how they arise during evolution. Genes that are found in different animals and that
arose from a single gene in the common ancestor of those animals are called orthologs. For
example, the divergence of insects has created orthologous labial genes in each insect species.
Genes that arose from gene duplication events in a single genome are called paralogs. For
example, the Hox genes of Drosophila are a complex of paralogous genes.

The difference between orthologous and paralogous genes is most easily depicted by
constructing a phylogenetic tree of related gene sequences. A gene phylogeny depicts the
evolutionary relatedness, or history, of members of a gene family. Figure 4.4 shows the relation-
ship between deuterostome engrailed genes. Each node on this gene tree represents
either an animal lineage bifurcation or a gene duplication. The divergence of animal lineages
created the orthologous engrailed genes found in the sea urchin, amphioxus, and the lamprey.
A gene duplication generated the paralogous En1 and En2 genes found in chicks, mice, and
humans. The En1 genes are more closely related to one another than to any En2 gene, indi-
cating a more recent common ancestry. Similarly, En1 genes and En2 genes are more closely
related to each other than to any of the *engrailed* homologs found in more basal vertebrate taxa. Fish *engrailed* genes have duplicated yet again to create four paralogs—*Eng1a*, *Eng1b*, *Eng2*, and *Eng3*.

**Mapping gene duplication events onto an animal phylogeny**

The relative timing of gene duplication events can be mapped onto an animal phylogenetic tree to provide a complete history of the evolution of a gene family. The presence of the same pair of duplicated genes in multiple animals indicates that the gene duplication event preceded the separation of their respective lineages; thus, this event must have predated their last common ancestor.

Continuing with the analysis of vertebrate *engrailed* genes, it is clear that the *engrailed* gene was duplicated early in vertebrate evolution, as humans, mice, and chicks have two paralogous *engrailed* genes (*En1* and *En2*). The *engrailed* gene phylogeny indicates that one
duplication occurred early in the vertebrate lineage after the divergence of lamprey (Fig. 4.5). A second duplication within the fish lineage generated the four *engrailed* genes found in zebrafish.

One critical step in establishing the relative timing of duplication events in a gene family is the careful choice of one or more outgroups (for example, animal lineages that diverged before a gene duplication) to help determine the ancestral condition for a group of organisms. In the case of *engrailed*, the identification of a single gene in lamprey and in several other basal deuterostome taxa is consistent with the ancestral deuterostome lineage having had a single *engrailed* gene.

In practice, decoding the pattern and timing of gene duplication is often more difficult than suggested by this scenario. The resolution of the branching pattern of gene phylogenies is not always definitive and can be consistent with more than one historical path. It can prove difficult to determine with certainty whether members of a gene family arose from a duplication in a common ancestor or from more recent, independent duplications in different animal lineages. The possibility of gene loss also can obscure the timing of a gene duplication event. Without a complete reconstruction of ancestral states, it is difficult to interpret whether
the absence of a gene in an extant genome reflects gene loss or whether it indicates that the
gene never existed in that animal lineage. For example, the first *engrailed* duplication in the
vertebrate lineage could have occurred before lamprey diverged, with one copy subse-
quently being lost in the lamprey lineage. If this scenario is correct, the real timing of the
*engrailed* duplication event probably lies deeper within the tree, at the time of large-scale
genomic duplications at the base of the vertebrate lineage.

**Gene divergence**

A gene duplication event may initially generate two redundant gene copies if both the
coding sequences and cis-regulatory control regions are duplicated. Redundancy can allow
one gene copy to be rapidly lost; indeed, gene loss may be a common result following gene
duplication. Physical gene loss can occur because of chromosomal deletions, or a duplicated
gene can be functionally lost because of the accumulation of deleterious mutations (becom-
ing a *pseudogene*). Large-scale genomic duplications, including tetraploidization, generate
large numbers of redundant genes that are not always maintained, as reflected in the loss
of some genes from vertebrate *Hox* clusters (Fig. 4.3). Nonetheless, large gene families of
paralogous genes have evolved due to the evolutionary fixation of duplication events. Why,
then, are duplicated genes retained within a genome?

Many duplicated genes persist because of functional divergence between the two paralogs.
Paralogs can diverge through changes in their coding sequences that lead to differences in
protein function, and they may accumulate changes in their cis-regulatory elements that gen-
erate differences in the timing or pattern of gene expression during development (Fig. 4.6).
Duplicated genes do not have to evolve new functions to be retained, if ancestral functions
are partitioned, or subfunctionalized, between the duplicate copies. Indeed, the modular

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### Table: Evolution of *engrailed* genes in deuterostomes

<table>
<thead>
<tr>
<th>Species</th>
<th>Number of <em>engrailed</em> genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Human</td>
<td>2</td>
</tr>
<tr>
<td>Mouse</td>
<td>2</td>
</tr>
<tr>
<td>Chicken</td>
<td>2</td>
</tr>
<tr>
<td>Zebrafish</td>
<td>4</td>
</tr>
<tr>
<td>Lamprey</td>
<td>1</td>
</tr>
<tr>
<td>Amphioxus</td>
<td>1</td>
</tr>
<tr>
<td>Sea Urchin</td>
<td>1</td>
</tr>
</tbody>
</table>

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**Figure 4.5**

The *engrailed* gene duplication events discussed in Figure 4.4 are mapped onto a deuterostome
phylogeny. The duplication event that created the *En1* and *En2* genes of higher vertebrates appears
below the divergence of tetrapods and fish, because both of these genes are present in
higher vertebrates, but above the lamprey
lineage. Other duplication events in the fish
lineage created the gene pairs *Eng1α/Eng1β* and
*Eng2/Eng3*, perhaps with a single genome-wide
tetraploidization.
nature of cis-regulatory elements may facilitate the partitioning of ancestral expression domains. For example, one gene copy may lose an element that is retained in the duplicate, whereas the other gene copy loses a different element. In addition, new elements may be individually gained to produce novel expression patterns.

One example of duplicated genes that have diverged primarily in their regulation involves the *Drosophila gooseberry/gooseberry-neuro* gene pair. These linked genes encode functionally redundant Pax family transcription factors, yet are expressed in different tissues in early development. The *gooseberry* gene, which is one of the *Drosophila* segment polarity genes, is expressed in stripes in the early embryo. By contrast, *gooseberry-neuro* is expressed at later stages, in the developing nervous system. The expression patterns of these related genes are controlled by different cis-regulatory elements. Changes in cis-regulation represent the primary evolutionary difference between *gooseberry* and *gooseberry-neuro*.

Of course, genes that have not experienced a recent duplication event are also susceptible to change due to mutation. Thus, orthologs in different lineages (and even within a population) also undergo diversification. Changes between orthologs are most evident in
sequence differences, though the vast majority of these differences do not apparently alter protein function. The constraint of maintaining a gene’s function most often results in stabilizing selection. Divergence in protein function of orthologs is possible, however, in cases of positive selection for a new function, or when interacting proteins coevolve in different lineages.

**Assembly of the toolkit: the first animals**

The growing list of genes shared by mice and flies (see Chapter 2 and Tables 2.1 & 2.2) and other bilaterians reveals that their common ancestor had an extensive toolkit of developmental genes. Basal animal lineages, including the diploblast phyla (Cnidaria, Ctenophora) and the Porifera (sponges), have much less developmental and morphological complexity than do bilaterians (Fig. 4.1). Does their simpler body organization reflect a smaller complement of toolkit genes? Or did the bilaterian toolkit predate the origin and radiation of animals all together?

We can track the assembly of the toolkit for animal development by comparing the genes shared among bilaterian phyla to the genes of cnidarians, sponges, and even other eukaryotic organisms. The genome projects for the yeast *S. cerevisiae* and the flowering plant *Ara- bidopsis*, in addition to the *C. elegans*, *Drosophila*, and various deuterostome species, have generated data that facilitates the direct comparison of fungal, plant, and animal genomes. Many protein domains that are characteristic of animal transcription factors or signaling molecules are found in yeast and plants, and thus have ancient origins predating multicellular life (Table 4.1). For example, homeodomains are present in all multicellular organisms, indicating that this protein motif is older than animals themselves. Other domains found in proteins that are crucial to animal development, however, have not been identified outside the animal kingdom. For example, TGF-β and Wnt signaling molecules, in addition to several families of transcription factor families, have been found only in animals (Table 4.1). Thus some toolkit genes evolved from ancient genes, whereas others appeared early in the animal lineage.

The set of genes present in the first animals may also be inferred by the complement of genes found in choanoflagellates, single-celled protozoa thought to be the closest outgroup to the metazoan clade, as well as basal animals (Cnidaria, Porifera). For example, choanoflagellates have genes involved in cell signaling and adhesion in animals, including tyrosine kinase signaling components, indicating that these genes predated the origin of animal multicellularity. Additional toolkit genes appear in the genomes of cnidarians, including components of the Wnt and TGF-β signaling pathways and many transcription factor families (Table 4.1). Cnidarians have several homeodomain-containing genes that are homologous to bilaterian toolkit genes (including *even-skipped*, * engrailed*, *Distal-less*, and *Hox* genes) and at least four members of the *Pax* family of transcription factors, several T-box genes (including *Brachyury*), a *Snail* homolog (zinc finger), a *Twist* homolog (helix-loop-helix), and a *mef2* (MADS-box) homolog. Clearly, at least some members of the bilaterian toolkit were present in the earlier animal lineages.

Although cnidarian genomes contain many of the gene families that contribute to the toolkit, cnidarians appear to have a relatively small number of toolkit genes. For example, bilaterians have at least one additional *Pax* gene and many more *Hox* genes than are found in cnidarians. A comparison of the toolkit genes shared among bilaterians with the genes present in cnidarians indicates that the bilaterian toolkit expanded through gene duplication after the divergence of the cnidarian lineage but before the radiation of bilaterian phyla.
CASE STUDY: EVOLUTION OF THE HOX COMPLEX

Expansion of the HOX complex

The best-characterized history of a family of developmental regulatory genes is the evolution of the HOX genes. HOX genes have been isolated from many metazoan phyla, allowing the timing of the tandem duplication events that created the HOX complex to be determined relative to the radiation of animal phyla (Fig. 4.7). Given these data, the history of the expansion of the HOX complex can be charted through the course of animal evolution.

Cnidarians have two definitive HOX genes, which are most closely related to the “anterior” group (lab/Hox1, pb/Hox2) and “posterior” group (AbdB/Hox9-13) genes found in bilaterians. Although some cnidarians have additional HOX genes due to cnidarian-specific duplication events, the common ancestor of Cnidaria and Bilateria most likely possessed only two HOX genes. By contrast, all bilaterians have at least two anterior group genes, multiple “central” group genes (Hox3 through Hox8, or their equivalents), and often multiple posterior group genes. Five HOX genes (lab/Hox1, pb/Hox2, Hox3, Dfd/Hox4, and Scr/Hox5), as well as at least one more central gene and one posterior gene, can be clearly identified in all three major bilaterian lineages (the Lophotrochozoa, the arthropod + onychophora + priapulid...
Chapter 4: Evolution of the Toolkit

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**Genome tetraploidization events**

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### Expansion of central Hox genes

- **Anterior:** Ubx, Abd-B
- **Central:** Lox5, Lox2/4, Post1/2
- **Posterior:** Lox5, Lox2, Lox4, Post1, Post2

**Figure 4.7**

**Evolution of metazoan Hox genes**

The relative timing of Hox duplication events is mapped onto a phylogenetic tree of metazoan phyla ([left](#)), as deduced from the distribution of Hox genes in metazoan phyla ([right](#)). Cnidarians have Hox genes representing only the most anterior and posterior group genes. Early expansion of the Hox complex at the base of the bilaterian lineage generated many of the central Hox genes. Before the bilaterian radiation, the Hox paralogy groups Hox1–Hox5 were established and fixed. The deuterostome Hox complex continued to expand to create Hox6–Hox13. During early vertebrate evolution, the entire complex became duplicated, thereby creating four complexes in tetrapods (representing the vertebrates). In the arthropod + onychophora + priapulid lineage, gene duplications and/or sequence divergence and fixation created the signature arthropod Hox genes Ubx and Abd-B. Similarly, the lophotrochozoans are characterized by Lox5, Lox2, Lox4, Post1, and Post2. Additional closely related central group genes are present in protostome phyla. Gray-shaded boxes delineate orthology assignments between phyla, when they can be clearly established.

clade, and the deuterostomes). This finding indicates that the last common bilaterian ancestor possessed at least seven Hox genes. Thus five or more Hox genes arose early in the bilaterian lineage, after the divergence from diploblasts but before the radiation of bilaterian phyla (Fig. 4.7).

It is important to recognize that not only did several new Hox genes arise before the radiation of bilaterian phyla, but also the homeodomain sequences of these Hox genes also diverged. The various Hox paralogy groups can be identified through their characteristic amino acid residues. Because these characteristic residues are conserved in long-diverged bilaterians, they must have been present and fixed in the ancestral Hox genes of the last common bilaterian ancestor. Thus the genes of the ancestral Hox complex included distinct sequences that have changed very little during the subsequent 500 million years or more of animal evolution.

The Hox complex continued to expand during bilaterian evolution. Its evolution in the protostome clade is characterized by sequence divergence of additional central and posterior group Hox genes. The two major protostome clades, the Lophotrochozoa (annelids, molluscs, flatworms, lophophorates) and the group containing the arthropods, onychophora, and priapulids, each have central and posterior Hox genes specific to their clades. The lophotrochozoans are characterized by the presence of Lox5, Lox2, Lox4, Post-1, and Post-2, and the arthropods + onychophora + priapulids by Ubx and Abd-B (Fig. 4.7). These genes may have arisen from independent gene duplication events or from sequence divergence that occurred early in each clade. The respective sets of lophotrochozoan and arthropod/onychophoran/priapulid Hox genes were represented in the last common ancestor of each lineage, before the radiation of protostome phyla took place.

The Hox complex also expanded early in the deuterostome lineage through tandem expansion of central and posterior group genes. There is a single Hox cluster with 10–14 genes in nonvertebrate deuterostomes, including echinoderms, ascidians (urochordates), and Amphioxus (cephalochordate) (Fig. 4.7). The deuterostome Hox6-8 (central group) and Hox9-13 (posterior group) genes do not have clear orthologs in the protostomes, indicating that they arose independently within the deuterostome clade.

Basal deuterostomes have sets of Hox genes that are very similar to those found in the vertebrates, but vertebrates have many more genes (Fig. 4.7). The latter development clearly reflects large-scale duplications or tetraploidization events early in the vertebrate lineage that generated at least four separate copies of the entire Hox complex. The presence of multiple Hox complexes in lampreys and sharks places the timing of the Hox complex duplication events at the base of the vertebrate lineage, after the divergence of cephalochordates (Fig. 4.8). A further round of duplication created additional Hox complexes in ray-finned (teleost) fish including zebrafish and Fugu (Fig. 4.8). This teleost-specific expansion is observed for other genes as well, including the engrailed example discussed earlier in this chapter, consistent with an additional large-scale or genome-wide duplication event within the teleost clade. The additional Hox genes in teleosts became subfunctionalized through the partitioning of expression domains and developmental functions.

The ParaHox genes: a sister complex to the Hox genes

The origin of the Hox complex is primarily a story of tandem gene duplication and divergence. Nevertheless, additional evidence indicates that the entire prebilaterian Hox complex of three
to four genes was duplicated early in metazoan history. The duplication of this “Proto-Hox” complex gave rise to the ancestral Hox complex and a sister “ParaHox” complex. The Para-Hox genes (Gsx, Xlox, and Cdx) have homeodomains with high sequence similarity to the anterior group, Hox3, and posterior group Hox homeodomains, respectively. The tight chromosomal linkage between these genes in Amphioxus (Fig. 4.9) and their colinear deployment in regions of the developing gut are reminiscent of the organization and colinearity of the Hox complex. The presence of both ParaHox and Hox genes in bilaterians and cnidarians suggests that the Proto-Hox complex duplication predates the divergence of cnidarians and the bilaterian lineage.

The relative timing of vertebrate Hox complex duplications is indicated on a phylogenetic tree of deuterostomes. Basal deuterostomes, including echinoderms and cephalochordates, have a single Hox complex. At least three complexes have been identified in the primitive jawless lamprey, indicating that Hox complex duplication occurred before the divergence of lamprey from higher vertebrates. Teleost fish have undergone an additional round of tetraploidization, creating additional Hox complexes in zebrafish and Fugu. Tetrapods, including humans, mice, and frogs, have four Hox complexes and a total of 39 Hox genes. Sharks possibly possess more than two complexes.

New functions for some insect Hox genes

While the Hox homeodomain sequences have generally been constrained against change, several Hox genes have evolved new developmental functions in the Drosophila lineage (Fig. 4.10) and no longer regulate regional identity along the A/P axis. The evolution of novel developmental roles correlates with rapid divergence in the sequence of the homeodomain, which may reflect the release from functional constraint as a Hox gene. In the first case, the ancestral Hox3 gene lost its Hox function during insect evolution and assumed a novel role in determining extra-embryonic tissue to become the zerknült (zen) gene in beetles, grasshoppers, and flies. More recently in the Drosophila lineage, the zen gene duplicated and diverged to create bicoid (bcd) as well, which in Dipteran flies acts maternally in A/P axis determination. A substitution at a key protein residue in the bcd homeodomain at position 50 has shifted the binding specificity of bcd, associated with the replacement of orthodenticle function with bcd in A/P axis determination in Diptera.

In the second case, the Drosophila fushi tarazu (ftz) pair-rule gene evolved from a central group Hox gene that is present in other arthropods. Functional tests of ftz genes from different
insects in *Drosophila* indicate that *ftz* has slowly lost its *Hox* function (retained somewhat in beetles and grasshoppers) and gained a pair-rule function. The expression of *ftz* in the central nervous system of several insects and a cirripede crustacean is more reminiscent of *Hox* genes, with a clear anterior boundary of expression in the thorax. The probable ancestral *Hox* expression pattern is reflected in the deployment of the *ftz* ortholog in a myriapod (centipede) and a chelicerates (mite), where this gene is expressed in a *Hox*-like pattern in the embryonic ectoderm.
Hox proteins can also acquire new functional motifs while retaining their Hox function. In the insect lineage the Ubx protein has gained a functional motif that is exploited as part of its role in regulating regional identity along the A/P axis. Insect Ubx proteins contain a carboxy-terminal glutamine and alanine-rich transcription repression domain that is not present in crustacean or onychophoran Ubx proteins, and which is implicated in the evolution of morphological segmental diversity in insects. We will go into more detail on the role of Hox protein evolution in arthropod body plan diversification in Chapter 5.

The evolution of the Hox complex illustrates several important mechanisms underlying the evolution and functional diversification of gene families. First, an early duplication of the small, ancestral Hox complex created the sister ParaHox complex. Second, tandem duplication events expanded the number of paralogous Hox genes within the complex. Third, the homeodomains of many Hox genes diverged and became constrained before the bilaterian radiation. Fourth, large-scale or genomic duplications at the base of the vertebrate lineage (and again in teleosts) increased the number of vertebrate Hox complexes. Fifth, rapid changes in homeodomain sequence accompanied the evolution of new developmental functions for some insect Hox genes. Sixth, evolution of new motifs has modified Hox protein functions.

**INTERPRETING THE TOOLKIT: INFERENCES ABOUT ANIMAL EVOLUTION**

**Expansion of the toolkit and the evolution of morphological complexity**

Is expansion of the genetic toolkit for development related to morphological evolution? Two significant periods of expansion of toolkit genes which occurred in the stem lineage of all bilaterians and in the stem lineage of vertebrates, do, in fact, correlate with periods of remarkable increases in developmental and morphological complexity. During the same interval in which the number of Hox genes expanded from two to seven distinct genes, the bilaterian stem lineage transitioned from a diploblast body organization to a bilaterally symmetrical triploblast animal. Several dramatic and crucial developmental innovations evolved during this period:

- The establishment of a regulated pattern of early cleavage to form an organized multicellular embryo
- The formation of a continuous gut
- The appearance of a new mesodermal tissue layer with an associated cavity (**coelom**)  
- The evolution of distinct anterior–posterior and dorsal–ventral axes

These features are the product of new or modified developmental genetic programs that may well have involved “new” developmental genes.

Large-scale genome duplications at the base of the vertebrate lineage greatly increased the number of Hox genes, as well as the number of other toolkit genes. Although these duplications created a great deal of genetic redundancy, many duplicated genes have been retained over a few hundred million years of vertebrate evolution and along the way acquired new or different roles in development. For example, in the mouse, three homologs of the *Drosophila hedgehog* gene (**Sonic hedgehog**, **Indian hedgehog**, and **Desert hedgehog**), are expressed in different tissues and play different roles during development.
During the same evolutionary period, the developmental and anatomical complexity of early vertebrates increased, as reflected by the greater number of differentiated cell types in vertebrates compared to other bilaterians. In particular, the vertebrate central nervous system became larger and more elaborate, which may have contributed to the ecological dominance exhibited by vertebrates since the Devonian (circa 300 Ma). The developmental complexity and success of the vertebrate lineage may reflect the exploitation of the dramatically larger number of developmental genes in the vertebrate toolkit (Table 4.1).

Although the expansion of the toolkit may be related to morphological complexity, no correlation appears to exist between toolkit expansion and animal diversity. The morphological diversity of vertebrates, from humans to hummingbirds, or from whales to snakes, evolved around a common set of developmental genes. For example, mammals, birds, and amphibians share the same set of 39 Hox genes. The story is similar for the invertebrate bilaterian phyla, all of which appear to have roughly comparable sets of Hox genes. In fact, the same Hox genes are found in all arthropods and in the most closely related phylum to the arthropods, the onychophora. The remarkable diversity of fossil and extant onychophorans, trilobites, myriapods, crustaceans, arachnids, and insects evolved around a shared complement of Hox genes. The apparent simplicity of some bilaterian body plans, such as the onychophora or the less glamorous flatworms and priapulids, belies the extensive genetic toolkit that these phyla share with more elaborate animal forms.

One interesting exception to the conservation of the toolkit for development is the apparent loss of some toolkit genes in the nematode *C. elegans*. While the uncertain phylogeny of nematodes within bilateria complicates the interpretation of the *C. elegans* toolkit, some genes that predate the common ancestor of the three major bilaterian lineages (deuterostomes, arthropods, and lophotrochozoans) are not found in *C. elegans*. For example, *C. elegans* has fewer Hox genes than other bilaterians. Furthermore, *C. elegans* lacks a clear ortholog of a hedgehog gene (although this apparent absence may reflect significant divergence of hh-like genes in the nematodes, as other members of the hedgehog signaling pathway are present). The origin of the hedgehog gene predates the bilaterian clade, so the absence (or divergence) of these genes represents a derived condition of *C. elegans*. Many nematodes are parasitic, which may create selective pressures that simplify development and morphology and facilitate the loss (or divergence) of critical developmental genes. Such dramatic gene loss appears to be the exception rather than the rule among bilaterians.

**Conserved genes, conserved biochemical functions**

The conservation of the genetic toolkit for developmental genes extends beyond primary protein sequences. Even though these genes are used in animals with radically different modes of development, *in vivo* comparisons of protein activity have revealed similarities in biochemical function between orthologous proteins. In particular, studies have demonstrated the ability of genes from evolutionarily distant species to recapitulate the activity of orthologous genes during development.

For example, overexpression of vertebrate Hoxb1 and Hoxb4 genes in *Drosophila* generates phenotypes that resemble the effects of overexpression of the *Drosophila* labial and Dfd genes, respectively. The sequence similarity between vertebrate Hox proteins and their arthropod orthologs is limited to the homeodomain and two other short peptides, and these sequences appear to be sufficient for most Hox protein functions (with some exceptions, as
noted in the previous section). As a general rule for Hox genes and many other toolkit genes, protein sequence conservation often reflects conservation of biochemical functions.

The functional conservation of transcription factors such as the Hox proteins probably derives from the constraints imposed by their regulation of potentially very large numbers of target genes. It may be very difficult for the sequence of a homeodomain to change, as a modification may affect its DNA-binding specificity and therefore alter the ability of the protein to properly regulate all of its target genes. A modification to a Hox protein that simultaneously disrupts multiple Hox-regulated networks would be catastrophic to development. Similarly, signaling pathways are used in a variety of tissues and at different stages of development; hence, a change in the ability of a ligand to interact with its receptor or a change in the interactions between other pathway components could have wide-ranging, dire consequences. For this reason, the evolution of toolkit proteins is constrained in order to maintain long-established ancestral biochemical functions.

Similarities in the biochemical functions of orthologs do not necessarily indicate that these genes are used for the same developmental function in their respective organisms, however. This point is most readily apparent for signaling proteins. Conserved biochemical functionality for a signaling protein simply requires conservation of its ability to bind to a receptor and trigger an intracellular signaling cascade. For example, in vertebrates and in fruit flies, the Hedgehog protein binds to the Patched receptor protein, thereby activating the Cubitus interruptus/Gli transcription factor (see Chapter 2 and Table 2.2). Nevertheless, the developmental outcome is different in each organism. In the vertebrate neural tube, a gradient of Hedgehog signaling generates different neuronal identities; in *Drosophila*, this pathway is used during segmentation and later in imaginal discs to pattern a variety of insect structures. Similarly, orthologous transcription factors may share the same DNA binding specificity and interact with the same cofactors, but perform different regulatory roles during development.

**Conserved developmental functions: rebuilding the bilaterian ancestor from phylogenetic inference**

Although the conservation of biochemical functionality may simply reflect evolutionary constraint on protein sequence evolution, similarities in the developmental function of homologous genes raise the possibility of deeper evolutionary significance. For example, how should we interpret the striking similarity in the developmental function of Hox genes in regulating regional identities along the A/P axis of both mice and *Drosophila*? One possibility is that this similarity is mere coincidence—that the transcription factors independently evolved roles in patterning the A/P axis of protostomes and deuterostomes. More interestingly, the Hox genes may have played a role in patterning the A/P axis of the last common ancestor of protostomes and deuterostomes—the hypothetical ancestor of all bilaterians, dubbed “Urbilateria.”

One way to distinguish between the possibility that observed similarities are convergent and the likelihood that they reflect conservation of features in a common ancestor is to examine other phyla. The more taxa that share a characteristic, the less likely that the similarity is due to convergence. The deployment of Hox genes along the A/P axis of *Amphioxus*, annelids, flatworms, and other bilaterians, for example, suggests that Hox colinearity is an ancestral feature of bilaterians.

Not only may we infer the ancestral function of toolkit genes from shared developmental functions among living bilaterians, but we may also infer some of the morphological
characteristics of Urbilateria. No fossils have been identified that provide direct knowledge about the morphological complexity of early bilaterian ancestors. Instead, we must use comparative developmental genetic data from living animals in our attempt to rebuild a picture of Urbilateria. The identification of conserved developmental regulatory genes, expressed in similar patterns or cell types among multiple phyla, implies that the same genes were utilized in a common ancestor to control pattern formation and cell differentiation; the open question concerns the complexity of the resulting structures in animal ancestors.

The logic that underlies inferences about the development and morphology of Urbilateria is illustrated by the conserved role of *Pax6* genes in eye development. Both the mouse *Pax6* gene and the *Drosophila* ortholog *eyeless* are at the top of the regulatory hierarchies that direct eye development in each organism. Other components of the *Pax6*-regulated circuit, including the *sine oculis*, *eyes absent*, and *opsin* genes, are also shared between flies and mice. Furthermore, *Pax6* orthologs are expressed during eye development in many other bilaterian phyla and a related *Pax* gene (with a *Pax6*-like homeodomain) is expressed in the eyes of a cnidarian (Fig. 4.11), providing more evidence that the functional similarities

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**Figure 4.11**

*Pax6* expression during eye development in different bilaterian phyla

Pax6 protein expression has been characterized in many bilaterian phyla. This protein is expressed in the eyes of vertebrates, urochordates, arthropods, annelids, molluscs, flatworms, and nemerteans (not shown). Thus the common bilaterian ancestor may have deployed an ancestral Pax6 gene in photoreceptor cells during the development of a primitive "eye" or light-sensing organ.
between the mouse Pax6 and Drosophila eyeless genes in eye development may be inherited from a common ancestor. The remarkable conservation of Pax6 expression and the Pax6-regulated circuit suggests that all bilaterian eyes share a common developmental genetic circuit and that this circuit was present in the bilaterian ancestor. Although we cannot say whether Urbilateria had eyes per se, it is likely to have possessed light sensing cells whose differentiation depended upon Pax6 function.

Several other developmental features of Urbilateria may be inferred from the conservation of developmental patterning roles of toolkit genes (Fig. 4.12). The similarity between D/V axis polarization by the short gastrulation/chordin genes and TGF-β signaling in insects and vertebrates, and A/P axis regionalization by Hox genes in many bilaterians, suggests that these genes also controled the patterning of the primary body axes of Urbilateria. A suite of common mesoderm patterning genes (including Brachyury, Twist, mef2, and Snail) is shared among insects and vertebrates. In fact, all of these genes and patterning systems have been identified in cnidarians as well, and their roles as developmental regulatory genes clearly predated Urbilateria. Cnidarians have a very different body organization, but do deploy the TGF-β and Wnt signaling cascades during formation of the primary body axis, Hox and ParaHox genes in restricted axial domains, and several anterior patterning genes (homologs of Brachyury, emx, forkhead, and paired-like genes) during head formation. Several bilaterian mesoderm patterning genes (Brachyury, Twist, mef2, and Snail) are expressed in striated muscles in the medusa stage of a hydrozoan, suggesting that bilaterian mesoderm may have an ancestral relationship to cnidarian striated muscle.

Using similar logic, additional patterning mechanisms present in Urbilateria can be inferred from other shared regulatory roles for toolkit genes in deuterostomes and protostomes. Anteroposterior patterning of the Urbilaterian endoderm, ectoderm, and nervous system may have been regulated by the ParaHox genes (Gsx, Xlox, Cdx), certain segmentation genes (barly, engrailed, Notch), and brain patterning genes (emx, otx, Hox), respectively. Even the expression patterns of the genes involved in dorsal–ventral patterning of the central nerve cord of flies and mice are similar (vnd/NK-2, ind, msh, netrins). Perhaps most surprisingly, similarities in the genetic regulatory mechanisms that control the development of cardiac tissue (tinman/Nkx2.5) and appendages (Dll/Dlx; bth/Meis + Exd/Pbx) in insects and vertebrates suggest that Urbilateria could have possessed a primitive contractile organ and structures that projected from the body.

Without fossils of early animal ancestors, we can only speculate about the complexity of the primitive bilaterian body plan. Perhaps the “organs” of Urbilateria were simple structures composed of a few specific cell types. For example, Urbilateria may have had a simple photoreceptor complex rather than an optically sophisticated eye, a contractile muscle regulating hemocoel fluids rather than a modern heart, and a simple outgrowth of the body wall or tentacle-like feeding structure rather than a modern locomotory appendage (Fig. 4.12). Although the anatomical details remain uncertain, the development of such structures could sufficiently constrain gene evolution to conserve the function of regulatory genes across Bilateria.

All of these features combine to build a fairly complex image of the “primitive” bilaterian, representing an animal that could move (or perhaps crawl) through sediment and could sense and interact with the environment. Perhaps most importantly, this type of ancestor would have many morphological features and genetic tools that might have facilitated a successful evolutionary response to changes in the natural history of early animal life, including
Figure 4.12: Rebuilding “Urbilateria”

The possible features of the common bilaterian ancestor are deduced from the conservation of genes and their developmental functions between arthropods (Drosophila) and vertebrates (mouse). (Top) Patterning of the D/V axis may have been controlled by ancestral genes of the short gastrulation (sog)/chordin and TGF-β families. The A/P axis was probably subdivided by nested, overlapping domains of Hox gene expression. (Middle) Different tissue layers were regionally patterned along the A/P axis, including the gut (ParaHox genes) and the nervous system (otd, ems, Hox). Segmentation may have evolved under the regulation of ancestral hairy, engrailed, and Notch genes. (Bottom) Primitive versions of a photoreceptor organ, a circulatory pump, and a body wall outgrowth might have been present in the bilaterian ancestor, under the regulatory control of the ancestral Pax6, tinman, and Dll genes, respectively.
climate change and predation. The inferred presence of ancestral patterning mechanisms and differentiated cell types, even in a simple or primitive form, demonstrates that the potential for much greater morphological complexity and diversity was largely in place in Urbilateria and was subsequently released by environmental and ecological forces (Fig. 4.13).

The revelation of developmental regulatory similarities between long-diverged bilaterians has complicated the assessment of the evolutionary relationships between all of the structures that deploy a shared regulatory gene. For example, it was once believed that “eyes” had evolved independently in arthropods, molluscs, and chordates, based upon morphological and phylogenetic considerations. The discovery of the role of Pax6 (and other genes in the circuit) in the development of all sorts of eyes suggests that eyes did not, in developmental genetic terms, evolve repeatedly “from scratch.” Regulatory circuits made up of the same sets of genes are not likely to have been constructed independently by chance, gene by gene, out of the entire repertoire of hundreds of transcription factors in the toolkit.

Figure 4.13
Evolution of animal morphological complexity
The evolution of diverse animal forms followed the radiation of bilaterian phyla in the Cambrian. The toolkit for animal development expanded in the stem lineage of bilaterians and was largely present in Urbilateria. The ecological forces of the Cambrian may have facilitated the evolution of animal morphology using an established set of developmental regulatory genes and networks to build diverse body parts and body plans.
Figure 4.14

Expression of Dll in bilaterian phyla

Dll protein is expressed in body wall outgrowths of many bilaterian phyla, including representatives from all three major bilaterian clades (arthropods + onychophora, lophotrochozoans, and deuterostomes). (a) Dll expression (green) in an onychophoran embryo marks the head appendages and the lobopod walking legs (inset). (b) In a polychaete (annelid) embryo, Dll is expressed in the growing parapodia (inset) and other appendages. (c) Dll is expressed in cells at the distal tip of tube feet and spines of a newly metamorphosed sea urchin (echinoderm). Dll expression marks the distal tip of proximo-distal outgrowths in these phyla, suggesting that a P/D axis was defined in Urbilateria and was marked by Dll.

In another example, the deployment of Dll during the development of body wall outgrowths in many bilaterian phyla (Fig. 4.14), as well as the function of \textit{hth}/\textit{Meis} and \textit{Exd}/\textit{pbx} genes in proximodistal limb patterning in arthropods and vertebrates, is not likely to have evolved de novo in each lineage. This similarity, however, does not mean that all bilaterian limbs are homologous, indeed, they are not. All bilaterian limbs did not evolve directly from an Urbilatariian appendage. Rather, they may be considered to be developmental “paralogs” of one another, products of the novel deployment and modification of an ancient and shared regulatory circuit in many different animal lineages over the course of animal evolution.

THE TOOLKIT AS DEVELOPMENTAL POTENTIAL

The deduced complexity of the genome of bilaterian ancestors is much greater than was once thought. Likewise, the image of Urbilateria as an animal with regionally differentiated body axes and a variety of specialized organs is more elaborate than earlier inferences. These combined genetic and developmental features push the origin of animal body patterning deeper into metazoan history. Indeed, the origin of bilaterian complexity—including the evolution of mesoderm, a patterned central nervous system, a regionalized through gut, and primitive eyes, heart, and limbs—corresponds to a period of early animal evolution that was marked by an increase in the number of developmental toolkit genes. The period between the last common ancestor of diploblasts and bilaterians and the radiation of bilaterian phyla represents an interval during which considerable genetic and morphological complexity evolved.

By contrast, the radiation of diverse bilaterian body plans is cast against a backdrop of shared genetic tools. Certainly, the genetic toolkit continues to evolve—gene duplication and divergence is a never-ending process. Notably, large-scale duplications early in the vertebrate lineage multiplied the contents of the vertebrate toolkit. Yet the basic components are widely shared among invertebrate bilaterian phyla, and the biochemical functions of the encoded proteins are surprisingly conserved across hundreds of millions of years. What, then, is the genetic basis underlying the morphological diversification of animal forms?

The remaining chapters of this book look not to the evolution of new genes, but rather to higher-order changes in developmental genetic programs. In particular, Chapter 5 focuses on evolutionary changes in gene regulation, including the pattern, timing, and level of gene expression and the genetic wiring of regulatory interactions, which enabled the diversification of arthropod and vertebrate body plans.

SELECTED READINGS

\textit{General Reviews}


Conservation of Developmental Regulatory Genes


Gene Duplication and Divergence: Assembly and Expansion of the Toolkit


Case Study: Evolution of the Hox Complex


Conserved Genes, Conserved Functions: Rebuilding the Bilateral Ancestor


We suggest that evolutionary changes in anatomy and way of life are more often based on changes in the mechanisms controlling the expression of genes than on sequence changes in proteins.

Mary-Claire King and Allan Wilson (1975)

CHAPTER 5

Diversification of Body Plans and Body Parts

Chapters 2 and 3 described the general principles of the underlying unity in developmental regulatory mechanisms, and Chapter 4 detailed the widespread conservation of the genetic toolkit for development among bilaterians. These chapters set the stage for the consideration of longstanding questions about body plan evolution. Given the extensive genetic similarities of living animals, how did new and vastly different forms evolve from a common bilaterian ancestor? What are the genetic differences that underlie the diversity of animal body patterns? This chapter focuses on ways in which evolutionary changes in the regulation of toolkit genes during development contributed to morphological change. Here we examine the relationship between body plan evolution and regulatory evolution, concentrating primarily on the diversification of repeated structures along the primary body axis and of homologous parts between lineages.

The identification of genetic mechanisms underlying body pattern diversity relies on a comparative approach encompassing both model organisms and their relatives. Our understanding of the genetic basis of the radiation of body plans within a phylum is mostly limited to arthropods (including the insect Drosophila melanogaster) and chordates (including vertebrate model systems such as chicks, mice, frogs, and zebrafish). Fortunately, these phyla are also exemplary regarding the degree of body plan diversification they display within the framework of a shared body organization. The incredible diversity of extant arthropods, particularly of crustaceans and insects, in combination with the many arthropod and onychophoran forms present in the Cambrian period (530 Ma), provides a rich experimental and historical foundation for a case study of body plan evolution. Similarly, the evolution and subsequent diversification of the axial morphologies of modern chordates provide dramatic examples of large-scale morphological diversification.
Much of our understanding of the role played by regulatory evolution in shaping animal body patterns comes from the \textit{Hox} genes. The \textit{Hox} genes are the best-characterized developmental regulatory genes within and between Metazoan phyla. The genes themselves predate the radiation of bilaterian body plans (see Chapter 4). In this chapter, we examine how \textit{Hox} genes are used during development in related organisms to understand how evolutionary changes in these and other selector genes contribute to body plan evolution. Comparative analyses of \textit{Hox} gene expression domains reveal that major transitions in body organization in both arthropods and tetrapods correlate with shifting spatial boundaries of \textit{Hox} gene expression. In particular, differences in the regulation of \textit{Hox} genes correlate with the diversity of the number and identity of repeated units, such as arthropod and annelid segments and appendages and vertebrate axial elements.

More closely related animal groups with a more conserved body organization, such as insects, exhibit fewer large-scale differences in \textit{Hox} gene expression. The diversification of homologous structures in the context of a stable body plan is largely characterized by regulatory changes downstream of \textit{Hox} (or other selector) gene function. For example, the morphological diversity of insect hindwings and of vertebrate forelimbs is a consequence of evolutionary changes in the assortment of target genes regulated by the \textit{Hox} (or other selector) genes that pattern these appendages. A second mechanism of diversification acting in organisms that share a particular body plan is the modification of \textit{Hox} expression patterns within developmental fields. In this chapter, we discuss the best-understood case studies that illustrate the relationship between morphological diversity and evolutionary changes in the regulation of the \textit{Hox} genes and of their downstream targets.

**DIVERSITY OF ANTERIOR/POSTERIOR BODY ORGANIZATION WITHIN ARTHROPODS AND VERTEBRATES**

The arthropods are the most successful animal taxa, with the insects alone accounting for roughly 75% of all known animal species. All arthropods share a body plan made up of repeated units—the segments that bear the paired, jointed appendages for which the phylum is named. Different arthropod classes have particular body plans characterized by the subdivision of the body into distinct regions (for example, head, thorax, abdomen) containing specific numbers of segments and by the distribution of appendages on those segments. Thus the dominant theme in the evolution of the various arthropod body plans has been the diversification of segment and appendage number, organization, and morphology (see Chapter 1, Fig. 1.6).

Extant arthropods include the following groups:

- The myriapods, which have many similar (homonomous) trunk segments
- The crustaceans, which display dramatic diversity in segment number and appendage shape and function
- The insects, which possess a stereotypical body organization consisting of a complex head, a thorax bearing six walking legs and two pairs of wings, and a limbless adult abdomen
- The chelicerates, which have a unique, two-part body subdivision into prosoma and opisthosoma
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By contrast, the sister phylum to the arthropods, the onychophorans, consists of animals that are much less diverse and have only a small number of segment and appendage morphologies. These characteristics likely reflect the primitive condition of the arthropod/onychophoran clade.

The arthropod fossil record is rich with diverse arthropod forms, and distinct arthropod and onychophoran body plans are readily apparent in the Cambrian period. The early divergence of this clade remains cryptic, as the segmental morphology and complexity of the earliest (Precambrian) arthropod ancestors have not been documented.

The evolution of different vertebrate body plans is also characterized by the divergence of repeated units—namely, the vertebrae and paired appendages (see Chapter 1, Figs 1.6 & 1.7). The mesodermal segments of vertebrates (that is, the somites) originate as identical, serially homologous fields that will eventually give rise to the vertebrae and associated processes of the axial skeleton (see Chapter 3). The number and morphology of the vertebrae that constitute each region of the tetrapod vertebral column (for example, cervical, thoracic, lumbar, sacral) differ among the major tetrapod groups. For instance, mammals typically have seven cervical (neck) vertebrae, as opposed to 13 or more in birds. Further, snakes can have hundreds of thoracic (rib-bearing) vertebrae. These differences in the number of specific vertebrae are significant in terms of the evolutionary adaptation of the various groups of tetrapods. The elongation of the thorax of snakes and some lizards, coupled with the loss of limbs, allowed them to exploit unique ecological niches. Likewise, the variations in the number of cervical vertebrae in birds and the loss of tail vertebrae in some primates are tied to their unique evolutionary histories.

Vertebrate limbs also exhibit a wide range of patterns and functions. The serially homologous forelimbs and hindlimbs of a single animal can have dramatically different morphologies, such as the wings and legs of birds. The shapes and functions of homologous limbs of different animals range from seal flippers to bat wings to human arms.

The evolution of different body plans within a phylum includes both the morphological diversification of repeated body parts within a single animal and the morphological diversification of homologous structures between animal lineages. In genetic terms, serially homologous body parts evolve in the context of the genome of a single species. By contrast, homologous structures that diverge among lineages, such as the forelimbs of different tetrapods or the hindwings of diverse insects, evolve in the context of independently evolving genomes.

**Evolution of the genetic control of segmentation in arthropods**

Underlying the general similarity of arthropod segmentation are conserved gene expression patterns, which suggests that there may be a common regulatory program for segmentation among arthropods. For example, segmentation in all classes—insects, crustaceans, myriapods, and chelicerates—passes through a stage when the *engrailed* gene is expressed in a stripe in each segment (Fig. 5.1). In *Drosophila*, this segment polarity gene—a member of the last tier in the segmentation gene cascade—functions to establish and maintain compartmental and segmental boundaries. In addition, a number of other genes involved in *Drosophila* segmentation, including *even-skipped*, *bairy*, *wingless*, and *runt*, are expressed in patterns that appear to be associated with segmentation in several insects, crustaceans, centipedes, and spiders. Thus, some of the members in the *Drosophila* segmentation cascade appear to be deployed among most arthropods during the segmentation process. However, a number of
striking differences are also apparent. These may be related to the differences in early embryonic development between arthropods. In *Drosophila*, segmentation depends on gradients of transcription factor activity in a syncytial embryo; in other arthropods, segmentation occurs over a longer period in a cellularized embryo.

Differences have evolved in earlier stages of segmentation in several distinct ways. First, genes have been independently recruited for a new role in, or potentially lost their role in, segmentation. Second, the regulatory interactions between members of the segmentation cascade have changed. Third, the evolution of new genes and gene functions has changed the
architecture of the segmentation regulatory hierarchy. We will examine selected examples of each type of evolutionary change in arthropod segmentation regulation.

Some of the mechanisms that regulate *Drosophila* segmentation appear to be more the exception rather than the rule for arthropods or even insects. For example, in spiders, *Notch* pathway genes are expressed in the developing segments and are required for segmentation. However, *Notch* signaling is not involved in segment formation in *Drosophila*. Also, although *even-skipped* is expressed in a segmental pattern in all arthropods, the pair-rule function of the *even-skipped* gene appears to exist only in *Drosophila* and a few other insect lineages. Other segmental gene roles appear to be unique to *Drosophila*, or at least higher dipterans. For example, the *Drosophila* pair-rule gene *ftz* does not have a pair-rule function in other insect orders and arthropod classes, in fact, *ftz* likely has an ancestral role as a *Hox* gene (see Chapter 4). Thus, some members of the segmentation regulatory hierarchy have changed between different arthropods, in part through the recruitment of genes for new roles.

The evolution of regulatory interactions in segmentation is illustrated by the different means by which the *hunchback* gene is regulated in beetles and flies. In the beetle *Tribolium*, the *hunchback* gene has an expression domain similar to the gap expression domain of *Drosophila hunchback*, but the beetle ortholog is regulated differently. The *Drosophila hunchback* cis-regulatory element that directs expression in the gap domain is activated by the Bicoid protein, one of the pivotal regulators of anteroposterior patterning during *Drosophila* embryogenesis. The equivalent *Tribolium hunchback* cis-regulatory element is regulated by the Caudal protein. Thus, similar *hunchback* gap expression domains are governed by cis-regulatory elements that bind different upstream regulators in the beetle and the fruit fly. Homologs of *bicoid* have not been identified outside of higher Diptera, suggesting a recent origin for this gene. Two of the targets of Bicoid in *Drosophila*, *orthodenticle* and *hunchback*, appear to be sufficient to regulate anterior patterning in *Tribolium* and this likely reflects the ancestral mode of anterior patterning and segment formation in insects. Following the evolution of the *bicoid* gene in the lineage that gave rise to the higher dipterans, Bicoid must have acquired the ability to regulate the zygotic expression of *orthodenticle* and *hunchback* and thus assumed its role at the top of the anteroposterior patterning hierarchy.

**Shifts in trunk Hox gene expression that mirror changes in arthropod body architecture**

The diversity of arthropod segmental identities suggests that evolution of the *Hox* genes has contributed to the radiation of arthropod forms. These genes have an ancient role in patterning the anteroposterior axis of bilaterians, and they likely regulate the development of distinct segmental identities in all arthropods and onychophora. As transitions in appendage morphology and in body regions occur at different axial positions between the major arthropod groups, the *Hox* genes are key candidates for the genetic players underlying arthropod diversity. Indeed, some of the morphological differences within the arthropod-onychophoran clade correlate with changes in *Hox* gene expression.

We are already familiar with the pattern of *Ubx* expression in *Drosophila*, where the anterior boundary initially falls within the first abdominal segment and later extends into the third thoracic segment. Posteriorly, *Ubx* expression extends through most of the abdomen. The anterior boundary of *Ubx* lies near the transition between the thorax, which bears the walking limbs and flight appendages, and the limbless abdomen. Genetic analysis in *Drosophila* has
shown that \textit{Ubx} regulates the distinct segmental identities of T3 and A1. In particular, the \textit{Ubx} gene represses leg development in A1 and is responsible for the morphological differences between the wing and the haltere (see Chapter 3). An anterior boundary of \textit{Ubx} expression in T3 is characteristic of all insects, reflecting the conserved insect body plan.

The anterior boundary of \textit{Ubx} gene expression, however, lies at a different segmental position within the trunks of crustaceans, myriapods, chelicerates, and onychophora (Fig. 5.2). For example, the expression pattern of \textit{Ubx} in a primitive crustacean lineage, the brine
shrimp *Artemia*, is quite different from *Drosophila* and other insects. In *Artemia*, the anterior boundary of *Ubx* falls at the anterior of the thorax, at the transition from the gnathal head segments (which bear feeding appendages) to the thoracic segments (which bear swimming appendages). Thus the expression of this *Hox* gene marks a transition in appendage morphology along the *Artemia* anteroposterior axis, but *Ubx* is expressed at a more anterior position relative to the insects.

Other, more derived crustacean lineages that possess specialized thoracic limbs exhibit different anterior boundaries of *Ubx* expression (Fig. 5.3). In some malacostracan and maxillopod crustacean lineages, one to three pairs of thoracic limbs are reduced in size and are used as feeding appendages (maxillipeds). In these organisms, the anterior boundary of *Ubx*/abd-A protein expression consistently lies to the posterior of any maxilliped-bearing thoracic segments. Interestingly, in some species, the loss of *Ubx*/abd-A expression in anterior thoracic limbs precedes the developmental transition from a larger limb morphology early in embryogenesis to a smaller maxilliped later in development. In these crustaceans, a shift in *Hox* gene expression anticipates the formation of specialized thoracic limbs.
The myriapods exhibit a body organization that differs from the body organizations of both the insects and crustaceans, and the boundary of Ubx protein expression occurs at a different segmental position (Fig. 5.2). In the centipede, the anterior boundary of Ubx/abd-A expression lies within the second trunk segment and extends through much of the homonomous trunk. Again, this boundary of Hox expression marks a transition in limb identity. Early in development, no Ubx protein is expressed in the developing poison claw on the first trunk segment, whereas it is expressed in all of the developing walking legs on the remaining trunk segments. In the fourth living arthropod class, the chelicerates, the anterior boundaries of Ubx and abd-A expression fall within the limbless opisthosomal segments (Fig. 5.2).

In summary, the relative positions of the anterior boundaries of Ubx and abd-A expression have shifted between the various arthropod classes, and even within orders of crustaceans. These shifts correlate with differences in body organization—indeed, with the morphology of thoracic limbs in insects, crustaceans, and myriapods.

In contrast to the highly diversified and patterned segments of arthropods, the onychophoran body plan includes fewer distinct segmental identities and a homonomous trunk. Where are the onychophoran orthologs of the trunk Hox genes Ubx and abd-A expressed? One might have guessed that their deployment would be similar to the arthropods, in which these Hox genes are expressed in most of the trunk segments. In reality, Ubx/abd-A protein expression is limited to the extreme posterior of the onychophoran embryo, in the last pair of lobopods and in the terminus (Fig. 5.2). Although Hox genes are conserved between onychophora and arthropods, clearly their expression patterns have changed significantly during their independent evolutionary history.

**Hox genes and the evolution of arthropod heads**

Modern arthropod classes have a more conserved organization of head segments than of trunk segments, with the exception of the chelicerates. Arachnids, for example, have paired walking appendages on their prosomal segments, the anterior body region of chelicerates. These prosomal segments exhibit patterns of Hox gene expression that correspond to the expression of the anterior Hox genes lab, pb, Dfd, and Scr in crustaceans and insects (Fig. 5.4). In arachnids, these genes have broadly overlapping expression domains, including a segmentally restricted Hox-like pattern of expression of the Hox3/zen ortholog. These Hox gene expression patterns suggest that the chelicerate prosoma is roughly equivalent to the head segments of insects and crustaceans.

The conservation of insect and crustacean head segmentation and patterning correlates with generally similar patterns of expression of anterior Hox genes (Fig. 5.4). The primary exception involves the regulation of the pb gene, which is expressed in different segments in insects and crustaceans and even among different insect orders. The insect gnathal head segments where pb is expressed form the mouthparts—namely, the mandible, maxilla, and labium. In crickets, beetles, and flies, the maxilla and labium are jointed and bear distal palps (compared with the stubby mandible); pb is expressed in both appendages. By contrast, hemipteran insects (including milkweed bugs and bedbugs) are characterized by specialized piercing-sucking mouthparts, where the mandibular and maxillary appendages form morphologically similar styles. The milkweed bug exhibits pb expression only in the labium. This more limited pattern of pb expression correlates with the coordinated development of the milkweed bug’s mandible and maxilla and reflects the loss of a patterning role for pb in the
maxillary appendages. The evolutionary shifts in \( pb \) expression observed in insects within the distal gnathal appendages break the usual colinearity of \( Hox \) expression and likely represent a derived condition relative to other arthropods.

**Annelid \( Hox \) expression patterns**

The annelids, or segmented worms, are a second protostome phylum characterized by a body plan made up of repeated units. Whereas the segments of the earthworm and the leech do not have many distinguishing external characters, the polychaete annelids are characterized
by diverse segmental identities that correlate with domains of \textit{Hox} gene expression (Fig. 5.5). Annelid \textit{Hox} genes are expressed in a nested, collinear fashion reminiscent of the expression seen in arthropods and vertebrates. Again, the anterior boundaries of orthologous \textit{Hox} genes fall at different segmental positions between annelid lineages. The expression domains of the leech orthologs of \textit{Hox1/\lab}, \textit{Hox4/Dfd}, and \textit{Hox5/Scr} are shifted to the anterior by at least one segment relative to the polychaete \textit{Chaetopterus} (Fig. 5.5).

**Correlation of vertebrate axial patterning with \textit{Hox} expression domains**

In vertebrates, changes in the number of vertebrae within regions of the vertebral column correlate with \textit{Hox} expression patterns in the paraxial mesoderm. The transition from one type of vertebra to another corresponds to the anterior limits of expression of specific \textit{Hox} genes (Fig. 5.6). For example, the anterior expression boundary of \textit{Hoxc6} falls at the transition of cervical vertebrae to thoracic vertebrae in mice, chickens, geese, and frogs. Each of these animals possesses a different number of cervical vertebrae, however, and the boundary of \textit{Hoxc6} lies at a different axial position relative to the head. In addition, the thoracic-to-lumbar transition, while occurring at different somite levels, appears to be associated with the expression of \textit{Hoxa9}, \textit{Hoxb9}, and \textit{Hoxc9}. The deployment of these \textit{Hox} genes consistently reflects both the relative position of vertebral identities and the number of vertebrae in each region. These observations suggest that changes in vertebral organization between different vertebrate orders evolved in concert with relative expansions or contractions in \textit{Hox} expression domains.

Another, more dramatic example of shifts in vertebrate \textit{Hox} expression domains is found in snakes. In most tetrapods, \textit{Hoxc6} and \textit{Hoxc8} are expressed in thoracic somites and are required to specify thoracic vertebral identity. The anterior boundary of \textit{Hoxc6} expression marks the cervical-to-thoracic transition; the posterior boundary of \textit{Hoxc8} falls at the position of the hindlimb, just anterior to the lumbar region. The vertebral column of snakes, however, does not have a clear cervical–thoracic boundary. In pythons, expression of \textit{Hoxc6} and

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**Figure 5.5**  
Expression patterns of \textit{Hox} genes in annelids  
The expression domains of the \textit{Hox} genes \textit{Hox1}, \textit{Hox2}, \textit{Hox3}, \textit{Hox4}, and \textit{Hox5} are depicted below the anterior segments of two annelids. The first nine segments of the polychaete (the A region) bear similar appendages; the B region has a different appendage morphology on each segment. The leech has no appendages, but anterior segments are grouped into R and M regions. Annelid \textit{Hox} genes are expressed in large overlapping domains in polychaetes and leeches, primarily in the nervous system. The anterior boundaries are generally not conserved between leeches and polychaetes, as the leech expression patterns tend to shift to more anterior segments.
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Figure 5.6

*Hox* genes and the evolution of tetrapod axial identities

Differences in the axial organization of tetrapods are reflected in shifts in *Hox* gene expression domains between animals. (a) The anterior boundaries of expression of several *Hox* genes in the paraxial mesoderm are shown beneath the somites (circles) and vertebrae (squares) of the mouse, chick, goose, and python body plans. In mammals and birds, which have distinct cervical (green) and thoracic (purple) axial regions, the anterior boundary of the *Hoxc6* gene lies at the cervical–thoracic transition, even though the axial position (somite number) of this transition falls at a different position in each organism. Similarly, the anterior boundary of *Hoxc8* expression lies within the thoracic region of chicks and mice; the *Hoxa9*, *Hoxb9*, and *Hoxc9* boundaries lie near the thoracic–lumbar transition, and the *Hoxa9* boundary lies near the lumbar–sacral transition. In the python, the *Hoxc6* and *Hoxc8* genes have a more anterior expression boundary, reflecting the expanded thoracic vertebral identities of the snake body plan. (b) Expression of the mouse *Hoxc8* gene in the thoracic region (the extent of high levels of expression is indicated by arrows). (c) Expression of the snake *Hoxc8* gene extends through the anterior of the axial skeleton (indicated by arrows and arrowheads).

Hoxc8 extends far to the anterior, up to the cranial region (Fig. 5.6). The posterior boundaries lie at the level of the (vestigial) hindlimb, as in other tetrapods. Within the domain of Hoxc6 and Hoxc8 expression, all of the python vertebrae bear ribs, indicating thoracic identity. Interestingly, a subset of cervical characters is present on the most anterior rib-bearing vertebrae of pythons, suggesting that thoracic identities overlie the ancestral cervical identities near the head. Thus the loss of the snake’s neck and the expansion of its rib-bearing vertebrae are correlated with the anterior shift in the expression of Hoxc6 and Hoxc8.

**How do Hox domains shift during evolution?**

Each of the bilaterian phyla that have metameric body organization exhibits relative shifts in Hox boundaries between lineages. These differences evolved through changes in the regulation of Hox gene expression during the radiation of arthropod, vertebrate, and annelid body plans. Mechanistically, shifts in Hox expression domains could result from changes in the expression or activity of the proteins that regulate the expression of Hox genes. Alternatively, changes could evolve within the cis-regulatory regions of the Hox genes themselves. The cis-regulatory elements that mediate Hox expression may evolve to respond differently to upstream regulators (activators or repressors), thereby changing the relative pattern or timing of Hox expression.

Circumstantial evidence from comparisons of vertebrate Hox genes suggests that evolutionary changes in Hox cis-regulation represent diversifying patterns of paralogous Hox genes. Vertebrate Hox gene homologs that arose from duplications of the ancestral chordate Hox cluster sometimes have different boundaries of expression. For example, the chick Hoxa9, Hoxb9, and Hoxc9 genes share a common anterior boundary of expression, but the expression of the Hoxd9 gene has shifted to the posterior (Fig. 5.6). These genes evolved from a single common ancestral Hox9 gene, and they initially had identical cis-regulatory regions following cluster duplication events. During their subsequent independent evolution, relative changes in their deployment apparently occurred because of sequence divergence within these cis-regulatory regions.

Direct evidence for cis-regulatory element evolution has been found for the vertebrate Hoxc8 gene. The anterior boundary of Hoxc8 expression lies at a different somite position in mice and chicks, but in a similar location within the thoracic region (Fig. 5.6). A comparison of homologous Hoxc8 cis-regulatory elements in chicks and mice indicates that the axial shift in Hoxc8 expression evolved because of a small number of sequence changes within the highly conserved Hoxc8 cis-regulatory region (Fig. 5.7). The activity of the chick cis-regulatory has also been tested in mice, and expression from the chick element appears to be initiated at a more posterior axial position than expression from the mouse element. This shift correlates with the more posterior somite location of the chick thoracic region and the boundary of native chick Hoxc8 expression. The functional difference between these homologous chick and mouse Hoxc8 elements suggests that the Hoxc8 cis-regulatory region evolved during the divergence of mammals and birds and reflects the different skeletal organizations of these groups.

**Morphological diversity within a conserved body plan**

Although some shifts in Hox expression correlate well with transitions in axial patterning (as discussed previously for arthropods, vertebrates, and annelids), morphological diversity also
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occurs among more closely related animals that share a particular body plan. The insects, which generally have conserved boundaries of Hox expression domains, exhibit differences in the shape, size, and pattern of their segments and appendages. How have the morphologies of different insects evolved at a genetic level? The answer to this question, again, lies in evolutionary changes in gene regulation.

Three mechanisms are implicated in the diversification of limb patterning between lineages. First, within a stable pattern of Hox gene deployment, regulatory changes downstream
of the *Hox* genes have led to the diversification of the shapes and patterns of homologous limbs. A similar process underlies the evolution of morphological differences between vertebrate forelimbs and hindlimbs. Second, some interesting developmental modulations of *Hox* expression patterns within fields have morphological and evolutionary consequences. These upstream differences in *Hox* regulation may provide a glimpse of how larger shifts in *Hox* expression evolve. Third, some *Hox* proteins have evolved novel functions and therefore novel patterning roles in different lineages that contribute to morphological diversification.

**Evolution of the limbless insect abdomen**

In addition to patterning arthropod appendage morphology, the *Hox* genes played a role in the evolution of the number of limbs in insects. The insect thorax bears three pairs of walking legs and two pairs of wings, but the abdomens of modern adult insects characteristically lack appendages. As many other arthropods have abdominal limbs, the insects are thought to have evolved from an ancestor with limbs on all trunk segments. Some primitive insect fossils have small abdominal leglets, which may represent an intermediary step in the evolution of the limbless insect abdomen. Thus limb development was repressed specifically in the abdominal segments of the insects.

The genetic mechanism underlying the repression of abdominal limbs has been elucidated in *Drosophila*. Abdominal limb development is prevented during embryogenesis via repression of the *Distal-less* (*Dll*) gene in abdominal segments. The Ubx and abd-A proteins directly repress an early-acting *Dll* cis-regulatory element, which drives expression of this gene in limb primordia in the embryonic thoracic segments. At this stage of embryogenesis, *Ubx* expression does not overlap *Dll* in the third thoracic segment, and *Hox* repression of *Dll* remains limited to the abdominal segments. In *Drosophila*, *Hox* proteins interact directly with the regulatory network controlling limb development.

In contrast to insects, other arthropods bear limbs on segments that express *Ubx* and *abd-A* (Fig. 5.8). In crustaceans and myriapods, expression of the Ubx/abd-A proteins overlaps with early *Dll* protein expression in trunk appendages; *Ubx* and *abd-A* do not appear to repress limb development or the *Dll* gene. In collembolans (a primitive insect), three abdominal appendages express *Dll* in the presence of high levels of *Ubx/abd-A*. Thus the repression of *Dll* must have evolved in the insect lineage. The evolution of *Dll* repression and the limbless insect abdomen appears to have involved both the evolution of Ubx and abd-A binding sites within *Dll* regulatory elements and the evolution of functional motifs within the Ubx protein. When expressed in *Drosophila*, the Onychophoran and brachiopod (*Artemia*) Ubx proteins lack the ability to repress *Drosophila Dll* and limb formation. Surprisingly, these orthologous Ubx proteins cannot repress *Dll* for apparently different reasons. In one case (*Artemia*), the carboxy-terminal region of the protein acts to block, or mask, the ability of *Artemia* Ubx to repress *Drosophila Dll* in *Drosophila*. By contrast, the onychophoran Ubx protein is unable to repress *Drosophila Dll* because it lacks a glutamine- and alanine-rich motif (which also maps to the carboxy-terminal region). This motif is conserved among all insect Ubx proteins and acts biochemically as a repressor (Fig. 5.9). The evolution of repressor activity within the Ubx protein, perhaps in concert with the evolution of Ubx and abd-A binding sites within *Dll* regulatory elements, may have sculpted the limbless abdomen of the insect body plan.
A comparison of different insect orders indicates that the repression of Dll by Ubx and abd-A may have evolved in at least two stages. Many insect embryos develop a small appendage on the A1 segment called a pleuropod. In beetles and grasshoppers, Ubx and Dll protein expression overlap during the development of this pleuropod (Fig. 5.10). In fact, Ubx is required to properly pattern the pleuropod limb. No limbs develop within the abd-A domain, and abd-A represses limb development in other abdominal segments. The more primitive condition suggests that abd-A repression of Dll may have evolved first. Then, in derived insect orders that do not have a pleuropodial appendage on A1 (such as diptera, which includes Drosophila), Dll became repressed by Ubx as well.

Crustacean Hox proteins have also gained novel functions that affect limb morphology. As described above, the carboxy-terminal region of the Artemia Ubx protein that modulates Ubx activity contains potential phosphorylation sites not found in insect Ubx proteins (Fig. 5.9). In another crustacean, the water flea Daphnia, the Antp protein has evolved the ability to...
repress Dll in ventral regions of the first trunk appendage. This appendage lacks a ventral comb-like feeding structure and is smaller than more posterior trunk appendages. The ability of *Daphnia* Antp protein to repress Dll maps to the amino-terminal region of Antp, which does not contain sequence similarity to other arthropod *Antp* genes. Thus, Hox protein evolution in multiple lineages contributes to shaping arthropod body plans.

**Evolution of insect wing number**

The scenario depicting the reduction and elimination of limbs on the insect abdomen may be paralleled by the history of the evolution of the number of insect wings. All modern winged insects bear two pairs of wings: forewings on the second thoracic segment and hindwings on the third thoracic segment. The invention of wings was a major event in the evolution of insects, catalyzing their radiation and domination of terrestrial habitats.

Insect wings are thought to have evolved only once, early in the insect lineage (the origin of insect wings will be discussed in more detail in Chapter 6). The insect fossil record suggests that, in the primitive state, wing-like structures formed on all of the trunk segments. Fossils of early insects from the Carboniferous (circa 325 Ma) include aquatic larval forms with segmentally iterated, wing-like projections that may have played a functional role in respiration. More recent fossilized mayfly nymphs had larger wing-like structures on thoracic segments...
and smaller winglets on the abdomen. The fossil record suggests that over time abdominal wing structures became reduced in size and ultimately disappeared, as did the wing-like structures on the first thoracic segment (Fig. 5.11).

The restriction of insect wings to the second and third thoracic segments suggests that the Hox genes sculpted the evolution of wing number and pattern. Indeed, genetic analysis has established that specific Hox genes repress the initiation of wing development in particular segments. In Drosophila, wing development is repressed in the first thoracic segment by Scr and in the abdominal segments by Ubx, abd-A, and Abd-B. The evolution of Hox repression of wing development over the course of insect evolution could explain the reduction and loss of wings on these segments (Fig. 5.11). Initially, Scr, Ubx, abd-A, and Abd-B may have modified the expression of wing-patterning genes to control the size of abdominal and T1 winglets. Over time, Hox regulation completely suppressed the formation of wings to sculpt the modern insect body plan. The evolution of Hox protein binding sites in the cis-regulatory regions of genes required for wing formation could account for the repression of wing development during evolution.

Figure 5.10
Dll and Ubx are coexpressed in the pleuropod appendages of beetles and grasshoppers

The evolution of Hox repression of abdominal limb development and Dll expression in insects evolved in at least two steps. First, abd-A gained the ability to repress limb formation in most abdominal segments (A2–A7) near the base of the insect clade. In the ancestral lineage of butterflies and flies, Ubx also evolved the ability to repress limb formation and Dll expression in the first abdominal segment (A1). Dipterans do not form an A1 appendage, and Dll (green) is not expressed in that segment (top right). In beetles and grasshoppers, a small pleuropod appendage forms in the A1 segment (bottom right). Both Dll (green) and Ubx (red) are expressed in these pleuropods (overlap in yellow), indicating that Ubx does not repress pleuropod formation or Dll expression in these animals.

The segmental domains of Hox gene expression did not change during insect evolution and were probably similar in ancestral insects (shown as abbreviations in each body segment: S, Scr; A, Antp; U, Ubx; AA, abd-A; B, Abd-B). The evolutionary progression of insect wing number reflects the modification or repression of wing development by Scr, Ubx, abd-A, and Abd-B. Primitive insects were wingless (a), and the first insects with wing-like structures were larval forms (b, c). Wing development became repressed in the first thoracic segment (red/Scr) and in abdominal segments (yellow/Ubx, green/abd-A, blue/Abd-B) of fossil (d) and modern (e, f) adult insects. In dipterans (f), Ubx also regulates the reduced size and modified shape of the haltere.

Diversification of insect wing morphology

Although all modern winged insects bear two pairs of wings, many structural, functional, and morphological differences exist between forewings and hindwings both within and between species (Fig. 5.12). For example, the wings of dragonflies (Odonata) appear rather similar, but the forewings of beetles (Coleoptera) have been modified into hardened coverings that protect the hindwings. Butterfly (Lepidoptera) forewings and hindwings are often of similar size, but have evolved different shapes and color patterns. In flies (Diptera), the hindwing has been reduced to the haltere, a small balancing organ with a different biomechanical function.

The Ubx gene regulates the differences between forewing and hindwing patterns within species. Ubx is expressed in the developing hindwing of all insects studied, and it controls hindwing-specific patterning (Fig. 5.13). Yet, the hindwings of different insects such as butterflies, beetles, and flies are all different from one another. This diversity reflects the regulation by Ubx of different sets of target genes in each animal.

Recall that in the Drosophila haltere, Ubx represses the wing-patterning genes that are required for the growth and flattening of the wing and for the development of wing veins and sensory organs (see Chapter 3). By contrast, Ubx does not repress the orthologs of these genes in the butterfly. The butterfly hindwing is large and flat, and it has similar venation and sensory organ patterning to the forewing; the developmental program that controls these features is deployed in both wing pairs. In certain butterflies, Ubx regulates target genes that control the wing shape, color pattern elements, scale pigmentation, and scale morphology—all features that have no counterpart in the fruit fly. Thus the homologous hindwings of flies and butterflies have evolved distinct morphologies in part because of independent changes in the regulatory connections between Ubx and genes in the wing hierarchy (Fig. 5.14).
Figure 5.13

**Ubx expression in insect hindwings**

Ubx is expressed in insect hindwings that have diverse morphologies. (a) In *Drosophila*, the wing is much larger than the balloon-like haltere. (b) Ubx protein (green) is expressed in the developing *Drosophila* haltere, but not in the wing. (c) In the butterfly *Precis coenia*, the hindwing is large and flat like the forewing, but has different color patterns and scale morphology. (d) Ubx protein (green) is also expressed in the butterfly hindwing.


Figure 5.14

**Evolution of insect hindwing morphological diversity**

Evolution of the Ubx regulation of hindwing development has contributed to the divergence of insect hindwing morphology. From an ancestral insect with similarly patterned forewings and hindwings (bottom), insect hindwings evolved that differ from other homologous hindwings in other species and from the forewings of the same species (top). This diversity is associated with evolutionary changes in the set of Ubx-regulated target genes in each lineage, in addition to changes in the wing developmental program itself.

Sets of \textit{Ubx}-regulated target genes may have diverged between species through changes in the \textit{cis}-regulatory elements that control gene expression in the wing field. For instance, the evolution of \textit{Ubx} binding sites within a wing-specific \textit{cis}-regulatory element could have modified gene expression in the hindwing in a given lineage. Alternatively, the evolution of a new activating \textit{Ubx} binding site could have created a novel expression pattern in the hindwing. 

The evolution of developmental responses to environmental cues has also contributed to the regulation and evolution of insect wing morphology and number. For example, in ants reproductive individuals have wings while the worker and soldier castes are wingless. This difference is regulated by hormonal and environmental signals that are independent of \textit{Hox} gene input. Evolution of \textit{cis}-regulatory elements responsive to these cues in genes that promote wing development may underlie the evolution of wing polyphenism in ants. Due to the modular nature of the wing developmental hierarchy, such evolutionary changes in \textit{cis}-regulatory elements may alter wing formation and morphology in some developmental programs without disrupting wing development entirely.

\textit{Modulations in Hox expression patterns within fields that contribute to insect diversity}

Homologous insect appendages have become diversified in the context of a highly conserved pattern of \textit{Hox} gene expression along the main body axis. Nevertheless, some important differences exist in \textit{Hox} deployment within the homologous limb fields of different insects.

We return to the subject of insect abdominal limbs to discuss an apparent evolutionary \textit{atavism}—that is, reversion to a more ancestral state. Recall that, in general, adult insects do not have abdominal limbs. However, butterfly caterpillars (and some other insect larvae) do have abdominal limb-like structures called \textit{prolegs}, which are used in locomotion. These stubby prolegs are not jointed like thoracic limbs, but their development is marked by \textit{Dll} expression in embryos. How does this \textit{Dll} expression escape repression by \textit{Ubx} and \textit{ Abd-A} in the abdomen?

In early stages of larval development, there is uniform expression of \textit{Ubx} and \textit{ Abd-A} and no sign of \textit{Dll} expression or proleg outgrowth in the butterfly abdomen. Later, however, the expression of \textit{Ubx} and \textit{ Abd-A} is turned off within small clusters of cells in the A3–A6 abdominal segments, just before proleg development begins. These cells then begin to express \textit{Dll} and to grow out from the body wall (Fig. 5.15). Thus the apparent evolutionary reversal that allows larval prolegs to develop in lepidopterans appears to be a segment-specific modification of abdominal \textit{Hox} gene expression. Interestingly, sawfly (\textit{Hymenoptera}) larvae, which also possess abdominal prolegs, circumvent \textit{Hox}-mediated limb repression differently than caterpillars, by initiating proleg formation independent of \textit{Dll} expression.

A second example of an evolutionary modification of insect limb patterning that is correlated with changes in \textit{Hox} expression within the limb field has been identified in several \textit{Drosophila} species. Subtle differences have been noted in the pattern of tiny leg hairs (trichomes) on the second thoracic legs of \textit{Drosophila melanogaster}, \textit{Drosophila simulans}, and \textit{Drosophila virilis} (Fig. 5.16). In \textit{D. melanogaster}, trichomes do not form in a certain region of the posterior femur of the T2 leg, resulting in a small “naked valley” of smooth epidermal cuticle. This “naked valley” is associated with high levels of \textit{Ubx} expression, and indeed \textit{Ubx} acts to repress trichome development. \textit{D. simulans} has a larger “naked valley” and high levels of \textit{Ubx} expression over an expanded region of the T2 femur. The more distantly related
Drosophila virilis lacks a “naked valley” in the T2 trichome array and exhibits only low levels of Ubx expression.

Careful genetic dissection of the differences in trichome patterning between D. melanogaster and D. simulans has indicated that evolutionary differences in the level and extent of Ubx expression map genetically to the Ubx locus. This finding indicates that the source of evolutionary divergence in trichome patterning between D. melanogaster and D. simulans lies within these fruit flies’ respective Ubx cis-regulatory control regions. Since the time of the last common ancestor of these species, sequence changes must have evolved in Ubx cis-regulatory elements that led to a relative increase in Ubx expression in the T2 femur of D. simulans. In this way, regulatory evolution at the Ubx locus contributed to a quantitative change in T2 leg morphology between Drosophila species.

This example demonstrates a potential link between the larger shifts in Hox expression seen among distantly related arthropods and the generally similar patterns of Hox expression between insects. That is, if even slight differences can evolve in domains of Hox gene expression between fairly closely related species, then one can begin to conceive of how larger shifts might arise over greater periods of evolutionary time.
Vertebrate limb diversity: regulatory changes downstream of other selector genes

The vertebrates also possess a wide diversity of limb types, ranging from fish fins to bat wings to human arms. The paired pectoral and pelvic fins of fish are serially homologous appendages that are homologous to tetrapod forelimbs and hindlimbs, respectively. The evolution of tetrapod forelimb and hindlimb characters is often linked. For example, digits
arose at the same time in both sets of paired appendages in Devonian tetrapods. Other, more recent adaptations such as hooves also appeared simultaneously in the forelimbs and hindlimbs of ungulates. Such features likely evolved in concert through the shared developmental genetic pathways that govern the growth and patterning of both limbs (see Chapter 3). The appearance of the same patterning elements in both the forelimb and the hindlimb likely results from the evolution of novel regulatory interactions downstream of a shared limb genetic hierarchy.

Of course, in many familiar cases the serially homologous appendages of vertebrates have evolved independent morphologies. For example, the forelimbs of birds and bats were independently modified into flight structures, having very different shapes and functions from the respective hindlimbs of these animals. Kangaroos have small, grasping forelimbs and large, muscular hindlimbs, as do many tetrapods that walk (or jump) primarily on two legs. In chickens and mice, the independent developmental characters of forelimbs and hindlimbs are controlled by limb selector genes that are differentially expressed (see Chapter 3). The same selector genes operate in other vertebrates to regulate the morphological diversification of serially homologous limbs.

The T-box transcription factor \( Tbx5 \) is expressed in the forelimbs of mice, chickens, and newts as well as in the pectoral fins of teleost and cartilaginous fish. A related gene, \( Tbx4 \), and the homeodomain-containing \( Pitx1 \) gene are expressed specifically in the hindlimbs of these vertebrates. As in the chick and mouse, these genes likely regulate different sets of target genes to pattern morphological differences between the serially homologous forelimbs and hindlimbs of the newt. Furthermore, as the forelimbs (wings) of birds are morphologically distinct from the forelimbs of mammals and amphibians, limb selector genes probably regulate different sets of target genes in these different lineages (Fig. 5.17). Extrapolating to other vertebrates, a common forelimb selector gene, \( Tbx5 \), probably regulates different target genes in bats and whales and humans.

The story of vertebrate limb diversification both between and within a species is analogous to the regulation of insect hindwing pattern by \( Ubx \). In both vertebrates and insects, field-specific selector genes differentially pattern serially homologous body elements. Homologous parts between lineages, such as vertebrate forelimbs or insect hindwings, express orthologous selector genes but have diverse morphologies, reflecting evolutionary changes in the array of their respective downstream target genes (Fig. 5.17).

**REGULATORY EVOLUTION AND THE DIVERSIFICATION OF HOMOLOGOUS BODY PARTS**

This chapter discussed several case studies of the genetic mechanisms underlying the divergence of arthropod and vertebrate body plans. In each case, morphological change is related to a change in the expression pattern of developmental genes. Shifts in \( Hox \) expression domains between arthropod, vertebrate, and annelid classes correlate with large differences in body organization—particularly transitions in appendage morphology and vertebral identity. Modifications in \( Hox \) expression domains within the limb fields of more closely related organisms underlie such morphological features as butterfly prolegs and fruit fly trichome patterning. When \( Hox \) or other selector gene domains are conserved, evolutionary changes in the regulation of downstream target genes facilitate the diversification of homologous parts
Figure 5.17
Homologous structures diverge through changes in the target genes that are regulated by conserved selector genes.

Schematized views of the evolution of insect hindwings (a) and vertebrate forelimbs (b). The conservation of selector gene expression in insect hindwings (Ubx) and vertebrate forelimbs (Tbx5) suggests that the ancestral hindwings of insects and ancestral forelimbs of vertebrates also expressed these genes. While the selectors regulated certain target genes (colored boxes) in the ancestral appendage (left), these sets of genes changed over the course of evolution, resulting in dramatically different morphologies of homologous limbs in modern lineages (right).
between organisms. For example, the *Hox* gene *Ubx* is expressed in all insect hindwings and the limb selector gene *Tbx5* is expressed in all vertebrate forelimbs, yet these genes regulate different sets of target genes in different lineages; the result is insect and vertebrate appendage diversity (Fig. 5.17).

The flexibility of regulatory networks and the modular nature of *cis*-regulatory control regions allow genetic interactions to evolve without changing the number of genes or even the primary sequence of the proteins encoded by animal genomes. Evolutionary modification of the regulatory interactions between developmental transcription factors and target genes allows different animal morphologies to be built using the same genetic and structural elements. Chapter 6 will extend our discussion of regulatory evolution to include the co-option of developmental regulatory genes in new fields or to pattern new structures. The evolution of novelty will be told as a story of using “old genes to perform new tricks.”

**SELECTED READINGS**

**General Reviews**


**Arthropod Segmentation**


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Chapter 5: Diversification of Body Plans and Body Parts


**Arthropod and Annelid Hox Gene Expression Domains and Segment Identities**


**Vertebrate Hox Expression Domains and Axial Patterning**


DNA to Diversity: Molecular Genetics and the Evolution of Animal Design


**Diversification of Insect Segmental Morphology: Wings and Legs**


Suzuki Y, Palopoli MF. Evolution of insect abdominal appendages: are prolegs homologous or convergent traits? *Dev Genes Evol* 2001; 211: 486–492.


**Vertebrate Limb Selector Genes**


CHAPTER 6

The Evolution of Morphological Novelties

Morphological novelties abound in the history of animal evolution and have been essential to the diversification of animal forms. The scope of novelties encompasses many familiar structures, from antlers to ear bones, from whale tails to panda thumbs. New structures and pattern elements are often key features that distinguish animal groups. For example, the vertebrates are characterized by an elaborate forebrain, the neural crest, and cartilage—all novelties that were critical to the success of the vertebrate lineage and that allowed the evolution of further innovations such as teeth and jaws.

Structural novelties (or “key innovations”) are associated with adaptive radiations into new ecological territories. The movement of vertebrates onto land is tied to the evolution of the tetrapod limb; the escape of insects into the air required the evolution of wings. The distinctive molar tooth shape has evolved independently several times in association with herbivory. Other novelties such as feathers and butterfly scales permit the display of colors and color patterns used for communication or predator avoidance.

New structures require the evolution of new developmental programs. To understand the origin of morphological novelties, we must look again to the genetic control of development. Have new developmental regulatory genes evolved to sculpt novel body parts? Or are “old” genes recruited for new patterning roles? And, if regulatory genes are reused, are they co-opted individually or as part of larger, preexisting circuits?

This chapter uses case studies to examine various ways in which new structures and pattern elements evolved through changes in developmental gene regulation. In several instances, shared aspects of development and regulatory gene expression reflect the evolution of novelties from preexisting ancestral structures. Such novelties evolved through downstream regulatory changes in developmental programs. In other cases, “old” developmental regulatory genes evolved a new role in the formation...
of novel pattern elements. In the best-documented examples, groups of genes or entire genetic circuits were recruited to carry out a new developmental function.

**WHAT IS MORPHOLOGICAL NOVELTY?**

The history of evolutionary biology is replete with operational definitions of novelty. For our purposes, which are primarily to understand the developmental and genetic basis of the evolution of novel animal forms and patterns, a “novelty” is defined as a structure or pattern element, or even an entire body plan, that has a new adaptive function. This chapter focuses on the best examples of morphological novelty for which developmental genetic knowledge has been elucidated. We do not address other forms of innovation, though they are fascinating in their own right, such as the evolution of physiological adaptations through protein evolution (for example, antifreeze proteins, lens crystallins, keratins, lactose synthesis, immune systems), because they do not concern morphological evolution per se.

Quantitative morphological variation, even when extreme, is not considered novel unless it encompasses a fundamental functional shift. Thus most homologous body parts are not considered novelties, even when the range of a vertebrate forelimb, for example, extends from the gigantic size of a whale flipper to the tiny arm of a tree shrew. Certainly, a gray area exists in terms of the degree of functional change considered sufficient to warrant classification as a novelty. Here, we limit our discussion to cases where a functional shift is accompanied by fundamental changes in development. These examples include the evolution of feathers, the origin of the insect wing and the spider spinneret, and the evolution of lepidopteran pigmented wing scales.

Other examples of novelty represent the evolution of a new structure, cell layer, or pattern element with no clear morphological antecedent. The cryptic origins of these innovations make them particularly interesting to evolutionary biologists. Several key innovations evolved in the chordate lineage, including the notochord, cranial placodes, the neural crest, and paired limbs. The wealth of developmental genetic knowledge about higher vertebrates and a few key outgroups, such as the cephalochordate amphioxus and urochordate ascidians, allows us to trace the evolution of some chordate novelties through comparisons of these taxa.

This chapter closes by analyzing radical transformations of body organization. Some ascidians have evolved direct development that eliminates the notochord, the defining structure of the chordate lineage. Snakes are “limbless tetrapods,” having an extended axial skeleton associated with a novel mode of locomotion. Although the loss of ancestral characters typically is not considered to be a novelty, in these examples key morphological characters disappear in the context of a new ontogeny. Other examples of body plan changes include turtles, which have an armored shell resulting from rearrangements of tetrapod characters combined with redeployment of specific cell types, and cephalopods and echinoderms, which exhibit novel body plans that are characterized by new structural components as well as reorganization of ancestral features.

The origins of most novelties have long puzzled biologists. Observing an anatomically and developmentally complex structure such as a bird feather, a butterfly wing, or the vertebrate brain does not provide insight into the means by which that structure evolved. Nevertheless, taking a comparative approach—integrating the study of paleobiology, comparative embryo-
Chapter 6: The Evolution of Morphological Novelties

...ology, and developmental genetics—provides a window into the history of some important morphological novelties. The evolution of these novelties involves changes in the regulation of transcription factors, signaling molecules, and structural proteins.

NOVEL FUNCTIONS FROM OLDER MORPHOLOGICAL STRUCTURES

The origin and evolution of avian feathers

True powered flight has arisen three independent times in vertebrate evolutionary history encompassing changes in the morphology and function of forelimb structures in the pterosaur, bird, and bat lineages. In all three cases, the leading edge of the wing is stabilized by the forelimb skeletal elements. But, in contrast to the pterosaurs and bats, which use a membranous lifting surface, the majority of the flight surface in birds consists of feathers. Feather-like structures have been found on fossils of several nonavian dinosaurs, suggesting that early feathers may have been used for camouflage, thermoregulation, or color display purposes. The subsequent co-option of feathers for flight involved changes in the shape and structure of feathers as well as numerous other changes in the anatomy and physiology of birds.

Like hair, nails, and scales, feathers are integumentary or epithelial appendages that arise from controlled proliferation of the epidermis. All epithelial appendages originate as epithelial placodes, or small regions of thickened epidermal cells, that will eventually give rise to individual outgrowths (Fig. 6.1a). The diversity of epithelial appendage morphologies belies the extensive similarities in their initial developmental stages. The spatial distribution of epithelial primordia in ordered arrays is established by a series of signals, involving the Shh, Wnt, FGF, Notch, and BMP (Bone Morphogenetic Proteins, members of the TGF-β superfamily) signaling pathways, between the mesenchymal (dermal) and epithelial (epidermal) tissue layers. The identity of epithelial appendages is determined by the underlying mesenchyme and affected by the timing and level of expression of these signaling molecules. Birds have both feathers and scales (on their feet) and feathers develop in response to high levels of Wnt and Notch signaling and low levels of BMP signaling, whereas scales develop in regions with inverse levels of these signals (Fig. 6.1b). Thus modulations in the strength or duration of these early signaling events appears to establish the identities of scales versus feathers.

Further development of the feather is regulated, in part, by Shh and Bmps. These two signaling pathways appear to be integrated in order to maintain a balance between proliferation (Shh) and differentiation (BMP) that is crucial to proper feather formation. Shortly after formation of the feather primordia in chicks and ducks, expression of Bmp2 and Shh becomes polarized with Bmp2 expression in the anterior and Shh expression in the posterior of each primordium. This same pattern of gene expression can be found in the developing scutate scales on the feet of chick and ducks and also in alligator scales (Fig. 6.1c). These data suggest that early Shh and Bmp2 expression was regulated in a similar manner in epithelial appendages of birds and crocodilians and was co-opted for further use in feather development.

After hatching, the feather primordia differentiate into the feather proper. A typical feather is a branched structure comprising a central rachis, a series of barbs attached to the rachis, numerous barbules connected to the barbs, and cilia or hooklets attached to the barbules. The Shh–BMP circuit is reused repeatedly during these later stages of feather development. The
Figure 6.1
Epithelial appendage development

(a) The early stages of epithelial appendage development (scales, feathers, and hair) begin with the formation of an epidermal placode. Signaling between the mesenchyme (dermis) and epithelium (epidermis) establishes the ordered array of placodes on the skin. Outgrowth of the placode is followed by the formation of a follicle, which supports growth and differentiation of feather and hair development. (b) The difference between the identities of avian scales and feathers reflects the timing and level of early signaling events. (c) Sequential redeployment of Shh-Bmp2 module during the development and evolution of a feather. Congruence between patterns of expression of the Shh-Bmp2 module in feathers and archosaurian scales (top), and the developmental theory of the origin and early evolution of feathers (bottom). The Stages I–IIIa of the developmental theory of feather evolution are proposed to have evolved by novel regulation of Shh-Bmp2 module expression. Stage I—the first, elongate tubular feather evolved from a primitive archosaurian scale by the derived distal Shh-Bmp2 co-expression (Event 1). Stage II—the first, branched plumulaceous feather evolved by the origin of derived longitudinal Shh-Bmp2 expression domains (Event 2) that created differentiated filaments from the tubular epithelium of the feather germ. The central–peripheral (c)–(ph) polarity of Shh-Bmp2 expression in the marginal plate epithelium between the barb ridges is shown in the inset. Stage IIIa evolved by the controlled dorsal (d) cessation and ventral (v) division of the longitudinal Shh-Bmp2 expression domains (Event 3) producing helical growth of barb ridges, indeterminate barb number, a rachis, serial fusion of barbs to the rachis, and a planar vane.

balance between the antagonistic activities of Shh and BMPs sets the number and spacing of the primary branches. Variations in the patterns of Shh and Bmp expression correlate with the changes in the organization of these branching structures in different types of feathers, suggesting that the co-option of these two signaling pathways has played a fundamental role in the origin and elaboration of feathers. The co-option of single genes, regulatory circuits, or entire regulatory hierarchies is a common theme in the development of evolutionary novelties.

The evolution of insect wings and spider spinnerets from ancestral gills

Arthropods were among the first metazoans to live on land. The transition from an aquatic to terrestrial environment required a number of changes in their morphology and physiology. Some of these changes represent the evolution of novel structures that led to adaptive radiations of specific arthropod lineages. For example, the evolution of insect wings is recognized as the key innovation behind the evolutionary success of the insects, which account for more than 75% of all known metazoan species. Likewise, the evolution and refinement of silk production from spinnerets was a key innovation that led to the evolutionary success of the spider lineage. The evolutionary origins of the insect wing and the spider spinneret are not clearly delineated in the fossil record, but molecular evidence suggests that these disparate structures may each be derived from the gills of ancestral arthropods.

Insect wings arose once during evolution, and the first wing-like structures, which may have been used for respiration or osmoregulation, appeared on each trunk segment of aquatic insect larvae in the early Carboniferous. Over the early course of insect evolution, wings became restricted to the adult thorax (see Chapter 5). At first, these structures might have allowed insects to skim across water surfaces; later, they evolved the ability to sustain gliding and powered flight. The earliest spider fossils document the presence of spinnerets on the opisthosomal segments, suggesting that silk was already in use during the early stages of spider evolution. Refinements in silk production, combined with changes in the positioning of the spinnerets and structure of the tarsal claws allowed spiders to use woven silk for multiple purposes. Just as flight allowed insects to escape predation, to catch prey, and to move to other habitats, silk provided spiders similar advantages via droplines, webs, and ballooning. The opportunities presented by these unique characters catalyzed the evolutionary radiation of the insect and spider lineages.

One hypothesis concerning the morphological origin of insect wings suggests that the wing evolved from a dorsal branch (epipodite) of a multibranched crustacean limb (Fig. 6.2a). In this scenario, the fusion of the base of the ancestral branched insect limb with the body wall led to the evolutionary displacement of the wing/epipodite away from the rest of the leg. This model also predicts that developmental regulatory similarities between insect wings and crustacean epipodites should reflect their common ancestry.

Indeed, the expression patterns of homologs of two Drosophila wing-patterning genes in crustacean appendages are consistent with the origination of insect wings from a dorsal limb branch in an ancestral arthropod. In the crayfish, the pdm gene is expressed in two distinct domains of the multibranched thoracic limb that are reminiscent of the two patterns of pdm (also known as nubbin) expression in the fly wing and leg, respectively. The pdm gene is expressed throughout a dorsal epipodite in crayfish thoracic limbs, much like the pattern observed in Drosophila wing imaginal discs, and in rings in the ventral, walking branch of crayfish thoracic limbs, much like the pattern seen in developing Drosophila legs (Fig. 6.2b,c).
Figure 6.2

**Evolutionary origin of the insect wing**

The insect wing and spider spinnerets may have evolved from a dorsal branch of an ancestral multibranched limb. (a) The proposed structural homology between the insect wing and (b) an ancestral dorsal leg branch (red). (c) In a crayfish, the Pdm protein is expressed throughout a dorsal branch of the third thoracic limb and in rings in the primary branch (walking leg). (d) The last common ancestors of all arthropods were aquatic creatures with branched appendages. The ventral branches (blue) of these appendages were used mostly for locomotion (e.g. legs), while the dorsal branches, called epipods (red), were used mostly for respiration and osmoregulation (gills). Endopods/legs are preserved in most arthropods. Epipods/gills are preserved in aquatic arthropods but modified or lost in terrestrial groups. In terrestrial arachnids (spiders and scorpions), a series of related primordia arise in posterior segments of the body. In spiders, the first primordium fails to develop further, the second gives rise to book lungs, the third gives rise to book lungs or to the lateral tubes of the tubular tracheae (depending on the group of spiders), and the more posterior ones give rise to the spinnerets. For simplicity, some appendages or appendage parts are not shown (for example antennae, exopods).

In the brine shrimp *Artemia*, *pdm* and a homolog of the insect wing dorsal compartment selector gene, *apterous*, are expressed in a dorsal lobe of the thoracic limbs. The parallels between the expression domains of *pdm* and *apterous* in crustacean epipodites and in insect wings suggest that these structures share a common history. This dorsal limb branch is used for respiration and osmoregulation in some extant crustaceans, consistent with the evolutionary origin of insect wings from a gill-like structure with respiratory function.

Comparisons with chelicerates suggest that dorsal limb epipodites have been repeatedly modified to form structures with highly divergent morphologies and functions. Horseshoe crabs, chelicerates that have retained an aquatic lifestyle, bear gill-like appendages on their opisthosoma segments. These book gills show uniform expression of *pdm*. Similarly, the tubular tracheae and book lungs (internal respiratory structures) of spiders express *pdm* as well as *apterous*. Thus, the respiratory role of the ancestral arthropod gill has been retained in some chelicerate as well as crustacean lineages. Spider spinnerets develop from the serially homologous opisthosomal primordia that give rise to these respiratory structures and they also show expression of *pdm* and *apterous* genes. These observations are consistent with the idea that spider spinnerets are derived from an ancestral respiratory structure, which was modified when early arachnids adopted a terrestrial lifestyle.

Examination of diverse arthropod species reveals that overlapping expression domains of *pdm* and *apterous* appear to be characteristic of structures that are derived from a specific epipodite or gill (Fig. 6.2d). Developmental genetic similarities between these disparate appendages suggest that regulatory evolution might explain the morphological differences between these structures. Given the similarities in the expression domains of the developmental regulatory genes *apterous* and *pdm*, downstream changes in target gene regulation may have contributed to the divergent morphologies of chelicerate respiratory structures, crustacean limb branches, spider spinnerets, and insect wings.

**Butterfly wing color scales: an evolutionary canvas**

The evolution of insect wings provided a new patterning surface that has been exploited in many ways by a variety of insect lineages. One of the most striking is the evolution of rows of pigmented scales that cover the wings of butterflies and moths (Lepidoptera). The shingle-like pigmented structural scales form the individual units of butterfly wing color patterns. Scales can exhibit a wide range of colors, shapes, and microarchitectures, both within and between butterfly species. Butterfly scales are also used for thermoregulation and predator avoidance. The array of thousands of overlapping scales on a butterfly wing surface represents a canvas on which a dazzling variety of colors and patterns has evolved.

Butterfly scales represent another morphological novelty that is derived from a preexisting structure—in this case, the innervated bristles that are common to all insects. The early cell lineage of butterfly wing scale development resembles the cell lineage that forms insect bristles (Fig. 6.3a). Each insect bristle consists of four distinct cell types: a bristle cell, a socket cell, a neuron, and a glial (sheath) cell. All four types originate from a single precursor, the sensory mother cell (SMC). The SMC divides to create the pIIa and pIIb cells; pIIa cells then divide again to create a bristle and a socket cell, and pIIb cells divide to create a neuron and a glial cell. By contrast, butterfly structural scales are made up of two cells, a scale cell and a socket cell, and are not innervated. The precursor cell for butterfly scales divides. While one daughter then divides again to create the socket and scale cells (similar to pIIa), the other daughter cell dies.
The evolutionary relationship between butterfly scales and insect bristles is reflected by regulatory gene expression within these cell lineages. The transcription factors that are expressed early in insect bristle development and that are required for bristle formation comprise the bHLH genes of the *Achaete-Scute* Complex (AS-C). A butterfly homolog of these genes (*ASH1*) is expressed both during butterfly sensory bristle formation and in the segregating scale precursor cells that are aligned in rows across the developing wing (Fig. 6.3b). The expression of *ASH1* during butterfly scale formation supports the hypothesis that early scale development is homologous to insect bristle formation.

At least three novel aspects of developmental gene regulation have generated the unique spatial distribution, cytoarchitecture, and pigmentation of butterfly wing scales. One involves the expanded expression of an AS-C gene in rows of cells across the entire surface of the developing butterfly wing. A second regulatory change is the evolution of a set of target genes deployed within scale cells that generates the flattened, rigid cytoarchitecture of butterfly wing scales. A third aspect of regulatory evolution is the recruitment of genes in the pigmentation pathway to be expressed in scale cells, thereby creating an array of colored
Chapter 6: The Evolution of Morphological Novelties

Evolution of the butterfly eyespot

The evolution of scale-covered lepidopteran wings provided a new landscape for displaying a striking variety of colors and shapes across the flat wing surface. These color patterns function to warn and confuse predators. For example, a flashing eyespot on a moving butterfly wing can startle a potential predator or redirect its attack away from the butterfly’s body. Butterfly eyespots, which are pattern elements composed of concentric rings of colored scales (Fig. 6.4), probably have a single evolutionary origin. Eyespot sizes and colors differ not only between butterfly species, but also between the forewings and hindwings of one animal, and even between the dorsal and ventral surfaces of a single wing. Despite the variation in the color, shape, number, and size of butterfly eyespots, the common features of eyespot development reveal key steps in the evolution of this novel pattern element—specifically, the co-option of genes and regulatory circuits for new patterning roles.

Eyespot development is controlled by a discrete organizer called the “focus,” a small group of epithelial cells at the center of the developing eyespot field. During butterfly pupation, focal cells induce the surrounding rings of scale-building cells to produce different pigments. Specification of the eyespot focus is indicated by the expression of the homeodomain-containing transcription factor *Distal-less* (*Dll*). A novel late pattern of *Dll* expression in focal cells follows an earlier conserved pattern of *Dll* expression (which is found in other insects...)

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Figure 6.4

Butterfly eyespot foci are marked by *Dll* expression

The adult hindwings of *Precis coenia* (a) and *Bicyclus anynana* (b) illustrate some of the variety seen in terms of number and coloration of eyespots between butterfly species. (c,d) *Dll* protein is expressed in larval stages along the wing edge in circular groups of cells of the presumptive eyespot foci. Two foci appear in *Precis* hindwings (c, arrows) and seven in *Bicyclus* hindwings (d).

as well) along the distal edge of the wing. The late pattern begins with the deployment of Dll in short rays of cells extending inward from the wing edge in each subdivision of the wing, then resolves to stable expression of Dll in circular fields of cells only in the subdivisions in which eyespot foci will form (Fig. 6.4). The continued expression of Dll in focal cells in specific wing subdivisions reflects the distribution of eyespots on adult butterfly wings. The conservation of Dll expression in the developing eyespot foci of different butterfly species suggests that the novel recruitment of Dll expression was an early event in the evolution of eyespots. Variation in eyespot size has been linked to the Dll locus, indicating the importance of this gene for eyespot evolution.

The evolution of butterfly eyespots entailed much more than the recruitment of Dll expression; indeed, an entire regulatory circuit was co-opted as well. The Hh signaling pathway, which has a conserved function in patterning along the A/P compartment boundary of insect wings (see Chapter 3), has evolved a second wing-patterning role during eyespot formation. Expression of genes in the Hh pathway (hedgehog (hh), patched (ptc), and Cubitus interruptus (Ci)) is modulated in unique patterns within or immediately surrounding eyespot foci (Fig. 6.5). These patterns suggest that this signaling pathway participates in the establishment of the eyespot focus. In addition, a known target gene of the Hh pathway at the wing A/P boundary—the late activation of engrailed (en) in some anterior compartment cells—is regulated by Hh signaling in the eyespot. En expression is upregulated within every eyespot

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**Figure 6.5**

Recruitment of the Hh signaling pathway in developing butterfly eyespots

The Hh signaling pathway plays a conserved role in patterning the A/P compartment boundary during insect wing development. In butterfly wings, this pathway is deployed in a novel pattern during the formation of eyespot foci. Co-option of Hh signaling requires changes in the regulation of members of the pathway including the hh and Ci genes (see text for details).

focus, even in foci that develop in the anterior compartment of the butterfly wing, where \textit{en} is not typically expressed (Fig. 6.5).

The recruitment of the Hh signaling pathway in the eyespot field is novel in two respects. First, the modulation of \textit{hh} expression in a novel pattern near developing eyespots is a unique feature of \textit{hh} expression in the butterfly wing. Second, the expression of \textit{ptc} and \textit{Ci} in the posterior compartment and of \textit{hh} in the anterior compartment of butterfly wings deviates from the compartmentally restricted deployment of these genes in the \textit{Drosophila} wing. In \textit{Drosophila}, expression of \textit{ptc} and \textit{Ci} is restricted to the anterior compartment because \textit{en} represses the expression of these genes in the posterior compartment. Consequently, \textit{en} repression of \textit{ptc} and \textit{Ci} must be relieved in butterfly wings so that the Hh pathway can be used in the posterior compartment. The combination of modulated \textit{hh} expression and release of the repression of \textit{ptc} and \textit{Ci} allows the Hh pathway to be activated in each developing eyespot field. The novel deployment of \textit{hh}, \textit{ptc}, \textit{Ci}, and \textit{en} in developing eyespots probably represents a critical subset of the regulatory changes that led to the co-option of a complete signal transduction circuit.

The diversity of color patterns within eyespots followed from regulatory changes in later stages of eyespot development. Early in butterfly pupal development, a signal from cells in the eyespot focus induces the pigmentation pattern in surrounding rings of scale cells. This inductive event is accompanied by rapid and dynamic changes in gene expression within the developing scale cells. Dll and En protein expression patterns arise again during this later stage in eyespot development, along with the regulatory protein Spalt (Sal). The expression patterns of these genes in eyespot scale-forming cells correlate with the future concentric color rings of the adult eyespot. Interestingly, the relative spatial domains of Dll, En, and Sal protein expression differ among species. No clear one-to-one correspondence exists between any of these proteins and a particular color output (such as an association between, for example, Sal and gold pigmentation in all eyespots). The differing relative spatial distributions of these proteins and eyespot color schemes reflect the remarkable evolutionary flexibility of the regulatory system governing eyespot development. It has been proposed that eyespots evolved from simpler spot patterns of uniform color. The expression of Dll, Sal, and En in the centers of eyespots appears to be a feature shared among eyespots and may offer a clue to eyespot origins. Namely, these proteins may have been expressed in simpler, uniform spot patterns. The evolution of multiringed spots may then have evolved the acquisition of signaling activity by cells expressing these genes and the recruitment of genes induced by this new signaling activity. The divergence of eyespot color schemes may have evolved through evolutionary changes in genetic responses downstream of signaling from the focus (Fig. 6.6).

The development and evolution of butterfly eyespots illustrate two recurring themes in the evolution of novel characters. First, conserved regulatory circuits can be recruited for new roles during the development of novel morphologies. This recruitment requires changes in the regulation of at least one gene, and subsequent deployment of other genes in the circuit may result from existing regulatory linkages, such as a signal transduction pathway. In this way, a large suite of genes may be deployed in a novel structure with just a small number of evolutionary regulatory changes. Second, evolutionary changes in target gene regulation can facilitate morphological diversification of a novel character. As regulatory evolution modifies the genetic interactions within a developmental program, new patterns can emerge both within and between species.
THE EVOLUTION OF VERTEBRATE NOVELTIES

The success and ecological dominance of the vertebrates can be attributed to several unique traits that arose during the evolution of vertebrates from a deuterostome ancestor. A limited list of novelties that evolved in the vertebrate lineage includes the following:

- The notochord
- Populations of cranial placodes and neural crest cells
- Jaws
- Paired pectoral and pelvic limbs

**Figure 6.6**

The evolution of eyespots from simple spots

It is thought that eyespots evolved from simpler spots of uniform color. Dll and En may have been expressed during the development of these primitive spots. As cells within this spot acquired signaling activity, surrounding cells acquired the ability to express different pigmentation genes. Further recruitment of signal-regulated genes led to the evolution of multiringed eyespots. Different responses to focal signaling among species may underlie different eyespot color schemes.
By comparing the morphology, ontogeny, developmental genetics, and fossil history of vertebrates and other chordates (such as the cephalochordate amphioxus and urochordate ascidians), it is possible to trace the evolutionary origins of these structures. Conserved regulatory genes control the pattern of these new developmental fields and tissues, revealing once again that new developmental programs have evolved through the co-option of existing regulatory genes and circuits and the expansion of ancestral patterning roles.

**Evolution of the notochord**

The presence of a notochord unites the chordates and is the distinguishing feature that inspired their name. The notochord is a stiff, axial rod of cells that represents the functional precursor of the vertebral column in basal chordates. It acts as an organizer for the early development of the CNS and adjacent axial mesoderm. Efforts to determine the origin of the notochord have often focused on the urochordates because of their relatively simple body plan and their basal position within the chordates. The larval ascidian tail consists of the notochord, muscle cells, and a dorsal nervous system. At metamorphosis, ascidian larvae lose their entire tail (including the notochord) and become sessile, filter-feeding adults.

The evolution of the notochord involved the co-option of an ancestral regulatory gene, *Brachyury* (*T*), a member of the T-box class of transcription factors (Fig. 6.7). In vertebrates, *T* is expressed in developing notochord cells and other mesodermal derivatives and is required for notochord differentiation. Similarly, the urochordate (ascidian and larvacean) *T* genes are expressed in cells that form the notochord and are sufficient to confer notochord fate. In larvaceans as well as hemichordates and invertebrates that lack a notochord,
homologs of the $T$ gene are expressed in posterior gut tissues. This expression may reflect the ancestral function of $T$ before the evolution of the notochord. This gene began to play a new role in notochord development after the chordate and hemichordate/echinoderm lineages became separated, as the hemichordate $T$ gene is not expressed in the stomochord (a possible precursor of the notochord).

*Sonic hedgehog* is also expressed in the notochord of cephalochordates and vertebrates and is required for maintenance of the notochord during development. By contrast, neither of the two ascidian *Hedgehog* genes are expressed in the notochord. In the ascidian *Ciona intestinalis*, *Hedgehog2* is expressed in the ventral portion of the nerve cord, similar to *Shh* expression in the floorplate of the neural tube in amphioxus and vertebrates. The expression of *Shh* in the mouse notochord and floorplate is controlled by distinct upstream regulators. Together, these data suggest that *Shh* was recruited for its role in the notochord after the divergence of the urochordate and cephalochordate/vertebrate lineages. Novel structures are likely to arise and become further elaborated through series of gene co-option events. Thus, changes in gene expression in the notochords of higher chordates may reflect new features of the notochord in the evolution of the chordate lineage.

**Placodes, neural crest, and jaws: rise of the predators**

One of the main vertebrate novelties was the elaboration of the vertebrate head, with an array of sense organs and features involved in an active predatory lifestyle. These structures are derived from two embryonic cell populations, cranial placodes and the neural crest, which arose near the base of the vertebrate lineage. Both of these pluripotent cell types undergo epithelial to mesenchymal transitions and have the capacity to migrate extensively within the body to form a variety of structures and cell types, including sensory neurons, the facial skeleton, connective tissue in the head and neck, peripheral neurons and glia, and melanocytes. The appearance of neural crest cells and cranial placodes and the resultant accumulation of special sense organs in the vertebrate head appears to have facilitated the evolution and radiation of the vertebrates. The subsequent invention of jaws enhanced the evolutionary success of the vertebrates, allowing a new mode of feeding.

Cranial placodes are discrete regions of thickened ectoderm that form in characteristic positions in the head of vertebrate embryos and contribute to specialized head structures. They make important contributions to the paired sense organs (nose, eyes, ears, and lateral line) and to cranial sensory ganglia, and are required for the formation of the sensory nervous system in the vertebrate head (Fig. 6.8a). Several lines of evidence suggest that at least some types of placodes are not unique to vertebrates but can also be found in basal chordates. Potential homologs of the olfactory placode have been identified in *Amphioxus* (corpuscles of Quatrefages) and in the colonial ascidian *Botryllus schlosseri* (neurohypophyseal duct). These specialized cell types form at the anterior end of the embryo, undergo morphogenetic movements, and express some of the same markers as vertebrate olfactory placodes. Another ascidian, *Halocynthia roretzi*, possesses structures (atrial primordia) that develop in a similar manner to, and express homologs of genes found in, the vertebrate otic placode. While placodes are apparently not unique to the vertebrates, they were, however, elaborated extensively during early vertebrate evolution and became indispensable for the formation of the vertebrate head.

The neural crest is derived from cells found at the interface of the lateral neural plate and epidermis along the dorsal side of the body (Fig. 6.8b,c). Basal vertebrate lineages such as hagfish and lampreys, as well as early fossil vertebrates, appear to possess most neural crest
Figure 6.8
Developmental origin of cranial placodes and neural crest

(a) In the chick embryo, cranial placodes (eight-somite stage fate map shown on right side) and neural crest contribute sensory neurons to the ganglia of cranial nerves (shown on left side). (b) Hox gene expression in the chick embryo at day 3 when the branchial arches (BA) are being colonized by neural crest cells originating from the posterior half of the midbrain (mesencephalon) and the rhombomeres (r1–r8). The arrows indicate the origin of the neural crest cells migrating to each BA. Expression of Hox genes is also indicated in the superficial ectoderm, the endoderm, and mesoderm. (c) Color-coded migration map of cephalic neural crest cells in the avian embryo. Posterior midbrain neural crest cells contribute to the anterodistal part of branchial arch 1 (BA1). The complementary portion of BA1 derives from r1/r2 together with a small contribution from r3. Other branchial arches (BA2–4) are populated by neural crest cells from more caudal rhombomeres.

derivatives, indicating that these cell populations arose early in the vertebrate lineage. Other chordates, such as amphioxus and ascidians, do not have homologous cell types. Nevertheless, comparisons of gene expression have suggested that members of the regulatory circuits that are required for neural crest induction in vertebrates are deployed in the neural tube in more basal chordates. Vertebrate neural crest cells express a suite of developmental regulatory genes, including transcription factors of the Dlx, Mnx, slug/snail, Pax2/5/8, and Pax3/7 families. Homologs of these genes are expressed in comparable positions in both amphioxus and ascidians. Thus, the spatial expression domains of these genes appear to have been established before neural crest cells evolved at the base of the vertebrate lineage. Conspicuously, other markers of neural crest cells are not expressed at the lateral edge of the neural plate in amphioxus, but can be found in neural crest cells in lampreys. The elaboration of the neural crest in the vertebrate lineage, with its diverse neuronal and non-neuronal cell types, may have its roots in a small ancestral population of cells in or near the chordate lateral neural plate. The changes must have occurred “rapidly” at the base of the vertebrate lineage.

The earliest vertebrates (agnathans) lacked jaws but jawed vertebrates (gnathostomes) appeared ~400 million years ago and subsequently radiated to occupy varied terrestrial and aquatic niches. During gnathostome development, both the upper and lower jaws are formed from the first set of branchial/pharyngeal arches, paired structures located ventrally in the cranial region of the embryo. The upper jaw will develop from the proximal portion of the arch while the lower jaw develops from the distal region. Neural crest cells from the midbrain and hindbrain migrate into the branchial arches (Fig. 6.8b,c) and differentiate into (among other things) the cartilage and bone of the craniofacial skeleton, including the jaws. In lampreys (agnathans), the early development of the neural crest and arches, including gene expression patterns, is similar to that of gnathostomes, except for the presence of lamprey Hox gene expression in the first arch. In vertebrates, Hox gene expression in the first arch represses jaw formation. The absence of Hox expression in the first arch of vertebrates may have allowed the formation of extra cartilage, which acted as a substrate for jaw formation. While shifts in Hox gene expression may have been permissive for jaw development and evolution, other patterning genes were recruited for the subsequent sculpting of jaw architecture. In mammals, the Dlx genes are expressed in nested domains along the proximodistal axis of the arches (Fig. 6.9a). Dlx3/7 are expressed at the distal tips of the arches while Dlx5/6 are expressed in a broad distal domain and Dlx1/2 are expressed and required throughout the arches. By contrast, lampreys do not show restricted expression of any Dlx genes along the proximodistal axes of the arches. The combinatorial expression of the Dlx genes acts to specify different domains along the proximodistal axis of the arches and regulates the identity of jaw elements (Fig. 6.9b). This regionalization of the first arch was likely a crucial step in elaboration of the upper and lower jaws in the gnathostome lineage.

The increased complexity of vertebrate cranial development, including the elaboration of placodes, the appearance of a neural crest cell population, and formation of jaws may be correlated with expansion of the vertebrate genome through large-scale duplication events (see Chapter 4). The additional developmental regulatory genes that were created by gene duplication evolved novel roles in controlling the differentiation of new cell types and the more complex organization of the vertebrate central and peripheral nervous systems. In vertebrates, gene duplication and divergence appears to have provided the potential for more complex morphologies.
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Fins to limbs: paired appendages and the tetrapod band

The adaptive evolution of vertebrates capable of surviving in aquatic, terrestrial, and aerial environments involved the acquisition and modification of paired pectoral and pelvic appendages. These limbs boosted vertebrates’ maneuverability and speed in water and later were used as the primary means of locomotion on land. The early history of paired
appendages remains uncertain, however. Diverse fin morphologies are present in Ordovician fish (463–439 Ma). Some of these ancient fish possessed median fins running the length of the body, some had paired fins near the head, and others had no fins at all. Based on the fossil record, median fins appear to have preceded the appearance of paired fins, and paired pectoral fins preceded the evolution of pelvic fins (Fig. 6.10).

The group of bony fishes that later gave rise to the tetrapod lineage included animals with similarly patterned pectoral and pelvic fins. This similarity may reflect the deployment of the same regulatory genes in each limb pair. Modern tetrapods and bony fish exhibit similar patterns of Hox gene expression during the development of pelvic and pectoral appendages. The posterior genes of both the HoxA and HoxD complexes (Hox9–13) are expressed in complex and dynamic patterns during tetrapod limb development (see Chapter 3). In unpaired fish fins, however, these Hox genes are not deployed. Consequently, primitive vertebrate appendages may not have been patterned by Hox genes, and Hox genes may have been recruited later to pattern paired fins.

Figure 6.10

Evolution of vertebrate paired appendages

Vertebrate paired appendages evolved in a series of steps. Paired appendages are first encountered in the fossil record as elongate fins extending laterally along the body wall or as paired pectoral fins in jawless fish (shown on basal branches). Jawed fish are characterized by multiple sets of paired appendages that often have a defined fin axis (shown in blue; top left). Tetrapods (top right) have a distal autopod, with digits branching from a bend in the limb axis. One step in the evolution of serially homologous paired appendages was the co-option of nested patterns of posterior Hox genes expression in both sets of paired appendages (lower right).

The evolution of posterior HoxA and HoxD gene expression in both pectoral and pelvic limbs may have occurred in two discrete steps (Fig. 6.10). First, axial patterns of posterior Hox gene expression may have been recruited during the development of the adjacent paired pelvic limbs. Later, the pectoral limbs may have co-opted the pelvic pattern of Hox deployment. The nested domains of Hox gene expression found along the primary body axis and within developing limbs suggest that the ancestral colinearity of Hox expression was maintained as these genes assumed novel roles in limb development.

Another novel feature of posterior Hox gene regulation is evident in the distal element of tetrapod limbs. The autopod, which consists of the hand (or foot) and digits, is unique to tetrapods. During autopod development, a late phase of Hox gene expression (phase III, see Chapter 3) appears that is not present in teleost fish (Fig. 6.11). This phase is characterized

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**Figure 6.11**

**The origin of tetrapod digits**

The evolution of the autopod is characterized by branching of digits from a bend in the limb axis and by the appearance of a third phase of Hox gene expression in the distal limb. A fin axis is present in the paired fins of bony fish (gray; first four fins on **left**). The distal tetrapod limb has a 90° bend in the equivalent limb axis (gray; **top right**). **Box** Proximal elements of the vertebrate limb branch to the posterior from the limb axis, whereas elements of the autopod branch to the anterior (red). The reversal in branching direction is also reflected in a reversal in the nested domains of phase III Hox gene expression (light blue, Hoxd11; dark blue, Hoxd13).

by a reversal in the anteroposterior order of Hox gene expression in the autopod compared with the earlier, more proximal expression domains in other developing limb elements. A single cis-regulatory element drives phase III expression of all of the posterior HoxD genes, indicating that several HoxD genes in the autopod may have been recruited simultaneously through the evolution of this element.

**EVOLUTION OF RADICAL BODY PLAN CHANGES**

Morphological novelty also encompasses the evolution of novel body organizations. Radical reorganization of body plans can include both the appearance of new structures and the loss of ancestral characters. The first two examples we discuss here have lost one or more key morphological features characteristic of their clade. The other examples illustrate the appearance of new characters and rearrangements of older features resulting in dramatically different body organization. As with the other types of novelties discussed in this chapter, these morphological changes are correlated with changes in genetic regulatory circuits.

**Loss of the ascidian tail, a chordate lacking the notochord**

Although the notochord is a defining feature of the chordates, several ascidian species have an abbreviated larval stage that has lost the notochord along with the entire larval tail. Of the more than 3000 ascidian species known, about a dozen are tailless. Loss or reduction of the larval tail has occurred at least six independent times in the ascidian lineage. Taillessness is associated with direct development, including a rapid progression to the sessile adult form.

In some cases, very closely related species differ dramatically in the extent of their tail development. For example, the urodele (tailed) larvae of *Molgula oculata* develop a notochord, a spinal cord, and striated muscle cells. By contrast, another *Molgula* species (*M. occulta*) has an anural (tailless) larval stage that lacks these characteristic chordate features. When these species are interbred in the laboratory, the crosses yield a short-tailed hybrid, complete with notochord (Fig. 6.12). Thus the tailless phenotype of *M. occulta* can be rescued by the *M. oculata* genome, suggesting that *M. occulta* may have lost some genetic function that results in its tailless larval form.

Changes in the tail development program that lead to an anural form occur downstream of *Brachyury* expression in the notochord. Notochord cells are specified normally in *M. oculata* and *M. tectiformis* (another tailless species) but these cells do not divide properly or undergo appropriate morphogenetic movements to form the notochord. The search for genes that are essential for *M. oculata* tail development, but that are downregulated in *M. occulta*, has led to the identification of the *Manx*, *Bobcat*, and *FoxA5* genes. The *Manx* gene is a zinc-finger transcription factor, *Bobcat* encodes an RNA helicase, and *FoxA5* encodes a winged helix transcription factor. The expression of these genes is largely restricted to the embryonic cells that will give rise to the *M. oculata* larval tail (including the notochord). These genes are similarly deployed in *M. oculata × M. occulta* hybrids, and inhibition of any of these genes results in an increase in programmed cell death in notochord and tail muscle cells that generates a tailless embryo. The requirement for *Manx*, *Bobcat*, and *FoxA5* gene function indicates that these genes are part of the developmental program for tail formation.
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The reduced expression of the Manx, Bobcat, and FoxA5 genes in the tailless species *M. occulta* suggests that the tail regulatory hierarchy is terminated early in development. During the evolution of this tailless form, the expression or function of an upstream regulator common to these genes may have been changed that led to the loss of expression of these genes. Interestingly, disruption of tail development seems to have relieved the selective pressure on downstream genes necessary for tail function. Several muscle actin genes, which in urodele ascidians are expressed in tail muscle cells, have become pseudogenes in several independent anuran ascidian lineages. The altered expression of a single regulatory gene required for tail development may be sufficient to evolve a novel larval form and to release selective constraint on the network of dependent genes.

The reduced expression of the *Manx*, *Bobcat*, and *FoxA5* genes in the tailless species *M. occulta* suggests that the tail regulatory hierarchy is terminated early in development. During the evolution of this tailless form, the expression or function of an upstream regulator common to these genes may have been changed that led to the loss of expression of these genes. Interestingly, disruption of tail development seems to have relieved the selective pressure on downstream genes necessary for tail function. Several muscle actin genes, which in urodele ascidians are expressed in tail muscle cells, have become pseudogenes in several independent anuran ascidian lineages. The altered expression of a single regulatory gene required for tail development may be sufficient to evolve a novel larval form and to release selective constraint on the network of dependent genes.

**Limbless tetrapods, or how the snake lost its legs**

The evolution of snakes, with their characteristic elongated bodies and reduced limbs, represents an adaptation of the tetrapod body plan to a novel form. Body elongation and limblessness are associated with burrowing and have evolved independently many times within the reptiles, including once at the base of the snake lineage. Among modern snakes, the python lineage retains some features of the tetrapod hindlimb, including a pelvic girdle and rudimentary hindlimbs, whereas more highly derived snakes are completely limbless. Comparisons between snakes and other tetrapods have revealed key developmental genetic
differences in axial patterning and limb development that correlate with the evolutionary transition to the limbless snake body plan.

The concomitant loss of snake forelimbs and expansion of thoracic axial identities suggest that a common genetic mechanism may underlie these evolutionary transitions. In Chapter 5, we discussed the correlation between the python axial skeleton identities and the anterior expansion of the expression domains of \( Hoxc6 \) and \( Hoxc8 \). In other tetrapods, the axial position of the forelimb is determined by the anterior boundary of \( Hoxc6 \) expression in lateral plate mesoderm at the cervical–thoracic transition (see Fig. 5.6). Thus the extension of the expression pattern of python \( Hoxc6 \) and \( Hoxc8 \) up to the cranial region may disrupt the \( Hox \)-regulated positioning of the forelimb. Without the proper axial patterning cues, this limb bud never forms.

The correlation between trunk elongation and limb loss in several other vertebrate taxa may be explained by similar changes in axial patterning caused by altered \( Hox \) expression patterns.

The fossil record and developmental studies indicate the reduction and loss of the snake hindlimb occurred via a separate series of evolutionary events. The initial development of the vestigial python hindlimb appears normal, indicating that the axial specification cues and early induction of python hindlimb development remain intact. As in other tetrapods, the posterior boundaries of \( Hoxc6 \) and \( Hoxc8 \) expression mark the python thoracic–lumbar transition at the position of the rudimentary pelvic girdle, and a limb bud does form. Ultimately, however, development of the hindlimb is arrested and no outgrowth is observed.

The python hindlimb bud lacks some of the characteristic features of other developing tetrapod limbs, such as an apical ectodermal ridge (AER) and a zone of polarizing activity (ZPA). Nevertheless, experimental manipulations have shown that the python limb bud mesenchyme retains the capacity to induce an AER and a ZPA; indeed, it can form a complete limb in response to an inductive signal from an exogenous AER. Thus much of the developmental program of limb formation is present in the python limb bud, but at least one essential element must be missing. The termination of python hindlimb development suggests that this field fails to initiate or maintain the signaling events that are required for AER formation and hindlimb outgrowth.

The loss of a structure is not necessarily accompanied by the loss of the genes—or even genetic regulatory circuitry—required to build that structure. The condition of limblessness can be reversed, as evidenced both by rare individuals in wild populations and by stable evolutionary lineages. For example, rare natural occurrences of cetaceans (whales and dolphins) with well-developed hindlimbs have been reported. Also, the snake fossil record shows that a limbless ancestor may have given rise to a group of snakes that had substantial limbs (Fig. 6.13). For example, the fairly well-developed hindlimbs of the snake fossils \( Pachyrhachis problematicus \) and \( Hassiophis terrasanctus \) may represent a reversal from a limbless state. Based on these cases of reversion to the ancestral limbed condition, it appears that the limb developmental program may still be present in limbless animals. As the genes required for limb patterning have other developmental functions, they are retained in the genomes of limbless taxa. More surprising, however, is that an otherwise cryptic regulatory circuitry required for limb development is retained in a limbless tetrapod.

### The evolution of the turtle shell

The turtle shell is the novel structure that defines the order Chelonia. The two main components of the shell are the dorsal carapace and the ventral plastron (Fig. 6.14a,b), which are linked
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The evolution of this bony casing encompassed a large number of modifications to the tetrapod body plan including rearrangements of the pectoral girdle, ribcage, vertebrae, neck, and skull. While dermal ossification is a primitive character for vertebrates as evidenced by numerous fossil fish specimens that display extensive exoskeletons, the turtle shell represents an extreme elaboration of the dermal skeleton among tetrapods.

The ventral plastron and dorsal carapace arise by different mechanisms during turtle development. Cells in the plastron express molecular markers of bone-forming neural crest cells, suggesting that the turtle plastron is derived from neural crest cells. However, in extant tetrapods trunk neural crest cells generally do not contribute to skeletogenesis. The turtle may have retained this ability or recruited the head neural crest cells to elaborate the ventral portion of this unique structure.

While formation of the ventral plastron in turtles might reflect the retention or redeployment of dermal armor found in early gnathostomes, the dorsal carapace arises from a novel structure, the carapacial ridge (CR), a bulge of ectoderm and mesoderm that arises on the dorsal flank of the turtle embryo. During turtle development, the ribs become trapped by the CR, enclosed within the dorsal dermis, and displaced laterally (Fig. 6.14d), later fusing with the dermis of the carapace (Fig. 6.14c). The tissue composition of the carapacial ridge is similar to that of the early vertebrate limb bud. Perhaps not surprisingly, the Fgf10 and Msx genes are expressed in the distal mesenchyme of the CR, similar to their expression patterns

Figure 6.13
Evolutionary transitions in snake limblessness

The phylogeny of snakes indicates that the loss of snake hindlimbs may have evolved once at the base of the snake lineage. The existence of well-developed hindlimbs in the fossils Pachyrachis and Haasiophis suggests that hindlimbs reappeared during the evolution of these species, as their ancestor may have been limbless. Alternatively, snake hindlimbs could have been independently lost in each limbless snake lineage.

in the growing limb bud underlying the AER. The expression of these genes in the CR, coupled with the similarities in morphology and development between the limb AER and the turtle shell CR, suggests that the same mechanisms may initiate the formation of these two structures. The CR does not express Fgf8 however, a crucial molecule required for limb outgrowth. The
CR may have arisen during evolution as the result of co-option of part of the limb initiation pathway.

**Evolution of the cephalopod body plan**

The molluscs appear in the fossil record of the early Cambrian about 530 million years ago. Extant molluscan species display a wide range of body plans but the typical mollusc has three defining features: a shell, a mantle, and a ventral locomotory foot (see Fig. 6.15a). Despite a close affinity to gastropod molluscs (snails, slugs, etc.), cephalopod molluscs (squids, nautiluses, octopuses, cuttlefish) have evolved striking modifications to these features. The cephalopod mantle has evolved new roles in respiration and locomotion while the ventral foot has been modified to form the arms of the brachial crown (tentacles) and funnel tube, used for locomotion. In addition, except for *Nautilus*, the cephalopod shell has been greatly reduced or lost (Fig. 6.15a,b).

Cephalopod squids (*Euprymna scalopa*) possess a complement of *Hox* genes typical of Lophotrochozoans: one anterior, one paralog group 3, five central, and two posterior group *Hox* genes. Six of these *Hox* genes are expressed in the central nervous system and while their expression patterns do not violate the principle of colinearity, they also do not display a canonical nested pattern. Strikingly, the *Hox* genes are expressed in morphological novelties unique to the cephalopods such as the arms of the brachial crown and the funnel tube. However, the anteroposterior expression domains of the *Hox* genes in these structures do not agree with *Hox* colinearity. Different subsets of *Hox* gene are expressed in nested domains within the funnel tube, while multiple *Hox* genes are expressed in dynamic combinations in the arms of the brachial crown (Fig. 6.15c). Other novel structures show the recruitment of individual *Hox* genes.

These radical changes in *Hox* gene expression domains suggest that the evolution of cephalopod morphological novelties was accompanied by, or even precipitated by, the deployment of *Hox* genes in unique patterns defining new structures. While collinear expression of *Hox* genes is evident in other molluscs, the loss of collinear regulation in cephalopods may be tied to the use of *Hox* genes to regulate developmental events other than anteroposterior patterning.

**Evolution of the echinoderm body plan**

The defining features of the Bilateria, including the evolution of bilateral symmetry with distinct rostrocaudal and D/V axes, arose before the radiation of most animal phyla. Although the echinoderms (sea stars, sea urchins, sea cucumbers, brittle stars, and crinoids) are a basal deuterostome lineage derived from a bilaterian ancestor, echinoderm adults have a radially symmetric body organization. Whereas echinoderm larvae exhibit a more typical bilateral form, adult echinoderms are radically rearranged, making it difficult to identify any vestiges of bilateral symmetry (Fig. 6.16). The fivefold symmetry of the adult nervous system, connected by a central nerve ring, has led to the proposal that each echinoderm “arm” represents a duplication of the bilaterian A/P axis. Alternatively, the distribution of mesodermal tissue in the adult may indicate that the orthogonal adult **oral-aboral axis** is derived from the larval A/P axis. The origin of the unique and highly derived body plan of adult echinoderms remains a puzzle.
The development of other structural novelties of echinoderms, including the water vascular system and the calcitic endoskeleton, appears to involve regulatory genes shared with other phyla. These genes have been recruited for deployment in echinoderm-specific tissues in the developing juvenile sea urchin. *Runt*, *Dll*, and *Otx* (in direct developers) are expressed in the tube feet (podia), which are extensions of the water vascular system. *Hox3* is expressed in the nascent tooth sacs, which give rise to the echinoderm mouthparts. The pattern of *en* expression in the ectoderm reflects the organization of the underlying endoskeleton. In short, these regulatory genes appear to have been co-opted for new patterning roles associated with the evolution of novel echinoderm structural elements.
Chapter 6: The Evolution of Morphological Novelties

Figure 6.16
Evolution of the radially symmetric echinoderm body plan
(a) Echinoderm lineages have bilaterally symmetric larvae and radially symmetric adults. (b) Two hypotheses regarding the orientation of the ancestral bilaterian A/P axis in the adult echinoderm body plan: (left) multiplication of the bilaterian A/P axis may have created the radial nerve cords that extend down each echinoderm arm; (right) the ancestral bilaterian A/P axis may have evolved into the echinoderm oral–aboral axis.

REGULATORY EVOLUTION AND THE ORIGIN OF NOVELTIES

The recurring theme among the diverse examples of evolutionary novelties described in this chapter is the creative role played by evolutionary changes in gene regulation. The evolution of new regulatory linkages—between signaling pathways and target genes, transcriptional
regulators and structural genes, and so on—has created new regulatory circuits that have shaped the development of myriad functionally important structures. These regulatory circuits also serve as the foundation of further diversification. The patterning of structures such as the tetrapod autopod and butterfly wing has diverged extensively throughout the radiation of the lineages in which these novelties first evolved.

The time scale over which new structures have evolved or body plans have diverged is considerable—of the order of many millions of years. One of the best-analyzed transitions, the evolution of tetrapod limbs from fish fins, is estimated to have transpired over the course of about 9 million years, for example. As we learn more about the architecture of the genetic regulatory differences between lineages, as well as the scope and time scale of anatomical evolution, we can better appreciate that changes in a potentially very large number of regulatory linkages accompany morphological evolution. Chapter 7 will examine the beginnings of that process—the origins of variations in morphological characters and the divergence of closely related species.

**SELECTED READINGS**

**General Reviews**


**Origin and Evolution of Feathers**


Prum RO, Brush AH. Which came first, the feather or the bird? *Sci Am* 2003; 288: 84–93.


Origin of the Insect Wing


Butterfly Scales and Eyespots


Evolution of the Notochord


Jeong Y, Epstein DJ. Distinct regulators of Shh transcription in the floor plate and notochord indicate separate origins for these tissues in the mouse node. *Development* 2003; 130: 3891–3902.


**Evolution of Cranial Placodes, Neural Crest, and Jaws**


Vertebrate Limbs and Digits


Loss of the Notochord in Ascidians

Swalla BJ, Just MA, Pederson EL, Jeffery WR. A multigene locus containing the *Manx* and *bobcat* genes is required for development of chordate features in the ascidian tadpole larva. *Development* 1999; 126: 1643–1653.

Loss of Limbs in Snakes


Origin of the Turtle Shell

DNA to Diversity: Molecular Genetics and the Evolution of Animal Design


The Cephalopod Body Plan


The Echinoderm Body Plan


CHAPTER 7

Morphological Variation and Species Divergence

The assembly and expansion of the genetic toolkit in early metazoans and its expansion in early vertebrates were two intervals in which gene duplication and divergence played major roles in morphological evolution. However, the diversification of protostomes, most deuterostomes, and the vertebrates (after genome expansion) occurred around ancient and largely equivalent toolkits of major developmental genes. These observations lead us to conclude that changes in developmental gene regulation are the predominant genetic mechanisms underlying large-scale morphological evolution. Most of the examples in Chapters 5 and 6, however, have focused on differences between groups at higher taxonomic levels, where gross differences in the expression of major regulatory genes are correlated with morphological diversification and innovation. One implicit assumption of the case studies examined thus far is that large-scale diversity is a product of the same evolutionary and developmental mechanisms that operate on the scale of individual species. To test the validity of this assumption, we must change the taxonomic scale of our analysis and the nature of the characters under study. Because the source of morphological evolution is morphological variation, it is important to understand the genetic architecture and the molecular basis of morphological variation within species and of phenotypic divergence between closely related species.

In order to approach phenotypic and developmental evolution at the level of individual species, analyses of more variable and rapidly evolving traits are necessary. Characters such as coat color in mammals, pigmentation in insects, bristle numbers in flies, and skeletal armor in certain fishes have diverged frequently and rapidly. Because genetic studies are sometimes feasible between closely related species, dissection of the underlying genetic, developmental, and molecular bases of the divergence of some characters is possible at a much deeper mechanistic level than are slowly evolving traits among more divergent groups. Studies of...
closely related or morphologically variable species have sought to address four general questions concerning the genetic and molecular mechanisms of morphological evolution, including:

1. How many and which specific genes underlie phenotypic divergence of a particular trait between species?
2. Are the same genes involved in intraspecific variation and interspecific divergence of a specific trait?
3. Are the same loci responsible for similar, independent phenotypic changes (convergence) in different lineages?
4. What is the molecular nature of genetic differences contributing to divergence (i.e. the relative contribution of regulatory and coding changes)?

This chapter uses case studies to examine the genetic architecture and molecular mechanisms underlying differences between related species or populations. The characters studied range from simple qualitative traits affected by single genes, to quantitative traits that may be influenced by variation at just a few to more than 20 loci. The architecture of genetic regulatory systems is a major determinant of the means through which evolutionary differences arise. In several cases, independent evolutionary changes in different lineages have occurred through evolution at the same genetic loci, providing striking molecular evidence for convergence. Both phenotypic divergence and variation are most often associated with noncoding, regulatory sequences, suggesting again that regulatory evolution in developmental genes is the predominant force in morphological evolution.

**EVOLUTION OF ANIMAL COLOR PATTERNS**

*Mammalian coat and bird plumage colors*

Some of the most striking and best understood examples of phenotypic divergence are the color patterns of vertebrates. Mammalian coat, bird plumage, and fish scale coloration schemes are wonderfully diverse. Much progress has been made in understanding the genetic control of color formation and of differences within and between species.

The most widespread pigment in the animal kingdom is melanin. It occurs in various chemical forms that when polymerized produce black, brown, buff, tan or even reddish pigments. Melanism, the occurrence of dark morphs within a population or species, is one of the most common forms of phenotypic variation. In mammals, two types of melanin are produced in melanocytes (the pigment cells of the epidermis and hair follicles), eumelanin and phaeomelanin, which produce black/brown and red/yellow coloration respectively. The relative amounts of eumelanin and phaeomelanin are controlled by the products of several genes. Two key proteins are the melanocortin 1 receptor (MC1R) and the Agouti protein. During the hair growth cycle, α-melanocyte stimulating hormone (α-MSH) binds to the MC1R, which triggers elevated cAMP levels and activation of tyrosinase, the rate-limiting enzyme in melanin synthesis. The Agouti protein acts as an antagonist of this process by blocking α-MSH action, inhibiting eumelanin production, and allowing the default pathway
Chapter 7: Morphological Variation and Species Divergence

of phaeomelanin synthesis. The true wild-type color of mouse fur arises from a banded coloration of each hair which is mostly black but with a subapical zone of yellow pigment. This zone is produced by a pulse of Agouti protein synthesis during a phase of the hair growth cycle. In the absence of Agouti function the hair is all black, while dominant gain of function agouti alleles produces a yellow coat. Mutations in the MC1R and agouti genes are associated with natural morphs of a variety of species and domestic breeds.

One striking example of melanism is found in the jaguar (*Panthera onca*) of Central and South America, which occurs in both the classic orange and a melanic phase (Fig. 7.1). Sequencing of melanic jaguar MC1R alleles revealed an in-frame five codon deletion with an adjacent amino acid replacement. All melanistic jaguars were found to be either homozygous or heterozygous for the deletion allele, while all orange animals were homozygous for the wild-type allele. Breeding studies have found that melanism in this cat is dominant, the melanistic allele appears to cause a constitutive activation of the MC1R protein. Similarly, melanism in the jaguarundi (*Herpailurus yaguarondi*) shows a dominant or semidominant
pattern of inheritance and is associated with an independent inframe deletion of *MC1R* that partially overlaps with the melanism-associated deletion in the jaguar. These findings demonstrate that dramatic changes in coat color can arise from mutations in a single gene, and be inherited and expressed in a dominant fashion in wild populations.

The factors that favor the spread of melanic forms of these cats are not clear. There is compelling field data to suggest that melanism involving the MC1R gene underlies adaptation in other species. For example, rock pocket mice are generally light-colored and live on light-colored rocks. However, in the American Southwest, populations of melanic mice are found that live on dark lava formations (Fig. 7.2) and field studies suggest that the dark coloration provides protection from predators such as owls. In an Arizona population, four amino acid replacements were found in the MC1R gene that showed a perfect association with dark coloration. Furthermore, a pattern of reduced nucleotide variation in the MC1R alleles of the

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**Figure 7.2**

**Melanism in the rock pocket mouse**

Light and dark *Chaetodipus intermedius* from the Pinacate region of Arizona are shown on light and dark rock backgrounds.

Clear associations between MC1R mutations and melanism have also been found in other species. For example, the bananaquit is a widely distributed bird of the Neotropics. Most individuals have a bright yellow breast plumage and a white eye stripe but on the islands of Grenada and St Vincent and a few other locales, melanic forms occur that are almost completely black. A point mutation in MC1R that changes a single amino acid is always associated with melanism. This mutation causes the identical amino acid replacement associated with melanism in domestic chickens and mice. Thus, the same MC1R mutation has occurred independently in multiple lineages, and other mutations in MC1R have occurred at different sites, all of which cause constitutive activation of the melanocortin receptor (Fig. 7.3). The association of many cases of melanism in various mammals and birds with the same gene demonstrates that, at least in the case of coloration, evolution can and does repeat itself.

There are examples of melanism caused by mutations in genes other than MC1R. In the domestic cat, a small deletion in the \textit{agouti} gene is responsible for a homozygous recessive form of melanism. Since Agouti acts as an inhibitor of melanism, homozygous loss of function mutations lead to melanism. Mutations in \textit{agouti} are also associated with black coat color in nine different horse breeds.

There are other, as yet unknown, evolutionary pathways to melanism in vertebrates. In a number of cat species, including the leopard, no association exists between \textit{MC1R} or \textit{agouti} and melanism. Perhaps most surprisingly, in a second population of melanic rock pocket mice in New Mexico, no association was found between melanism and \textit{MC1R}. These cases demonstrate that while similar phenotypic changes in different populations can be due to changes in the same gene, phenotypic convergence can also occur via independent genetic mechanisms. In the case of the rock pocket mice, the lava flows on which the two populations live are less than one million years old, indicating that convergence has occurred by different mechanisms in a relatively short timeframe.

Variation at the \textit{MC1R} locus is also associated with other coat color phenotypes in addition to melanism. In the Pacific Northwest, two populations of the black bear occur. In addition to the more prevalent black morph, there is a striking white-phase “Kermode” bear (Fig. 7.4). Once thought to be separate species, it is now understood that the white-phased bear carries a single amino acid replacement in MC1R (Fig. 7.3). In this case, the black allele is dominant and the white allele is recessive, suggesting that white is due to a loss of MC1R function. Interestingly, the replacement in the Kermode bear MC1R protein occurs very near to a terminal deletion in MC1R that is associated in dogs with the red or yellow coat colors of yellow labradors, golden retrievers, and Irish setters, also thought to be due to a loss of MC1R function (Fig. 7.3). The different specific coat colors among these dog breeds are due to additional modifier genes that have not yet been identified.

It is important to note that all of these examples of coat color differences that are brought about by mutations in \textit{MC1R} or \textit{agouti} affect the entire coat. Of course, many coat colors are patterned in mammals, in stripes, spots, bands, etc. The molecular genetic basis of these patterns is generally not understood so it is not known whether coding changes in pigmentation pathway components, or differences in their spatial regulation, underlie the phenotypic divergence of more complex patterns. There is some suggestion from the mouse \textit{agouti} gene as to how more complex coat patterns may be determined. One of the most common coat color schemes in rodents and other mammals is the lighter coloration of the underside. The
Figure 7.3
Location of mutations in the MC1R protein associated with melanism and other coat color forms

Partial diagram of the MC1R protein shows the location of amino acid replacements (orange colored circles) or deletions (dotted lines) associated with melanism in species indicated. The inset at left shows the general topology of the MC1R protein, with mutations nearer the amino terminus associated with melanism, and those nearer the carboxy terminus associated with white (Kermode Bear) or orange, yellow, or red coats (in various dogs).

agouti gene contains two different promoters, one which is active only in the dermal papilla of hair follicles on the ventral side of the animal, and one which is active in the midpoint of the hair cycle all over the body. The presence of this ventral-specific element suggests that it is likely that differences in the regulation of *agouti* in different body regions and other pigmentation pathway components could evolve through changes in *cis*-regulatory elements and contribute to coat color pattern diversity. The genetic architecture of the evolution of spatial coat color patterns is likely to be complex and, other than in the mouse, would be difficult to examine experimentally in most mammals. However, many groups of insects also display quite diverse color patterns, and the genetic basis of variation and divergence is beginning to be understood in model groups.

**Pigmentation pattern diversity in Drosophila**

Insect pigmentation offers some particular advantages as models for understanding the genetic, developmental, and molecular basis of character evolution. Foremost among these
is the wealth of genetic and biochemical knowledge of pigmentation pattern development in *Drosophila melanogaster*. The fruit fly subfamily Drosophilinae to which *D. melanogaster* belongs displays a vast spectrum of pigmentation patterns, usually a mixture of light and dark pigments distributed in discrete spatial patterns on the thorax, abdomen, and sometimes, the wings. The cell biology of pigmentation is entirely different in insects and vertebrates, but some of the pigment chemistry is very similar. Black coloration is a polymer of dopamelanin and brown is a polymer of dopamine melanin. Other hues are created by chemical modifications of dopa and dopamine precursors. Pigmentation patterns are determined by a variety of transcription factors that control the expression of enzymatic components of the pigmentation pathway. A number of studies have identified individual genes associated with pigmentation variation and divergence among various groups of *Drosophila* species. Regulatory changes in both structural and regulatory genes contribute to pigmentation divergence, and in some cases, the regulation of the same gene has been modified many times independently in different lineages to produce divergent or similar patterns.

**Evolution through a modular cis-regulatory element of a pigmentation gene**

The *yellow* and *ebony* genes encode proteins that promote and inhibit black melanin formation, respectively, in *D. melanogaster* (the genes’ names reflect the phenotypes of mutants). In many groups of related species, the degree of pigmentation of body parts differs considerably, and differences in Yellow and Ebony protein expression are associated with the pattern and/or degree of pigmentation among particular taxa. For example, in *D. subobscura* and *D. virilis*, dark pigmentation is more broadly distributed on the abdomen than in *D. melanogaster*. The level and pattern of expression of the Yellow protein correlates with dark pigmentation (Fig. 7.5).

The expression of *yellow* in various body parts (wing, body, bristles, larval mouthparts) is controlled by different cis-regulatory elements. Interspecific differences in Yellow expression are due, at least in part, to the evolution of a cis-regulatory element of the *yellow* gene. The function of the “body” enhancer of the *yellow* locus has diverged between *D. melanogaster*, *D. subobscura*, and *D. virilis* such that higher levels and a broader pattern of Yellow are produced in the latter two, more darkly pigmented species. This can be demonstrated using transgenic techniques by introducing the respective *yellow* genes or regulatory elements from different species into *D. melanogaster*. Because the cis-regulatory elements are independent modules, the pattern, level, and timing of *yellow* gene expression and pigmentation can evolve in selected body regions independently of other body parts.

Darker body pigmentation is not always associated with the *Yellow* protein. In two closely related species, the lightly colored *D. novomexicana* and its darkly colored sister species *D. americana*, the darker coloration is not associated with evolution at the *yellow* gene, but rather at the *ebony* gene. The darker *D. americana* expresses lower levels of the Ebony protein which allows for darker pigmentation. These two species can be crossed and hybrids backcrossed. Genetic association studies indicate that several other unidentified loci also contribute to the difference in pigmentation. Even though pigmentation is a relatively less complex character than the morphology of body parts, and single genes can have large effects, multiple loci are probably involved in the evolution of pigmentation differences between species.
Chapter 7: Morphological Variation and Species Divergence

Evolution at a regulatory locus associated with diverse sex-specific and segment-specific pigmentation patterns

Most members of the melanogaster species group exhibit sexually dimorphic pigmentation where the last two abdominal segments are heavily pigmented in males but not in females (Fig. 7.6, top). This sex-specific and segment-specific pattern is a fairly recent innovation, as the abdomens of both sexes are identically pigmented in most other members of the subgenus Sophophora (e.g. D. willistoni, Fig. 7.6). In D. melanogaster, the sexually dimorphic pattern is controlled by the bric-à-brac (bab) locus, which encodes two related transcription
factors that act as repressors of pigmentation. In *D. melanogaster*, female Bab proteins are expressed throughout the developing abdominal epidermis, but in males the Bab proteins are repressed in the fifth and sixth abdominal segments (Fig. 7.6), allowing for pigmentation of these segments. The sex- and segment-specific regulation of *bab* is controlled by the Doublesex and Abdominal-B proteins of the sex determination and homeotic segment identity systems, respectively. In males, Abd-B represses *bab* in the fifth (A5) and sixth (A6)
abdominal segments while in females the female-specific isoform of the Doublesex protein activates \textit{bab} in A5 and A6 to sufficient levels to suppress pigmentation.

Comparison of Bab expression in a large number of species reveals that, in most species with male-specific pigmentation, Bab is expressed in a similar sex- and segment-specific pattern. In species in which pigmentation is identical between the sexes, Bab is expressed throughout the abdomens of both sexes. This suggests that evolutionary changes in the regulation of \textit{bab} expression in the \textit{melanogaster} species group played a key role in the origin of sexually dimorphic pigmentation. Because DSX and Abd-B expression are conserved across species, the favored model for this innovation would be the evolution of binding sites for the DSX and Abd-B transcription factors in the \textit{cis}-regulatory control element(s) of the \textit{bab} gene that govern expression in the developing abdomen.

Further support for a central role of the \textit{bab} locus in the evolution of abdominal pigment patterns comes from studies of both intraspecific variation in \textit{D. melanogaster} and interspecific divergence throughout the entire subfamily Drosophilinae. In \textit{D. melanogaster}, females exhibit considerable variation in the degree of pigmentation in the posterior abdomen. Quantitative genetic analysis has indicated that variation at the \textit{bab} locus accounts for approximately 70% of the phenotypic variation in females. This finding has two important implications. First, it suggests that variation at \textit{bab} has a major effect on pigmentation and that selection on the \textit{bab} locus is a likely scenario for the origin of sexually dimorphic pigmentation. Second, it also indicates that other loci contribute to variation and divergence.

Bab’s role in the diversification of pigmentation patterns is broader than just within the \textit{melanogaster} species group. A survey of a large number of species from across the subfamily with conspicuous and diverse patterns of abdominal pigmentation revealed a correlation between \textit{bab} expression and melanism patterns in a majority of species. This indicates that evolutionary changes in \textit{bab} have occurred multiple times in independent lineages to create different patterns. For example, in \textit{D. tripunctata}, \textit{bab} is repressed in circular patches of cells along the midline of the A4, A5, and A6 segments in both sexes, which correlates very well with the development of intense pigmentation in these patches (Fig. 7.6). In \textit{D. funebris} and other species, Bab is expressed at high levels along the midline of the abdomen, which correlates with repression of melanin formation (Fig. 7.6). Dimorphic regulation of \textit{bab} expression in the posterior abdomen has occurred several times in the course of \textit{Drosophila} evolution.

Importantly, there are exceptions where \textit{bab} is not associated with abdominal pigmentation, even for sexually dimorphic patterns. These findings illustrate that while evolution at \textit{bab} is the most frequently exploited path to abdominal pigmentation pattern diversification, it is not the only path. The genetic architecture of pigmentation regulation in fruit flies is such that variation at other loci can be selected upon if evolution at \textit{bab} is constrained, just as we saw for evolution of the \textit{MC1R} and \textit{agouti} genes in the evolution of mammalian fur colors.

\textbf{NODAL POINTS IN REGULATORY NETWORKS AND THE EVOLUTION OF CHARACTER NUMBER AND PATTERN}

In the examples above of pigmentation evolution in vertebrates and insects, genes such as \textit{MC1R}, \textit{bab}, or \textit{yellow} were frequently involved in phenotypic divergence, but there were
other genetic routes to similar phenotypic changes. This illustrates that an apparently simple developmental readout (such as melanic pigmentation) can be modified by a variety of mechanisms.

Other characters, however, may be governed by genetic regulatory architectures that may have fewer or greater possible genetic inputs, and therefore different degrees of genetic constraints on evolution. For example, innervated bristles and noninnervated hairs are two types of epithelial projections that adorn most surfaces in insects and other arthropods, and whose formation requires the activity of the regulatory genes of the Achaete-Scute Complex (see Chapter 2) and the shaven-baby gene, respectively. Each of these genes acts as a nodal point (see Chapter 3) that integrates spatial information and whose expression and function is absolutely required for formation of the respective pattern elements. Therefore, diversification in the number or pattern of these structures will always be associated with differences in the expression of these regulatory genes. The key question then is the degree to which diversity is achieved through changes at these regulatory loci, or at other loci whose products affect the expression of these regulatory genes.

In most Drosophila species, there are intricate patterns of hairs covering large portions of the larval body. In D. melanogaster, the formation of these hairs requires the activity of the aptly-named Shavenbaby (Svb) transcription factor (mutants in svb lack hairs, appearing “shaven”), whose expression precisely foreshadows the position of denticles and hairs in the larval epidermis. In the melanogaster species group, all members except for D. sechelia bear rows of fine hairs on the dorsal part of larval segments (Fig. 7.7). The close relationship of members of this group allow for some interbreeding in the laboratory, which revealed that the difference in D. sechelia is entirely due to changes at the svb locus, and is correlated with the lack of svb transcription in the region where the hairs form in other species. These findings indicate that cis-regulatory evolution at svb underlies this discrete phenotypic difference.

As in the case of pigmentation pattern diversity, the loss of larval hair patterns is not restricted to just one species, but has also occurred repeatedly in the distantly related D. virilis group (which some 40–60 Ma diverged from D. melanogaster). In each instance, regulatory changes within the svb locus are associated with the phenotypic change. This is not expected because changes in the deployment of an activator of svb would also be a plausible mechanism for hair loss. The parallel, independent loss of dorsal hairs via regulatory evolution at the svb locus demonstrates that morphological evolution may occur through the same developmental mechanism and that such parallel changes in different lineages may be more prevalent than has been anticipated.

The bristle patterns of adult fruit flies also exhibit considerable intraspecific variation and interspecific divergence. Drosophila melanogaster is covered with sensory bristles that serve mechano- and chemoreception functions. Both the number and pattern of bristles on various body parts can vary between individuals as well as between populations. The most successful approaches to understanding intraspecific variation exploit genetic methodologies for assessing the number and identity of genes involved in morphological differences between populations. Much is known about the genes that control the bristle pattern and their functions in the developmental processes that give rise to it. Consequently, the variation and evolution of bristle patterns offers great potential for exploiting our emerging knowledge of developmental genetics so as to better understand morphological variation.
Figure 7.7
Evolution of trichome patterns via regulatory evolution at the shavenbaby gene

The dorsal hair patterns of five members of the melanogaster species group shown schematically. The distribution of short denticles (small blue projections), fine hairs (curved thin lines), and large denticles (large blue projections) on an individual segment are depicted (anterior cells are open rectangles, posterior cells are shaded blue). In D. sechelia, the fine hairs are absent due to differences at the shavenbaby locus (see photographs on right).

Source: Courtesy David Stern and adapted from Sucena E, Stern DL. Proc Natl Acad Sci USA 2000; 97: 4530–4534.
Bristle number is an example of a quantitative trait that varies continuously in populations. Other quantitative traits include features such as body size, organ size, and life span. Quantitative genetic techniques seek to elucidate the number and identity of those loci (quantitative trait loci (QTLs)) that contribute the bulk of the genetic variance underlying the morphological variance in characters under study. A useful distinction is to discriminate between genes of “small effect,” many of which may combine to account for some portion of the variance in a trait, and genes of “large effect,” which may account for 5–10% (or much more) of the variance in a given trait.

Studies of Drosophila bristle variation have shown that many loci affect bristle number but only a few loci have large effects that cause most of the variation. Some QTLs are genes known to be involved in bristle patterning, including the achaete-scute, scabrous, and Delta genes. The achaete and scute genes encode transcription factors required for neural precursor formation, whereas the Delta and scabrous genes encode ligands involved in the Notch-mediated signaling pathway that regulates cell interactions in proneural clusters, and ultimately, achaete and scute expression. Because mutations in these genes have dramatic effects on bristle development, their identification as QTLs in bristle number variation implicates them as playing a role in the evolution of bristle number.

The variation at all three loci appears to occur within regulatory regions. In the achaete-scute region, for instance, DNA insertions are strongly associated with variation in bristle number in natural populations. These insertions do not disrupt the protein coding regions, but they do appear to affect the expression of the achaete and scute genes. Two sites within the Delta gene are also associated with bristle variation. Each of these sites is located in an intron, rather than a coding segment of the gene. Similarly, sites associated with bristle variation in the scabrous gene appear within particular regulatory regions. These studies of Drosophila bristle variation have also revealed that the molecular basis for the genetic variation in loci that contribute to morphological variation can involve two or more sequence differences in a gene, not just a single nucleotide substitution.

Another emerging and powerful experimental approach to studying the genetics of morphological variation involves artificial selection on traits of interest. Rather than relying solely on the expressed quantitative variation present in natural populations, it is possible to derive populations with much greater divergence in traits through repeated selection over several generations for individuals with character states at either end of a continuum.

Experiments have been performed to select lines of flies with greater or fewer bristles. Genetic crosses between fly lines with “high” and “low” bristle numbers can be used to estimate the number of loci involved in the divergence and to map the quantitative contributions of genetic regions and individual loci. Many of the candidate loci identified in these experiments encode proteins known to play roles in the patterning and genesis of bristles. Although the specific molecular bases of the genetic differences responsible for the selected morphological divergence remains unknown, we do understand one crucial finding in both natural and artificially selected populations—that is, many genetic loci typically contribute to differences in such modest traits as bristle number on a single fruit fly body part. Because the products of these genes interact through regulatory networks, the effects of genetic variation are usually not strictly additive. Combinatorial interactions—so powerful in developmental processes—clearly play major roles in morphological variation and evolution.
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QUALITATIVE AND QUANTITATIVE ASPECTS OF SKELETAL EVOLUTION IN STICKLEBACK FISH

Pigmentation, hairs, and bristles are all characters displayed on the body surface, so perhaps it is not surprising that extensive variation and divergence occurs in these traits on very similar or identical body plans. One might expect that traits such as elements of the vertebrate skeleton would be more constrained and slower to evolve. This notion is shattered by the remarkable evolutionary history of the threespine stickleback fish (*Gasterosteus aculeatus*) in North America.

This species colonized many newly created lakes and streams at the end of the last ice age. In the course of just 15,000 years, rapid parallel evolution has taken place that has produced similar pairs of differentiated species in numerous locales. In general, in each location benthic forms with reduced body armor, increased body size, and reduced gill rakers (for filtering ingested food) have differentiated from limnetic forms with more extensive body armor, a longer body, and increased number of gill rakers (Fig. 7.8). The two forms are reproductively isolated in the wild, but can be intercrossed in the laboratory to allow for detailed investigation of the genetic architecture of anatomical and behavioral differences between body forms.

Figure 7.8
Evolution of limnetic and benthic forms of the threespine stickleback
Limnetic (top) and benthic (bottom) forms of the threespine stickleback from Priest Lake, British Columbia. Differences in spine length, armor plate number, and gill raker number have evolved as adaptation to different niches.
Source: Photographs courtesy of Katie Peichel and David Kingsley.
While some of these differences appear to be due to variation at many loci of small effect, a small number of loci were found to account for much of the variance in other characters. Interestingly, the sets of QTLs affecting the length of the first and second dorsal spine were distinct, while those affecting the second dorsal spine and pelvic spine overlapped with one another. These observations reveal that genes with a range of magnitude of effects have contributed to evolutionary change, and that very similar morphological features can be affected differently by the same gene. A key outstanding question presented by the threespine stickleback, in addition to the identity of genes involved in divergence, is whether the same loci are involved in the independent evolution of similar forms in different locales.

MORE VARIATION THAN MEETS THE EYE: CRYPTIC GENETIC VARIATION AND THE POTENTIAL FOR MORPHOLOGICAL EVOLUTION

In addition to the overt, phenotypic variation in the characters described here, developmental geneticists have uncovered more variation lurking in individuals than is generally appreciated (“cryptic variation”). The importance of gene interactions to morphological variation has been underscored by some fascinating studies of the phenotypes that arise when mutations are introduced into individuals with different genetic backgrounds.

While a single mutation is often described as causing a particular phenotypic change in laboratory studies, the effect, in practice, depends on the context of the genetic background in which a mutation is studied. For example, it is well established that in laboratory lines of *D. melanogaster*, introduction of dominant mutations in the Sevenless tyrosine kinase receptor and EGF receptor (*DER*) induces a roughening of the surface of the adult eye. This outcome occurs because of a perturbation of developmental events in eye patterning that are dependent on the Sevenless and DER pathways. By contrast, introduction of these mutations into wild-type flies of different origins (and different genetic backgrounds) produces a considerable range of severity in phenotypes. In some genetic backgrounds, the mutant phenotypes are suppressed as compared to their effects in laboratory strains; in others, they are enhanced (Fig. 7.9). The enhancement observed in some backgrounds sometimes exceeded that caused by the combination of Sevenless or DER mutations, and mutations in additional components of the Sev or EGF-R signal transduction pathways. This finding indicates that considerable genetic variation occurs in the wild in loci affecting the function of major pathways.

Similar observations have been made from introducing homeotic mutations into different genetic backgrounds. Both extreme modification and suppression occurs. In some cases, this disparity reflects the presence of single genes of large effect that modify the homeotic phenotype, but have no stand-alone effect on wild-type development. Such studies demonstrate that there may be widespread genetic variation that does not discernibly affect phenotypes unless certain other interacting mutations are present. Evolutionary and developmental geneticists are only now beginning to truly appreciate this cryptic variation. Importantly, this hidden variation implies that underlying phenotypic stability, the quantitative aspects of genetic regulatory inputs may vary extensively. Furthermore, it suggests that the existing genetic variation available for the selection of new phenotypes during evolution may be significantly greater than previously thought.

One striking illustration of this latter idea has come from artificial selection for homeotic phenotypes in *Drosophila*. It has long been known that homeotic phenotypes can sometimes
be obtained by environmental insults to embryos during sensitive periods. Treatment of developing wild-type flies with ether vapor, for example, can induce mimics or phenocopies of homeotic transformations caused by mutations in the Ubx gene. C. H. Waddington showed decades ago that if one repeatedly selected for individuals demonstrating the bithorax phenocopy, the resulting populations would show a greater frequency of response to the treatment, including individuals that displayed the phenotype independent of treatment. The selection therefore uncovered genetic variation that affects development.
Developmental and molecular genetic analysis of the Ubx gene in response to selection for the ether-induced bithorax phenocopy revealed that ether induces loss of Ubx expression in patches of cells in the haltere imaginal disc. In selected populations, the frequency of lost gene expression increases and is heritable. Therefore, genetic differences must exist, either in cis or trans to the Ubx gene, that influence the susceptibility of the Ubx gene to loss of expression following exposure to ether. One site that responded to selection for phenocopy induction was mapped downstream of the Ubx coding region in a large cis-regulatory region. This finding suggests that genetic variation in a cis-regulatory element of the Ubx gene affects the fly population’s susceptibility to loss of gene expression in response to ether.

Cryptic variation is also detectable between species. D. melanogaster and D. simulans adults exhibit identical patterns of thoracic bristles, a pattern that is highly stereotyped and rarely variant among individuals. Yet, when interspecific hybrids are produced, the hybrids lack a variable number of bristles. The loss of bristles is associated with decreased transcription of the achaete and scute genes and genetic experiments suggest that divergence in achaete/scute cis-regulatory elements and in interactions with trans-acting genes (regulators of achaete/scute) has occurred between species. These observations suggest stasis in adult patterns is maintained by compensatory interactions within each species and that this compensation is disrupted when alleles of each species are combined in hybrids.

The phenotypic variation unmasked by artificial selection or interspecific hybrids suggests that there is a reservoir of cryptic genetic variation among developmental genes that can be tapped during evolution by natural selection. Changes in the environment or introduction of a new mutation into a population can quickly uncover genetic variation as observable phenotypic variation. The increase in phenotypic variation provides a broader range of fitnesses upon which natural selection may act.

REGULATORY EVOLUTION AND SPECIES DIVERGENCE

There are four major concepts that can be distilled from the case studies presented in this chapter. First, the major determinant of the genetic architecture of trait evolution is the genetic architecture governing trait development. When traits are governed by many genes that act at different steps of a developmental process, such as coloration in vertebrates or melanin patterning and bristle development in insects, there are multiple evolutionary paths to phenotypic divergence. However, when the formation of a pattern element or structure is, as in the case of Drosophila hairs or trichomes, governed by a single gene, evolution at that regulatory gene is a primary mechanism of phenotypic diversification.

The second major idea concerns the phenomenon of parallel evolution. In several cases, similar but independent evolutionary changes in different lineages have arisen through evolution at the same loci, even when there are alternative paths to the same final pattern. This has not been generally anticipated and suggests that similar genetic regulatory changes may account for the diversification of particular patterns in groups of species.

Third, phenotypic divergence involving pleiotropic genes is most often associated with evolution of gene regulation. This was demonstrated or inferred to involve evolution in modular cis-regulatory elements governing gene expression in discrete body regions. We saw in Chapters 5 and 6 how the diversification of metameric body plans and serially homologous body parts has occurred through evolution of Hox gene regulation and downstream target
genes. In this chapter we described an example of a Hox-regulated target gene (bab) and a character (abdominal pigmentation) that exhibit both intraspecific variation and interspecific divergence. It is a reasonable extrapolation that the sorts of genetic regulatory mechanisms underlying variation and divergence within and between species are responsible for the larger-scale divergence we see at higher taxonomic levels and over greater spans of time.

In Chapter 8, we will examine how variation in gene regulation arises, its prevalence in natural populations, and how this variation provides the raw material for the evolution of diverse animal forms.

**SELECTED READING**

**General Review**


**Melanism in Vertebrates**


Millar SE, Miller MW, Stevens ME, Barsh GS. Expression and transgenic studies of the mouse agouti gene provide insight into the mechanisms by which mammalian coat color patterns are generated. *Development* 1995; 121: 3223–3232.


Rieder S, Taourit S, Mariat D, Langlois B, Guérin G. Mutations in the agouti (ASIP), the extension (MC1R), and the brown (TYRP1) loci and their association to coat color phenotypes in horses (*Equus caballus*). *Mamm Genome* 2001; 12: 450–455.


**Pigmentation Diversity in Drosophila**


**Trichome and Bristle Development and Evolution**


Long AD, Lyman RF, Langley CH, Mackay TFC. Two sites in the *Delta* gene region contribute to naturally occurring variation in bristle number in *Drosophila melanogaster*. *Genetics* 1998; 149: 999–1017.


**Evolution in Stickleback Fish**


**Cryptic Variation: Quantitative and Developmental Genetic**


CHAPTER 8

From DNA to Diversity: The Primacy of Regulatory Evolution

In the final chapter of this book, we consider why regulatory evolution is the major creative force underlying morphological diversity across the evolutionary spectrum, from variation within species to body plans. The link between DNA sequence evolution and phenotypic diversity often involves cis-regulatory elements acting as units of evolutionary change. Here, we take an in-depth look at the function of cis-regulatory elements and molecular models and examples of their evolution. We will examine a few experimental approaches that have begun to reveal the direct connections between changes in regulatory DNA, the evolution of gene regulation, and the origins of morphological diversity.

cis-regulatory elements have properties that make their evolutionary dynamics distinct from coding sequences. The turnover of transcription factor binding sites within elements with conserved functions reveals that compensatory mutation is a critical feature of cis-regulatory DNA evolution. Both theoretical considerations and experimental evidence suggest that the modification of cis-regulatory element function can readily occur through point mutations that create or destroy transcription factor binding sites. Variation within species at such sites appears to be widespread. A growing body of evidence supports the view that regulatory evolution at the species level is sufficient to account for the larger-scale patterns of morphological evolution, including the origin of humans.

WHY IS REGULATORY EVOLUTION A PRIMARY FORCE IN MORPHOLOGICAL EVOLUTION?

Many case studies in Chapters 5, 6, and 7 illustrated that selected changes in gene regulation in one part of the developing animal,
independent of other parts, underlie the morphological diversification of serially homologous structures, the origin of novelties, and the divergence of traits between species. Regulatory evolution is the enabling genetic mechanism for the modular organization and diversity of larger bilaterian phyla and for the emergence of new morphological characters.

The argument made for the central importance of regulatory change in morphological evolution is not a new one. Indeed, the creative potential of regulatory change and the comparatively greater constraints on protein evolution were recognized early in the history of molecular biology. Now there is considerable empirical evidence which demonstrates that changes in the regulation of genes that affect morphology are implicated much more frequently in the evolution of diversity than are new genes or functional changes in protein sequences. Furthermore, our current understanding of genetic regulatory hierarchies, networks, and circuits in development is capable of revealing why this is so.

Specifically, regulatory evolution is powerful because of the following characteristics:

1. **Regulatory evolution enables pleiotrophy of toolkit genes.** The same transcription factor usually controls different target genes in different tissues at different stages of development. The DNA binding activity of a given protein usually remains the same throughout development, however, it is evolution within the regulatory sequences of target genes that enables these genes to respond in a context-specific fashion.

2. **Regulatory evolution enables developmental modularity.** For anatomical structures to become different from serial homologs within an animal or from homologs in other animals, changes must evolve in the regulatory hierarchies that operate during the development of these structures. Due to their modular organization, changes in *cis*-regulatory elements allow changes in gene expression to occur in one structure independently of another (this independence is also referred to as dissociation). In metazoans, the great success (in terms of species diversity and adaptation to terrestrial, aquatic, and aerial environments) of highly modular body plans such as the arthropods, vertebrates, and annelids suggests that the modularity of developmental regulatory mechanisms at both the anatomical (for example, fields) and molecular (*cis*-regulatory elements) levels have facilitated this diversity.

3. **Regulatory evolution is a rich and continuous source of variation.** The *cis*-regulatory regions may occupy vast stretches of DNA and contain many independent functional elements. Changes in regulatory sequences within individual elements may subtly affect the level, timing, or spatial pattern of gene expression, perhaps very selectively in terms of the tissues and stages of development involved. Without any change in protein sequences, these quantitative, temporal, and spatial changes in the deployment of regulatory genes may affect the level, timing, and spatial expression of other developmental genes. Regulatory changes are often cryptic and have little effect on morphology, but they nevertheless create genetic variations with the potential to produce the morphological variation that is the raw material of evolution. Regulatory DNA is a rich and continuous source of potential genetic, developmental, and phenotypic variation, and thus evolutionary change.

4. **Regulatory evolution creates novelty.** The exploration of new morphologies is facilitated by new combinations of gene expression that can arise without changes in protein function.
Chapter 8: From DNA to Diversity: The Primacy of Regulatory Evolution

THE FUNCTION AND EVOLUTION OF \textit{CIS}-REGULATORY DNA

\textbf{Functions of \textit{cis}-regulatory elements}

Several examples of \textit{cis}-regulatory elements and \textit{cis}-regulatory regions were presented in Chapter 3 in the context of understanding the genetic regulatory networks that orchestrate pattern formation. To better understand the role played by \textit{cis}-regulatory DNA in evolution, it is important to appreciate several general features of \textit{cis}-regulatory systems.

First, most elements are regulated by a minimum of four to six transcription factors of various structural types. Transcription factors virtually never act alone, so the output of individual \textit{cis}-regulatory elements is determined by the integration of multiple diverse inputs.

Second, the spatial relationship of binding sites for transcription factors within a \textit{cis}-regulatory element can be of utmost functional importance. Within the few hundred base-pairs of a typical element, the proximity of binding sites can determine whether transcription factors interact cooperatively. For example, multiple lower-affinity sites may have a greater effect on transcription through cooperative interactions than a single high-affinity site. Most metazoan transcription factors bind a family of DNA sequences with some degree of degeneracy in the sequences recognized. The “flexibility” of transcription factor binding opens up greater opportunities for cooperative interactions.

Third, repression—not activation—is generally the ground state of gene expression. That is, the chromatin that packages DNA has generally repressive effects on transcription. The \textit{cis}-regulatory elements represent sites where protein complexes are assembled that attempt to overcome this repression.

Fourth, spatial boundaries of gene expression in a cellular field are usually set by both positive and negative inputs into \textit{cis}-regulatory elements. Loss of positive inputs (in \textit{trans}) or of their binding sites (in \textit{cis}) eliminates or contracts the spatial domains of gene expression. Conversely, loss of repressors or their sites expands spatial domains. The sites where activators and repressors bind may overlap; in those cases, competition for sites may determine gene activity. In other cases, they may not overlap and the activators and repressors modulate the output of the element through short-range (fewer than 100 bp) or long-range (more than 100 bp) effects on transcriptional functions.

Fifth, expression of terminal differentiation genes, such as those encoding structural proteins that carry out functions specific to certain cell types, are often regulated by \textit{cis}-elements that respond largely or entirely to positive transcriptional activators. Hence, genes found progressively farther downstream from the regulators that establish spatial domains in the developing embryo may not require negative inputs to prevent their expression in inappropriate positions or cell types.

In summary, \textit{cis}-regulatory elements are evolved devices. The number, type, and topology of transcription factor binding sites within any element is shaped by very different kinds of functional considerations than those affecting coding sequences. These distinct functional features govern the rate and means by which new elements arise and extant elements vary and evolve.
Evolution of cis-regulatory elements

Evolutionary changes in a given gene’s expression may arise through a variety of mechanisms. That is, they may arise directly from alterations in the cis-regulatory DNA of the gene or they can occur indirectly, through changes in the deployment or activity of upstream transcription factors that regulate the gene (which ultimately reflect changes in the cis-regulatory elements of genes encoding these transcription factors or their regulators). In the remainder of this chapter, we focus on the developmental and evolutionary consequences of changes in cis-regulatory elements.

The cis-regulatory DNA function may evolve through any of several molecular mechanisms. Here, we distinguish between two major potential sources of cis-regulatory DNA evolution:

- The de novo evolution of cis-regulatory elements through changes in nonfunctional DNA (Fig. 8.1a).
- The evolution of cis-regulatory elements from preexisting functional elements (Fig. 8.1b–d).

Within the latter category, we distinguish between three modes of sequence evolution:

- Duplications and DNA rearrangements involving existing functional elements that create additional copies of elements or new elements (Fig. 8.1b).
- Modifications of existing cis-regulatory elements, for example, through the gain or loss of binding sites for positive and/or negative regulators (Fig. 8.1c).
- The special case of co-option of an existing element that expands its developmental function (Fig. 8.1d).

First, we examine, from a theoretical viewpoint, the factors affecting the probabilities of these different mechanisms of cis-regulatory DNA evolution. Next, we analyze some illuminating case studies of cis-regulatory element evolution.

De novo evolution of cis-regulatory elements from preexisting nonfunctional DNA sequences

The vast majority of animal genomes are composed of noncoding DNA. Consequently, extensive stretches of DNA sequences exist that could potentially function to regulate the expression of adjacent genes. Much of this DNA consists of various lengths of repetitive elements, some of which may have architectural functions (for example, the chromosome centromere), but most of which are believed to be nonessential and not involved in the regulation of specific genes. In addition, an abundant amount of single-copy DNA occurs between coding regions.

In principle, such unconstrained sequences could evolve into functional cis-regulatory elements. The key question is, How “difficult” is it for random DNA sequences to evolve into a new regulatory element? To answer this question, we must consider two major factors. The first factor is the probability that one or more binding sites evolve for individual specific DNA-binding proteins. The second consideration is the minimal input required for cis-element
Chapter 8: From DNA to Diversity: The Primacy of Regulatory Evolution

**Figure 8.1**

Molecular mechanisms of cis-regulatory DNA evolution

Four modes of cis-regulatory element evolution are depicted. (a) De novo evolution of an element from nonfunctional sequences could occur through mutations ($\Delta_i$, $\Delta_{ii}$, and so on) that create binding sites for transcription factors (TF) in proximity to a transcription unit. (b) New elements may evolve by duplications of genes or of cis-regulatory elements, followed by sequence divergence (left), or DNA rearrangements that move elements to new sites in the genome (right). (c) Modifications of existing cis-regulatory elements can occur through the gain, loss, or change in affinity of sites for activators (TF) or repressors (R). (d) A special case of cis-regulatory element modification is co-option of an existing element to function in a new domain through the gain of sites for specific factors (TF).
function; more specifically, a novel element requires a degree of function upon which positive selection can act.

The probability that binding sites for transcription factors will evolve depends on the properties of DNA-binding proteins. Individual DNA-binding proteins typically recognize a 5–9 bp core sequence, usually with some flexibility (that is, degeneracy) in the identity of bases found at particular positions. Assuming an equal frequency of all four bases in a genome and random distribution, we can calculate the frequency with which a given length site will appear in a genome (Table 8.1). For example, a specific 6-bp sequence will occur, on average, once every 4096 bases. We can also relate these frequencies to genome size and to the approximate number of genes in various animal genomes (Table 8.1). For example, in a genome the size of that of *Drosophila melanogaster* (approximately 1.15 × 10⁸ bp of single copy DNA) or of humans (approximately 1.66 × 10⁹ bp of single copy DNA), a given 6-bp sequence should occur roughly 28,000 and 400,000 times, respectively. Based on the number of genes in these species, these calculations demonstrate that single binding sites for proteins with a six-base core recognition sequence will occur, on average, at least once per gene.

These frequency calculations apply only to a single genome at a given time. Obviously, species are composed of potentially very large populations of individuals and have characteristic generation times. Over the course of evolutionary time, the steady occurrence of random mutations will increase the probability that a given binding site will arise in a given binding genomic location at some time in a population. Whether that site is preserved (“fixed”) depends upon a number of factors, including the effective population size of the species and the selective advantage or disadvantage of alleles bearing that site relative to other alleles in the population.

We are most interested in the question of whether the appearance of any binding site is functionally significant. Answering this question requires that we consider the requirements for *cis*-regulatory element function, the diversity of the transcription factor repertoire, and the level of gene expression upon which selection can act. The *cis*-regulatory element function

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**TABLE 8.1 The probable frequencies of various length sites for DNA-binding proteins in animal genomes**

<table>
<thead>
<tr>
<th>Binding Site Length (bp)</th>
<th>Frequency of site (1/no. of base pairs)</th>
<th>Approx. Number of Sites in Genome; per Gene</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Drosophila*</td>
<td>Homo sapiens†</td>
</tr>
<tr>
<td>5</td>
<td>1024</td>
<td>112,500; 8.2</td>
</tr>
<tr>
<td></td>
<td>4096</td>
<td>28,125; 2.1</td>
</tr>
<tr>
<td>6</td>
<td>16,354</td>
<td>7031; 0.5</td>
</tr>
<tr>
<td>7</td>
<td>65,536</td>
<td>1758; 0.1</td>
</tr>
<tr>
<td>8</td>
<td>262,144</td>
<td>439; 0.03</td>
</tr>
<tr>
<td>9</td>
<td>1,048,576</td>
<td>110; 0.008</td>
</tr>
<tr>
<td>10</td>
<td>16,777,216</td>
<td>7; 0.0005</td>
</tr>
</tbody>
</table>

* Assumes approx 1 × 10⁸ bp genome; 15,000 genes
† Assumes approx 5 × 10⁹ bp genome; 50,000 genes
typically depends on a host of binding sites for diverse regulatory proteins acting in some proximity to one another. Gene activation must overcome the generally repressive effects of chromatin through the recruitment of co-activator proteins by transcription factors. In most elements, single binding sites or single transcription factors are not sufficient to confer significant transcriptional activation. Likewise, gene activation typically involves the synergistic effects of two or more activators bound to nearby sites. If these requirements need to be met to achieve the levels of gene activation upon which selection can act, then our calculations in Table 8.1 must be extended to account for the probability of two or more sites arising within an interval of the typical length (about 200 bp) of a \( \text{cis} \)-regulatory element (Table 8.2).

For example, target gene activation by several Hox proteins involves interactions with the Exd homeodomain protein bound to an adjacent binding site. A typical Hox/Exd composite site contains a critical 10-bp core sequence. A specific 10-bp sequence will occur at random approximately once every 1 million bp (Table 8.1). If DNA is measured in 200-bp intervals, such a rare site would appear at random only once per any 5000 intervals (Table 8.2). In comparison, combinations of shorter sequences would arise more frequently. For example, two specific 5-bp sequences are 200 times more likely to appear in a 200-bp interval than they are in a single 10-bp site (Table 8.2).

Experiments have shown that transcription factors can often interact synergistically with a variety of other transcription factors. This finding implies that for a new binding site to function within an evolving element, the only requirement may be that other transcription factors bind nearby. The identity or type of transcription factor may not be critical. For example, classic studies of steroid receptor-binding sites have demonstrated that a wide array of transcription factors can interact synergistically with receptor–DNA complexes to induce gene activation. Thus a steroid receptor-binding site (or, in principle, a site for any other DNA-binding activator protein) that arises at random could impart activity upon a nearby gene if nearby DNA sequences are also bound by other transcription factors.

To analyze the probability that nearby binding sites will be occupied, we must have an understanding of the transcription factor repertoire in a typical animal cell. Such information is just emerging from genome sequencing and genome-wide screens of gene expression. As
a result, we are beginning to get an idea of the size and complexity of the entire transcription factor toolkit.

In *C. elegans*, for example, of the roughly 19,000 protein coding genes in the genome, approximately 400 are predicted to be DNA-binding transcription factors. To calculate the probability that one or more of these transcription factors will bind to sites near each other or the site of another transcription factor, we must make some inferences about the diversity of transcription factors expressed in any given cell. The 400 or so transcription factors in *C. elegans* are expressed in a total of about 30 cell types. Although some fraction of these transcription factors may be expressed in all cells, the distribution of most likely remains restricted to a few cell types. For discussion purposes, we’ll assume that each transcription factor acts in five cell types on average. We can extrapolate that if each of these 400 different factors is expressed in five cell types, then the average cell contains $400 \times \frac{5}{30}$ or roughly 60 different transcription factors.

Assuming an average 6-bp recognition sequence, for any 20 transcription factors with non-identical binding specificities, at least one site probably exists for one of the 20 transcription factors every 200 bp of DNA sequences. With 60 factors per cell, then at least three sites for the average set of transcription factors would exist in every 200 bp. These calculations suggest that for any randomly evolved site, a few other potential sites for different transcription factors are likely to exist nearby. A new site appearing in this context may be sufficient to affect gene activity.

The theoretical considerations discussed in this example largely focus on the probabilities surrounding the evolution of 6-bp sites. In reality, many transcriptional regulators, such as homeodomain proteins, recognize shorter core sequences and exhibit a range of measurable affinities for families of related sequences.

Homeodomains, for example, have evolved to be able to interact with many different DNA sequences. An important implication of this flexibility of homeodomains (and potentially other regulatory proteins) is that they can bind to shorter, more abundant, lower-affinity sites. Such sites may consist of functional intermediates in the course of *cis*-regulatory DNA evolution upon which positive selection could act.

**Evolution of cis-regulatory elements from existing elements**

General chromatin repression and the calculations given in the previous example suggest that limitations and uncertainties surround the probability of de novo element evolution. Surmounting these limitations through the accumulation of single, initially neutral base changes is not the only means of evolving new functional elements. Indeed, one way to evolve new elements more readily might be to derive them from preexisting functional elements. The three other *cis*-regulatory element evolution scenarios depicted in Fig. 8.1(b–d) are all variations on this theme, albeit with important distinctions that are delineated below.

**Duplications and DNA rearrangements** Perhaps the simplest way that genomes can “experiment” with the expression of individual genes is by shuffling *cis*-regulatory DNA in the genome (Fig. 8.1b). Tandem duplications expand the number of *cis*-regulatory elements, and recombination, transposition (for example, of mobile elements), inversions, and translocations all create new juxtapositions of DNA sequences. If functional *cis*-regulatory DNA (including
all or part of an element) becomes newly juxtaposed to another cis-regulatory element or transcription unit, the rearrangement may affect gene expression. In addition, new combinations of regulatory sequences may be created that have properties differing from those of either the new or resident sequence.

Because regulatory DNA is not constrained to maintain any reading frame, spacing, or polarity, cis-regulatory DNA is more likely than coding regions to tolerate and thus accumulate partial duplications and insertions that expand the overall element. Local duplications of elements or entire genes can create functional redundancy that may be retained by positive selection because of the duplication’s effects on gene expression levels. Expanded elements with duplicated sites may be more tolerant both to substitutions that can cause functional divergence and to DNA rearrangements that separate or subdivide parts of an element. Acting in this manner, existing cis-regulatory DNA elements could give rise to new elements much more readily than could the gradual accumulation of single base changes. Indeed, it has been estimated that the rate of gene duplication is 0.01 per gene per million years, which is the same order of magnitude as the mutation rate per nucleotide site. This suggests that smaller local duplications arise at a significant frequency and contribute to the evolution of regulatory DNA.

Transposable elements themselves appear to contribute to regulatory DNA evolution. In human genes, nearly a quarter of promoter regions contain sequences derived from one of the several most abundant types of transposable elements found in humans. Approximately 3% of cis-regulatory elements also contain sequences derived from transposable elements. Some of these elements may provide attachment of DNA regions to the nuclear scaffold or matrix. These observations suggest that transposable elements (which comprise 45% of the human genome) have been “domesticated” to serve functional roles in the regulation of individual genes.

Modification of existing cis-regulatory elements Any cis-regulatory element is subject to the constant forces of mutation and selection. Changes in nucleotide sequence may be neutral if they meet any of the following criteria:

- They fall outside of any site recognized by an essential regulator of an element.
- They do not alter the affinity of a critical binding site.
- The change in a binding site has negligible effects on the overall function of the element (which may be due to other compensatory changes that occur).

Changes may not be neutral if they affect the timing, level, or spatial domain of gene expression controlled by the element. Changes in the affinity or the gain or loss of activator sites and repressor sites can subtly or more dramatically affect cis-element function.

For cis-regulatory elements that regulate gene expression patterns within a particular field, the domain of expression controlled by individual elements can expand or contract simply by a net decrease or increase in activator sites or a net decrease or increase in repressor sites (Fig. 8.1c). For example, for a regulatory element that is controlled by a signal or transcription factor that is expressed in a concentration gradient, gene expression could broaden or narrow, respectively, if a gain/reduction in the number or a change in the affinity of activator sites...
occurs. Alternatively, gene expression could be excluded from all or part of a field through the evolution of sites for a spatially localized repressor.

**Co-option of an existing element and expansion of its developmental function**

The evolution of a novel cis-regulatory function (for example, controlling gene expression in a new pattern or cell type) could also arise through modification of an existing element (Fig. 8.1d). We consider this possibility here as a special case of modification of existing cis-regulatory elements, which may also subsequently involve mechanisms described for duplications and DNA rearrangements.

Given a functional element, if new sites for additional factors evolve, these new sites—acting in conjunction with functional sites bound by other transcription factors—might be able to co-opt an existing element such that it also functions in a new context (Fig. 8.1d). Such a bifunctional or multifunctional element could then control gene expression in two or more tissues or developmental stages.

There are no mechanistic grounds to preclude the evolution of multifunctional elements. We can postulate, however, that functional pressures might potentially lead to additional evolutionary changes in multifunctional elements. Multifunctional elements lack the advantage of two or more modular elements. The necessity to “fine-tune” gene expression in two or more contexts may favor the subdivision of an element into multiple independent elements. The most facile way for this subdivision to occur is for additional modifications to accumulate, such as internal duplications and the addition of factor binding sites that enhance the functional robustness of the element and make its subdivision possible. A subdivision could arise through DNA rearrangements, including the interposition of sequences that functionally isolate cis-regulatory domains.

**The evolution of gene repression versus gene activation**

From these various mechanistic scenarios, we can see that the modification of gene expression patterns within a field can be readily explained by the accumulation of single base changes in cis-regulatory DNA or through DNA rearrangements. A corollary of these observations may be especially pertinent to the trends in body plan diversification described in this book. That is, the selective repression of part of an existing gene expression pattern may be more readily evolved at the cis-regulatory level than are novel tissue-specific patterns of gene expression.

We have emphasized the trend in vertebrate, arthropod, and annelid evolution marked by the increasing diversity of serially homologous structures. The diversification of somites, segments, appendages, and other parts often entails the selective repression of developmental regulatory networks in one homolog versus another (for example, limb and wing repression in the insect abdomen). If the evolution of single repressor binding sites in cis-regulatory elements is sufficient to modify or eliminate gene expression in a field, it would appear to involve fewer binding sites and transcription factors than is required for the evolution of new patterns of gene expression. If, indeed, gene repression is “easier” to evolve than is gene activation, then the morphological diversity that follows from selective repression of gene expression may be one of the most important evolutionary correlates of the logic of metazoan cis-regulatory element function.
Chapter 8: From DNA to Diversity: The Primacy of Regulatory Evolution

Case studies in cis-regulatory evolution

The identification of specific cases representing any of the four modes of cis-regulatory element evolution has been a considerable challenge. While the preceding treatment outlines several conceivable possibilities, the analysis of the evolution of cis-regulatory elements of developmental genes has lagged behind the study of the evolution of gene number or protein sequence evolution. This lag is partly explained by the facts that knowledge of cis-regulatory element function in development is far from complete and that it is difficult to identify all of the diverse inputs into any given element.

The paucity of evolutionarily informative data also reflects methodological challenges that make it difficult to distinguish functional from nonfunctional changes in DNA sequences taken from different species. Because cis-regulatory DNA can better tolerate changes in sequence length and the number, spacing, and orientation of functional motifs, two homologous elements with a high degree of sequence divergence may nevertheless drive the same pattern of gene expression during development. A considerable body of comparative data exists for the sequences of homologous cis-regulatory elements taken from multiple taxa. Although these comparisons make it possible to identify conserved sequences, which are often sites for critical DNA-binding proteins, it remains difficult to determine which, if any, of the remaining sequences that are not shared between elements from different species has functional and therefore potential evolutionary significance.

Despite these obstacles, a rapidly increasing number of comparative studies of cis-regulatory elements have illustrated some general principles concerning the conservation, dynamic turnover, and functional modification of cis-regulatory DNA sequences in evolution. These principles are discussed next.

Conservation of functional elements among widely divergent taxa

Given a typical mutation rate of 1 nucleotide change per $10^9$ bp per generation, sequences that are not under any functional constraint (that is, selection) will diverge substantially over millions of generations. Thus, comparisons of sequences from sufficiently distant taxa can reveal sites that are under functional constraint. It is now common practice to isolate homologs of genes of interest from multiple taxa. Comparisons of their noncoding regions often prove helpful in identifying probable functional cis-regulatory sequences.

For example, comparisons of the mouse, chicken, and pufferfish Hoxb1 gene regulatory regions have revealed sequences that are shared among these three classes of vertebrates. In elements that control gene expression in the r4 rhombomere, four common sequence blocks include three binding sites for the Hoxb1/Pbx protein complex (Fig. 8.2). These shared sites indicate that this element has controlled Hoxb1 autoregulation since the common ancestor of vertebrates.

Similar comparisons have been carried out for the cis-regulatory elements of a wide variety of toolkit genes. Although these investigations often identify a pattern of small islands of sequence conservation in elements from long-diverged taxa, some elements demonstrate a remarkable degree of sequence conservation. For example, the element that controls expression of the Drosophila melanogaster vestigial gene along the dorsoventral boundary of the imaginal wing disc (see Chapter 3, Fig. 3.13) exhibits extensive sequence conservation with its counterpart in Drosophila viridis. In one stretch, 119 bases are perfectly conserved.
Some cis-regulatory elements may even exhibit conservation between phyla. For example, the labial gene of Drosophila, a member of the same Hox paralogy group as Hoxb1, is auto-regulated by a Lab/Exd protein complex. The similar functional sites in the autoregulatory elements of vertebrate Hoxb1 genes and the Drosophila lab gene could be due to conservation of an autoregulatory element from their last common ancestor.

The dynamics of sequence turnover in cis-regulatory DNA

The vast evolutionary distances between the taxa in the preceding examples allow us to confirm the significance of the conservation of particular sequences; they do not, however, elucidate the dynamics of cis-regulatory DNA evolution. For a better picture of that process, we must compare a greater number of more closely related species.

As an example, consider the homologs of the well-studied Drosophila melanogaster even-skipped stripe 2 element (Fig. 8.4a), which have been isolated from a host of other Drosophila species. These elements all drive an accurately positioned stripe of reporter gene expression in the eve stripe 2 domain in D. melanogaster, yet considerable sequence divergence occurs among the elements. Interestingly, some of the sequence changes in the other Drosophila species abolish sites that are known to be essential in the D. melanogaster element (Fig. 8.4b). For instance, the third Bicoid binding site (bcd-3) and the first Hunchback (hb-1) binding site found in D. melanogaster lack counterparts in D. pseudoobscura, D. erecta, and D. yakuba.
Chapter 8: From DNA to Diversity: The Primacy of Regulatory Evolution

The bcd-3 site is required for proper eve stripe 2 formation in *D. melanogaster*. This analysis suggests that the bcd-3 site may be a new site in *D. melanogaster*, whose function became essential in the context of other changes within the element.

Of the 17 known binding sites for transcriptional regulators in this element in *D. melanogaster*, only three are perfectly conserved among the other *Drosophila* species studied (Fig. 8.4b). Thus most binding sites exhibit a considerable degree of substitution. As each full element from the heterologous species drives an accurate eve 2 stripe in *D. melanogaster*, it appears that the various changes in binding sites compensate for one another, such that the net output of the element is the same (in *D. melanogaster*).

One evolutionary explanation for the observed pattern of functional conservation and sequence changes in the eve stripe 2 enhancer is that stabilizing selection governs the
Conservation of Transcription Factor Binding Sites in *Eve* Stripe 2 Enhancer

<table>
<thead>
<tr>
<th>Sites</th>
<th>Kr</th>
<th>Bcd</th>
<th>Hb</th>
<th>Gt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Species</td>
<td>1</td>
<td>2</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td><em>D. simulans</em></td>
<td>P</td>
<td>P</td>
<td>P</td>
<td>P</td>
</tr>
<tr>
<td><em>D. yakuba</em></td>
<td>P</td>
<td>S</td>
<td>P</td>
<td>P</td>
</tr>
<tr>
<td><em>D. erecta</em></td>
<td>P</td>
<td>S</td>
<td>P</td>
<td>P</td>
</tr>
<tr>
<td><em>D. pseudobscura</em></td>
<td>S</td>
<td>S</td>
<td>S</td>
<td>S</td>
</tr>
<tr>
<td><em>D. picticornis</em></td>
<td>W</td>
<td>S</td>
<td>W</td>
<td>S</td>
</tr>
</tbody>
</table>

Conservation

- **P** = perfect
- **S** = strong (1-2 changes)
- **W** = weak (3 or more changes)
- **A** = absent

Figure 8.4

*Evolutionary dynamics of transcription factor-binding site evolution in a conserved cis-regulatory element*

(a) Binding sites for the Krüppel (Kr), Giant (Gt), Bicoid (Bcd), and Hunchback (Hb) proteins in the 670-bp *D. melanogaster* eve stripe 2 *cis*-regulatory element are shown. (b) The conserved binding sites in five different *Drosophila* species are tabulated. The degree of sequence conservation within each site is indicated (P, S, W, A). Note that certain sites such as Kr5, Kr6, and Bcd5 are perfectly conserved, whereas other sites such as Hb1 and Bcd3 are absent from certain species.

dynamics of sequence turnover in the element. Because multiple sites exist for each regulatory protein, as well as both positive and negative inputs, slightly deleterious substitutions may accumulate if **compensatory mutations** are also occurring that maintain the overall function of the element.

The sequence variation seen in the *eve* elements is similar to those observed in other complex elements. A survey of eight *cis*-regulatory elements for early developmental genes in several *Drosophila* species revealed that among 104 transcription factor binding sites found in *D. melanogaster*, approximately 30–40% were not present in orthologous elements in *D. virilis*. The rate of turnover of binding sites was approximately 1% (gain or loss) of sites per million years.

Similarly, a survey of 51 human *cis*-regulatory elements found that approximately 32–40% of the human transcription factor binding sites are not present in the homologous elements of rodents, also indicating a high turnover rate in mammalian *cis*-regulatory elements. Together, these data suggest that binding site turnover is a general feature of *cis*-regulatory sequence evolution, and that the balancing effects of mutation and selection are likely to apply, in general, to the functional conservation of *cis*-regulatory elements.

Indeed, in some cases, it appears that *cis*-regulatory element function can be maintained despite nearly complete turnover of transcription factor binding sites. The regulation of the *Endo16* of the sea urchin *Strongylocentrotus purpuratus* has been analyzed in extensive detail, revealing a modular architecture of 5′ *cis*-regulatory elements (comprising seven modules named A–G). In *Lytechinus variegatus*, a species some 35 million years diverged from *S. purpuratus*, only the A module is conserved, the remaining elements cannot be identified by sequence similarity. Yet, the pattern of transcription of the *Endo16* gene during development is largely conserved. Experimental evidence suggests that the set of transcription factors that interact with the upstream elements of *Endo16* has also diverged. These findings suggest that orthologous genes expressed in homologous positions in two well-diverged species may not be regulated by homologous *cis*-regulatory sequences or transcription factors. This demonstrates both the remarkable potential fluidity of *cis*-regulatory DNA sequences, as well as the challenge of understanding the mechanisms of regulatory divergence between genes of long-diverged taxa.

**Co-evolution of *cis*-regulatory elements and the transcriptional factors that bind them**

The high rate of transcription factor binding site turnover suggests that compensation is not restricted to changes in *cis*-regulatory elements. Another means of compensation is the co-evolution of transcription factor specificities. Detailed examination of the evolution of elements regulated by the Bicoid protein has uncovered evidence for significant functional sequence evolution in the Bicoid protein in Diptera, apparently driven by the continuous process of enhancer restructuring.

One of the best characterized targets of the Bicoid protein is the P2 promoter of the *hunchback* gene (see Chapter 3, pp. 64–65). There are seven Bcd binding sites upstream of the *hb* P2 promoter in *D. melanogaster* (Fig. 8.5). The number, spacing, sequence, and orientation of Bcd sites differs in other Dipterans (Fig. 8.5), as does the sequence of the Bcd homeodomain. The differences in the *hb* enhancer and Bcd protein are of functional significance, because components combined from different species fail to interact as effectively as those
Figure 8.5
Evolution of Bcd binding sites in the hunchback genes of higher Diptera

The distribution of sites with TAAT core sequences (red hexagons) and different core sequences (blue ovals) in the upstream 1 kb of the hunchback genes of five species are shown. The number and topology of sites has diverged among species, and the Bcd protein has co-evolved with these regulatory elements.

from the same species. For example, the Bcd protein from *D. melanogaster* binds with a higher affinity to the *D. melanogaster* *hb* sequences than to the *Musca* *hb* sequences, and the *Musca* Bcd protein binds to the *Musca* *hb* with higher affinity than to the *D. melanogaster* *bg* element. Consistent with these observations, the *Drosophila* Bcd protein rescued embryos from *bcd* mothers more efficiently than did *Musca* Bcd. These data are consistent with compensatory coevolution occurring between the Bcd transcription factor and the *hh P2* promoter.

**Functional modification of cis-regulatory DNA sequences**

Many of the comparative studies presented in Chapters 5, 6, and 7 inferred a change in *cis*-regulatory function but, with the exception of Hoxc8 element evolution in vertebrates (see Fig. 5.7), the specific functional changes within elements have not been identified. Nevertheless, some well-analyzed cases of the evolution of tissue-specific *cis*-regulatory elements do illustrate the likely mechanisms underlying the functional modification of such elements of developmental regulatory genes.

Co-option or modification of *cis*-regulatory elements is the most likely explanation for the evolution of lens crystallins (a diverse group of water-soluble proteins that have been recruited to function in eye lenses in the refraction of light onto the retina). Various enzymes or proteins related to enzymes such as lactate dehydrogenase B, argininosuccinate lyase, α-enolase, glutathione-S-transferase, and small heat-shock proteins are expressed at very high levels in the lens in particular species. The recruitment of these proteins as crystallins has occurred independently in various lineages, suggesting that many gene co-option events have occurred during the evolution of lens crystallins.

Analyses of *cis*-regulatory elements of a wide variety of crystallin genes have revealed that certain transcription factors are frequently involved in high levels of protein expression in the lens. They include the Pax-6 and retinoic acid receptor proteins, which play major roles in eye development in most animal phyla. The implication of these proteins in crystallin gene regulation suggests a scenario for the molecular genetic basis of lens crystallin recruitment. Namely, the evolution of sites for Pax-6, retinoic acid receptors, and other transcription factors within extant *cis*-regulatory elements of these genes may lead to abundant levels of expression in developing eye tissue. In general, high levels of gene expression in a given cell type or tissue could reflect the evolution of sites for abundant transcription factors within extant *cis*-elements.

Tissue-specific losses of gene expression also occur. For example, blind mole rats have degenerated, nonfunctional eyes. In the superspecies *Spalax ehrenbergi*, the small heat shock protein/α-crystallin gene enhancer has selectively lost activity in the developing lens, while its activity in the heart and muscle have been conserved. This appears to be due to relatively few changes in the mole rat gene relative to other rodents.

A second, well-analyzed case involves the glucose dehydrogenase (*gld*) enzyme, which is expressed in particular reproductive tissues of many *Drosophila* species. In *D. teissieiri*, for example, *gld* expression does not occur in a subset of these tissues. Comparisons of the homologous *cis*-regulatory regions of the *gld* gene of seven different *Drosophila* species have revealed extensive sequence conservation between all seven genes, including the perfect conservation of certain motifs between all seven species. On the other hand, the *D. teissieiri* *gld* element lacks all copies of a sequence motif that occurs three times in a *D. melanogaster* *gld* regulatory element. This motif directs gene expression in *D. melanogaster* in the tissues
from which gld expression in *D. teissieiri* is absent. Thus evolutionary changes in the *D. teissieiri* gld element correlate with tissue-specific loss of gld expression in this lineage.

The gain or loss of gene expression can, in some cases, be as specific as involving a single cell. In the two nematode worms *C. elegans* and *C. briggsae*, the expression of the *lin-48* gene is shared in hindgut cells and neuronal support cells but is present in the excretory duct cell only in *C. elegans*. Examination of the respective *lin-48* cis-regulatory elements of both species suggests that this difference is due in part to the presence of binding sites for the CES-2 transcription factor in the *lin-48* element in *C. elegans* that were either gained in the *C. elegans* lineage or lost in the *C. briggsae* lineage.

**Variation in regulatory DNA and gene expression**

The evolutionary divergence in gene expression between species associated with morphological diversity, the turnover of binding sites in cis-regulatory DNA, and the association of some quantitative trait variation with noncoding DNA sequences all point towards the importance of variation in regulatory DNA sequences as a major component of morphological evolution. How widespread is functional variation in regulatory sequences in natural populations?

Variation in gene expression has also been examined at the level of individuals, populations, and closely related species. For example, in the fish *Fundulus heteroclitus*, 18% of loci surveyed differed in expression between individuals within a population. Among different strains of *D. melanogaster*, a range of 2–6% of genes surveyed exhibited differences in expression levels at a single developmental stage. Pairwise comparisons of *D. melanogaster* and two closely related species, *D. simulans* and *D. yakuba*, suggest that 10% of genes exhibit expression differences at a single stage, and that 27% of all genes differ in expression levels between at least two strains or species. Perhaps even more compelling, comparisons of gene expression profiles from the brains of humans, chimpanzees, orangutans, and macaques suggest that a substantial fraction of genes, perhaps a quarter, are expressed at different levels in different primates.

We are only beginning to directly survey nucleotide variation and its attendant effects on gene expression. Allelic variation in gene expression has been best studied thus far in mice and humans. In general, polymorphisms with affects on the order of 1.5–4-fold on the level of transcription have been found for as many as half of all genes surveyed. These findings suggest that regulatory variation is widespread within natural populations. Furthermore, small-scale nucleotide variation is sufficient to impart potentially functionally significant differences in levels of gene expression. Indeed, it may be that humans, for example, are heterozygous at more functional cis-regulatory sites than amino acid positions, underscoring the substantial potential of regulatory variation to underlie phenotypic variation.

The emerging picture, then, is that regulatory variation is abundant at the level of individuals, populations, and species. The future challenge will be to link this regulatory variation to discrete phenotypic characters.

**THE EVOLUTION OF REGULATORY DNA AND MORPHOLOGICAL DIVERSITY**

In this chapter, we have examined the evolution of regulatory DNA from three perspectives—theoretical considerations of cis-regulatory element function, comparative analyses of
known *cis*-regulatory elements, and a handful of genetic investigations into the nature of morphological variation. All of these approaches support the claim that regulatory DNA is the source of genetic diversity that underlies morphological diversity. The greater role of regulatory DNA in morphological evolution (as compared to coding DNA) is enabled by three critical factors:

- The degree of freedom in *cis*-regulatory sequences
- The modularity of *cis*-regulatory elements
- The combinatorial action of the transcription factor repertoire in cells

The degree of freedom in regulatory DNA is important because it imparts a greater tolerance of regulatory DNA to all types of mutational change. Regulatory DNA does not need to maintain any reading frame, and it can function at widely varying distances from or orientation to the transcription units it controls.

The modularity of the elements that make up the *cis*-regulatory systems of genes facilitates evolution because individual elements can evolve independently. The ability of regulatory DNA to readily evolve creates a rich source of genetic variation and, therefore, potential morphological variation.

The importance of the combinatorial nature of transcription regulation cannot be overemphasized. The transcription factor repertoire is sufficiently diverse and the stringency of DNA binding sufficiently relaxed such that sites for most transcription factors can evolve at a significant frequency in animal genomes. As new combinations of sites arise in existing functional elements, and potentially within nonfunctional DNA as well, variations in the timing, level, and spatial domains of gene expression may evolve and generate phenotypic variations, which serve as the raw material for selection and morphological change. The very structure of the *cis*-regulatory regions of toolkit genes—that is, its composition from multiple, independently regulated *cis*-elements—is the product of these evolutionary processes. It constitutes persuasive evidence that the diversification of regulatory DNA, while generally maintaining coding function, is the most available and most frequently exploited mode of genetic diversification in animal evolution.

**Extrapolating from bristles to body plans**

This book has explored and sought to explain the genetic basis of the morphological diversity of animals. Three kinds of assumptions are implicit to our approach and the conclusions we have drawn.

First, we assume that we can extrapolate from the present to the past. Ayshaiea (see Fig. 1.1a) and Acanthostega (see Fig. 1.5e) are no longer walking the Earth, but we assume that we can extrapolate from genetic and developmental knowledge of their modern descendants and thereby infer processes that occurred in the Cambrian or the Devonian periods.

Second, we assume that we can extrapolate from the particular to the general. Very few genetic regulatory circuits or networks are known in great detail, a modest number of *cis*-regulatory elements have been thoroughly dissected at the DNA sequence and transcription factor level and compared between relevant taxa. Nevertheless, we assert that the specific knowledge we do have is likely to apply to the general case.
Our third, and perhaps most controversial, assumption is that we may extrapolate from the observable small genetic changes underlying fine-scale morphological variation to the largest changes that have happened in animal history. Jacques Monod once asserted that what was true of the bacterium *Escherichia coli* was true of elephants. Analogously, we suggest that what is true of bristles is true of body plans. If dozens of regulatory changes and gene interactions underlie the difference of just a few bristles between fruit fly populations, then the number of genetic regulatory differences underlying the full range of morphological differences between a fly and a butterfly, or between a mouse and a human, must be staggeringly immense. But, do we need to invoke any additional or special genetic or evolutionary mechanisms beyond those illustrated for bristle variation to explain morphological diversity at higher levels? Is there any role, for example, for the sorts of dramatic variants described by Bateson in the evolution of large differences between taxa? The emerging evidence suggests not. Although, in theory, any morphological or genetic variation may potentially be selected for, Bateson’s monsters (for example, homeotic mutants) are generally less fit than other individuals and would be selected against in interbreeding populations. Furthermore, the case studies and mechanisms described in this book suggest that not only are such large steps improbable, but unnecessary. We have described many examples of morphological evolution that involved homeotic genes, but not homeotic mutations. Dramatic morphological changes, such as the fin-to-limb transition in vertebrates or the evolution of the insect haltere, involved many regulatory, developmental, and anatomical modifications that could not and did not evolve instantaneously. Instead, these structures were sculpted by regulatory evolution over millions of years.

Does variation exist in animal forms that might provide the basis for large-scale morphological evolution? Indeed, variants are found with significant frequency within and between natural populations for characters that are more significant to the evolution of body plans and body parts than just bristle number. For example, detailed analysis of a single population of newts has revealed a surprising array of variation in limb skeleton morphologies in approximately 30% of individuals examined (Fig. 8.6). Most interestingly, these variant patterns often resembled the standard limb skeletal morphologies found in other species. Some of these patterns were similar to more “ancient” patterns; others reflected more derived conditions. These observations suggest that within this single population of animals, many potential limb morphologies are expressed that represent both potential novelties and atavisms (return to an ancestral state). As the evolution of limb skeletal morphology is important to the functional adaptation of these and other tetrapods, the variation documented in this population suggests that the evolution of new forms or the reversion to “old” forms can arise through readily available genetic variation in the genes that affect limb morphology. Little, if any, morphological variation in this population is likely to be due to intraspecific variation in protein function.

Other well-documented examples of striking intraspecific morphological variation have been described in centipedes and certain fishes. Among geophilomorph centipede species, the number of leg-bearing segments varies from 29 to 191; in other centipede orders, this number remains relatively constant. Within geophilomorph species, there may be a range of variance of 12–14 segments. Given the considerable range in segment number observed in fossil and extant arthropods, the mechanisms underlying variation in such a central feature of the arthropod body plan are of immense interest.
Remarkable body pattern variation is also observed between populations of the threespine stickleback fish. This group of species has experienced repeated postglacial episodes of colonization of freshwater lakes and streams. Skull, body, and appendage shapes differ extensively between populations (Fig. 8.7) that have become isolated relatively recently (for example, circa 13,000 years ago). It appears that, in the stickleback, colonization and ecological diversification have led to a radiation of morphologically diverse, but closely related species.

These surveys of natural populations offer striking support for the idea that the morphological variation required for the evolution of new body patterns is available, at least in some groups. When this information is coupled with new insights into the scope of genetic variation that can lurk beneath the surface of phenotypic stability, it appears that the developmental,
genetic, and potential morphological diversity of any large interbreeding group is much greater than has been realized by biologists, or is realized in the course of evolution. This potential is generated from the creative power of regulatory change and interaction; and it is realized through ecological interactions at many levels.

We arrive at a view — while perhaps made more sophisticated by virtue of modern embryology, genetics, and molecular biology—that does not stray far from the spirit, if not the heart,
of Darwin’s original ideas. Darwin chose to open *The Origin of Species* with a discussion of domesticated species, arguing persuasively about the power of selection upon variation, on a scale and landscape familiar to his readers. Then, in closing his great work, he urged his audience to extrapolate from dog and pigeon breeding to the larger landscape of life’s history:

“we are always slow in admitting great changes of which we do not see the steps. . . . The mind cannot possibly grasp the full meaning of the term of even a million years; it cannot add up and perceive the full effects of many slight variations, accumulated during an almost infinite number of generations.”

Today, we are beginning to identify the genetic steps underlying the evolution of specific traits, and to conceive and reconstruct some of the innumerable steps underlying the great changes in animal designs that have unfolded from our Precambrian ancestors. We have been able to elucidate some of the general mechanisms that have been at work throughout animal history. Armed with increasingly more powerful tools for analyzing variation and comparing genomes and gene expression, the relationship between the evolution of DNA and animal diversity is drawing increasingly within our grasp.

**SELECTED READINGS**

**General Reviews**


**Regulatory Evolution: Early Ideas**


**Functions of cis-Regulatory Element and Transcription Factor Interactions**


Origins of cis-Regulatory Elements


Conservation and Turnover of Binding Sites in cis-Regulatory Elements


Divergence of cis-Regulatory Element Function


Coevolution of Binding Sites and Transcription Factors


**Regulatory Variation**


**Population Genetics of cis-Regulatory Elements**


**Morphological Variation within Species**


**Extrapolation from Microevolution to Macroevolution**


activator A protein that positively regulates transcription of a gene.

Annelida Phylum comprising about 15,000 species of segmented worms; includes polychaetes, earthworms, and leeches.

anterior group Hox genes Hox genes expressed in the anterior region of bilaterians and located toward the 5’ end of Hox clusters; include the Hox1 and Hox2 genes in vertebrates.

anural The tailless condition in chordates.

apical ectodermal ridge (AER) A thickening of the ectoderm at the distal extent of developing tetrapod limbs.

artificial selection Selection imposed by humans for one or more traits, usually in a laboratory, captive, or domesticated population.

atavism The evolutionary reversion of a character to an ancestral state.

autopod The distal portion of the vertebrate limb, including the hand/foot and digits.

axes The polarity of animals can be defined in three dimensions. In bilaterians, the two main axes are the anteroposterior (head-to-tail or rostrocaudal) axis and the orthogonal dorsoventral axis. Projections from the body wall, such as appendages, also possess a third proximodistal axis.

basal In phylogenetic terms, a group that branches at the base of a clade.

battery A group of genes regulated by the same transcription factor.

Bilateria The bilaterally symmetrical animals, including all protostomes and deuterostomes, but not sponges, cnidarians, or ctenophores.

blastopore An invagination on the surface of embryos into which the mesoderm and endoderm move during early embryogenesis.

Cambrian The geological period from 544 to 490 million years ago, during which the diversity of modern animal phyla expanded.

carapacial ridge A prominent outgrowth of ectoderm and mesoderm that arises on the dorsal flank of the turtle embryo and gives rise to the dorsal carapace (shell).

cell autonomous Genes whose products affect only the differentiation or behavior of cells in which they are expressed. Contrast with genes that encode signaling ligands that act on other cells (i.e., non-cell-autonomously).

central group Hox genes Hox genes expressed in the central region of bilaterians and located in the middle of Hox clusters; include the Hox3 through Hox8 genes in vertebrates.

cephalochordate Phylum composed of only approximately 25 marine species characterized by a notochord and small anterior brain vesicle. Also known as the lancelets, due to their body shape, they constitute the nearest phylogenetic outgroup to the vertebrates.

chromatin The material into which DNA in the cell nucleus is packaged with proteins.

circuit A regulatory pathway that includes particular target genes.

cis-regulatory element A discrete region of DNA that affects transcription of a gene.

clad A group of species descended from a common ancestral species. Also known as a monophyletic group.
Cnidaria  Phylum comprising approximately 8000 living, mostly marine species, all of which possess nematocyst cells. Their diploblastic, radially symmetric body organization consists of an outer cell layer (ectoderm) and an inner cell layer (endoderm). Cnidaria includes jellyfish, sea anemones, and corals.

coadministrator  Protein that interacts with the activator in control of transcription. Some co-activators possess activities that modify local chromatin conformation.

coeleom  A body cavity formed from the mesoderm that lies between the gut (endoderm) and the body wall (ectoderm) in many triploblast animals.

colinearity  The correlation between the order of Hox genes on a chromosome and the anteroposterior order of gene deployment in the embryo.

combinatorial regulation  The control of gene transcription by two or more transcription factors. The spatial patterns of gene expression are often delimited by the combined action of transcription factors.

compartment  A discrete subdivision of a developing field that contains populations of cells that share a lineage restriction. Cells of one compartment do not intermix with adjacent cells in other compartments.

compensatory mutation  A mutation that restores or maintains the function of a gene or cis-regulatory element by counteracting the effects of one or more different mutations.

cooperativity  The enhanced binding of a protein to a cis-regulatory DNA sequence due to interaction with other bound proteins.

co-option  The recruitment of genes into new developmental or biochemical functions.

corepressor  A protein that interacts with a repressor in the control of transcription. Corepressors may modify the local conformation of chromatin.

cryptic genetic variation  The genetic variation in components of developmental programs that exists in the absence of phenotypic variation. Perturbations to developmental programs may unmask this variation and lead to phenotypic variation.

deuterostomes  The group of animals including echinoderms, ascidians, cephalochordates, and vertebrates in which the mouth forms at a site separate from the blastopore.

Devonian  The geological period from approximately 409 to 362 million years ago, during which terrestrial forms invaded the land.

diploblasts  Animals possessing two cell layers—the outer ectoderm layer and the inner endoderm layer; include cnidarians.

disparity  The variety of designs in body plans.

diversity  The number of species in a clade; the divergence of morphology.

echinoderm  A deuterostome animal with a pentaradially symmetrical adult body plan.

ectoderm  The outer tissue layer of animal embryos that gives rise to the epidermis and the nervous system.

Ediacara  The fauna that lived 575–544 million years ago, named for their fossil deposits in the hills of South Australia.

endoderm  The inner tissue layer of animal embryos that gives rise to the gut and other associated organs.

exon shuffling  The process through which new genes are generated by recombination of one or more exons of other genes.

extant  Living, as opposed to extinct, species.

field-specific selector gene  A specific class of selector genes that control the formation and patterning of morphogenetic fields such as the eye, leg, and wing.

focus  The developmental organizer at the center of the butterfly eyespot field.

gap genes  The genes that subdivide the *Drosophila* embryo into several regions encompassing many segments. They encode transcription factors that regulate the expression of other gap genes, pair-rule genes, and homeotic genes.

gene complex  A group of adjacent genes related by gene duplication.

gene divergence  The process through which two or more genes created by a duplication event(s) acquire distinct functions. Changes in regulatory and/or coding regions lead to differences in gene function.

gene duplication  The creation of additional genes from the template of one gene.

gene family  Two or more genes with related sequences that are derived from a common ancestral gene. They are not necessarily linked, may be widely dispersed in the genome, and may be greatly diverged in function.
**genetic architecture** Refers to the number and interactions among genes whose products govern a particular trait.

**genetic redundancy** Two genes are redundant when the absence of the function of one gene causes little or no effect because a second gene can operate in its place.

**germ layers** The regions of early animal embryos that give rise to the different tissue layers. Bilaterians have three germ layers, which form the ectoderm, mesoderm, and endoderm.

**gradient** Formed when the concentration of a regulatory substance varies with the distance from a source.

**haltere** The hindwing of two-winged insects such as flies and mosquitoes.

**homeodomain** The protein domain encoded by the homeobox.

**homeotic genes** Genes that regulate the identity of body regions. Mutations in homeotic genes cause the transformation of one body region or part into the likeness of another.

**homolog** When referring to genes, indicates genes that share similarities in their sequence because they evolved from a common ancestral gene. When referring to morphological structures, indicates features that are shared due to common ancestry.

**homonomous** A body plan or body part composed of repeating, similar parts.

**housekeeping genes** Genes that encode proteins required for basic functions required in all cells.

**Hox genes** Homeotic, homeobox-containing genes found in linked clusters in all bilaterians.

**imaginal disc** Small sets of epithelial cells that are set aside during *Drosophila* embryogenesis, subsequently proliferate in the larva, and give rise to adult structures.

**individualization/individuation** The process of morphological differentiation of similar body parts.

**lateral plate mesoderm** Mesoderm in vertebrate embryos that gives rise to the blood, kidneys, and heart tissues.

**leucine zipper** A common transcription factor motif in which two subunits of the protein interact through leucine-containing repeat loops.

**Lophotrochozoa** A great clade of protostomes including the annelids, molluscs, flatworms, and other phyla; many phyla within this group produce a particular form of larva (trochophore) or feeding structure (lophophore).

**maternal effect** Genes that are expressed during oogenesis and whose products function in the development of the embryo.

**meristic variation** Differences in the number of repeating structures.

**mesoderm** The tissue layer between the ectoderm and the endoderm that gives rise to the musculature of all bilaterians.

**Metazoa** Multicellular animals, including diploblasts and triploblasts.

**modularity** The organization of animals into developmentally and anatomically distinct parts. Also, the organization of gene regulatory regions into discrete *cis*-regulatory elements.

**molecular clock** Differences in the sequences of RNA, DNA, and protein molecules accumulate as species diverge. In theory, geological information can be used to calibrate the rate of sequence changes between lineages. The relative sequence divergence between species can be used to infer phylogenetic relationships.

**morphogen** A substance whose concentration varies across a tissue or field and to which cells respond differently at different concentrations.
mophogenetic field A discrete region of an embryo that will give rise to a structure and within which pattern formation is largely independent of other developing structures.

network Two or more regulatory circuits that are linked by regulatory interactions between components of each circuit.

neural crest A population of cells in vertebrates that migrates from the edge of the neural plate to different regions of the body and gives rise to tissues such as the facial skeleton, melanocytes, and peripheral neurons.

notochord A structure that runs from head to tail in chordates and lies beneath the developing central nervous system.

novelty New morphological characters of adaptive value.

onychophora A phylum of soft-bodied animals possessing unjointed appendages; closely related to the arthropods.

oral–aboral axis The axis orthogonal to the pentaradial arms of echinoderms.

Ordovician The geological period from approximately 490 to 439 million years ago, which followed the Cambrian and preceded the Devonian.

organizer A signaling center in a developing embryo or field that induces the development of surrounding tissues. Examples include the Spemann organizer of amphibia embryos and the focus in butterfly imaginal wing discs.

orthologs Homologous genes in different species that arose from a single gene in the last common ancestor of these species.

outgroup A taxon that diverged from a group of other taxa before members of that group subsequently diverged from one another.

pair-rule genes Drosophila genes that are usually expressed in seven transverse stripes across the future segmented region of the early embryo, one stripe for every two segment primordia.

Parahox genes A small cluster of genes that arose from an early duplication of a primitive Proto-Hox gene cluster. Parahox genes function in endoderm development and may be involved in the evolution of the through gut in bilaterians.

paralogs Homologous genes that are related by duplication of an ancestral gene.

parasegment Units of the Drosophila embryo composed of the posterior part of one segment and the anterior part of the adjacent segment.

paraxial mesoderm That part of the vertebrate mesoderm that gives rise to the somites.

pathway In cell–cell signaling, the components required for the sending, receiving, and transduction of a signal, including one or more ligands, membrane-associated receptors, intracellular signal transducers, and transcription factors.

phenocopy A mimic of a genetic mutant phenotype, usually caused by environmental changes.

phylogenetic tree A depiction of the evolutionary relationships among species and their order of “branching” from common ancestors.

pleiotropic Genes or proteins with multiple functions.

polychaete One class of annelids distinguished by the presence of chaetae, projections from the sides of each body segment.

Porifera Phylum containing the sponges, the most primitive animal group that does not contain true germ layers.

posterior group Hox genes Hox genes that are expressed in the posterior or caudal region of bilaterians and are found at the 5′ end of Hox clusters; include the Abd-B-related and Hox9-13 genes.

primordia A discrete field of cells that will give rise to a particular organ, appendage, or tissue type.

proleg The leg-like abdominal appendages found on the larva of certain insects, particularly Lepidoptera.

promoter The region of a gene near the start site of transcription to which the general transcriptional machinery binds.

proneural cluster A group of cells from which one or more neural precursor cells will segregate, divide, and differentiate.

Proterozoic The geological era from approximately 2500 to 544 million years ago.

protostomes One of two clades of bilaterians characterized by the origin of the mouth at the blastopore.

pseudogene The remnant of a gene that has been rendered nonfunctional through the accumulation of mutations.
quantitative trait A character that exhibits continuous variation in a population.

quantitative trait loci (QTL) Genetic loci that contribute to the variation in a quantitative trait in a population.

radiation The evolutionary divergence of a lineage into a variety of forms; often used to describe rapid diversification.

region-specific selector One class of selector genes that regulates the identity of contiguous body regions (e.g., *Hox* genes).

regulatory circuit A signaling pathway and one or more of the target genes regulated by the pathway in a given cell, tissue, or field.

regulatory evolution Evolutionary changes in gene regulation.

reporter gene A gene whose expression is used to visualize the activity of a heterologous, linked *cis*-regulatory element in vivo.

repressor A transcription factor that negatively regulates the expression of a gene, often by binding directly to DNA sequences in a *cis*-regulatory element.

rhombomere A subdivision of the vertebrate hindbrain.

secondary field Discrete units of development that are specified in the developing embryo after the primary axes are established and that give rise to appendages and organs. Pattern formation within secondary fields occurs independently of other fields.

segment polarity gene Genes that act within developing *Drosophila* segments to regulate the anteroposterior polarity of each segment.

selector gene A gene that controls cell fate.

serial homologs Repeated structures of a single organism that share a similar developmental origin.

somite A segmented subdivision of the vertebrate mesoderm that gives rise to vertebrae and associated processes, selected muscles, and the dermis.

stabilizing selection Selection that acts to keep a character constant in a population.

stem lineage Generally refers to fossil taxa that possess some, but not all, of the characters that define a later group.

syncytium An embryo or tissue containing nuclei that are not separated by cell membranes.

synteny Similarity in the linkage order of orthologous genes between two or more species.

tetraploidization A process of whole-genome duplication caused by doubling the number of chromosomes per cell from diploid (two copies) to tetraploid (four copies).

tetrapod The four-limbed terrestrial classes of vertebrates, including amphibians, reptiles, birds, and mammals.

trace fossil The impressions left in sediments by the meanderings and burrowing activities of ancient animals.

transcription factor A protein that regulates the transcription of genes, often, but not exclusively, by binding to *cis*-regulatory elements.

triploblasts Animals composed of three germ layers; includes all bilaterians.

Urbilateria The hypothetical last common ancestor of all bilaterians.

urochordate A marine phylum in which the notochord (or urochord) is found only in the larval tail.

urodele The tailed form of chordates.

zinc finger A distinct class of transcription factors in which a DNA-binding polypeptide loop or “finger” forms through a coordination complex of zinc with cysteine and histidine residues at the base of the loop.

zone of polarizing activity (ZPA) The organizer at the posterior margin of the developing vertebrate limb that regulates anteroposterior polarity.

zygote The fertilized egg. It contains the genetic contributions of both the male and female parents.
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