9. The genetics of axis specification in Drosophila

Thanks largely to the studies by Thomas Hunt Morgan's laboratory during the first decade of the twentieth century, we know more about the genetics of Drosophila than about any other multicellular organism. The reasons for this have to do with both the flies and the people who first studied them. Drosophila is easy to breed, hardy, prolific, tolerant of diverse conditions, and the polytene chromosomes of its larvae are easy to identify (see Chapter 4). The techniques for breeding and identifying mutants are easy to learn. Moreover, the progress of Drosophila genetics was aided by the relatively free access of every scientist to the mutants and the techniques of every other researcher. Mutants were considered the property of the entire scientific community, and Morgan's laboratory established the database and exchange network whereby anyone could obtain them.

Undergraduates (starting with Calvin Bridges and Alfred Sturtevant) played important roles in Drosophila research, which achieved its original popularity as a source of undergraduate research projects. As historian Robert Kohler noted (1994), "Departments of biology were cash poor but rich in one resource: cheap, eager, renewable student labor." The Drosophila genetics program was "designed by young persons to be a young person's game," and the students set the rules for Drosophila research: "No trade secrets, no monopolies, no poaching, no ambushes."

But Drosophila was a difficult organism on which to study embryology. Although Jack Schultz and others attempted to relate the genetics of Drosophila to its development, the fly embryos proved too complex and too intractable to study, being neither large enough to manipulate experimentally nor transparent enough to observe. It was not until the techniques of molecular biology allowed researchers to identify and manipulate the genes and RNAs of the insect that its genetics could be related to its development. And when that happened, a revolution occurred in the field of biology. The merging of our knowledge of the molecular aspects of Drosophila genetics with our knowledge of its development built the foundations on which are current sciences of developmental genetics and evolutionary developmental biology are based.

Early Drosophila Development

In the last chapter, we discussed the specification of early embryonic cells by their acquisition of different cytoplasmic determinants that had been stored in the oocyte. The cell membranes establish the region of cytoplasm incorporated into each new blastomere, and it is thought that the morphogenetic determinants then direct differential gene expression in these blastomeres. During Drosophila development, however, cellular membranes do not form until after the thirteenth nuclear division. Prior to this time, all the nuclei share a common cytoplasm,
and material can diffuse throughout the embryo. In these embryos, the specification of cell types along anterior-posterior and dorsal-ventral axes is accomplished by the interactions of cytoplasmic materials within the single, multinucleated cell. Moreover, the initiation of the anterior-posterior and dorsal-ventral differences is controlled by the position of the egg within the mother's ovary. Whereas the sperm entry site may fix the axes in ascidians and nematodes, the fly's anterior-posterior and dorsal-ventral axes are specified by interactions between the egg and its surrounding follicle cells.

**Cleavage**

Most insect eggs undergo superficial cleavage, wherein a large mass of centrally located yolk confines cleavage to the cytoplasmic rim of the egg. One of the fascinating features of this cleavage type is that cells do not form until after the nuclei have divided. Cleavage in a *Drosophila* egg is shown in Figure 9.1. The zygote nucleus undergoes several mitotic divisions within the central portion of the egg. In *Drosophila*, 256 nuclei are produced by a series of eight nuclear divisions averaging 8 minutes each. The nuclei then migrate to the periphery of the egg, where the mitoses continue, albeit at a progressively slower rate. During the ninth division cycle, about five nuclei reach the surface of the posterior pole of the embryo. These nuclei become enclosed by cell membranes and generate the pole cells that give rise to the gametes of the adult. Most of the other nuclei arrive at the periphery of the embryo at cycle 10 and then undergo four more divisions at progressively slower rates.

During these stages of nuclear division, the embryo is called a syncytial blastoderm, meaning that all the cleavage nuclei are contained within a common cytoplasm. No cell membranes exist other than that of the egg itself.

Although the nuclei divide within a common cytoplasm, this does not mean that the cytoplasm is itself uniform. Karr and Alberts (1986) have shown that each nucleus within the syncytial blastoderm is contained within its own little territory of cytoskeletal proteins. When the nuclei reach the periphery of the egg during the tenth cleavage cycle, each nucleus becomes surrounded by microtubules and microfilaments. The nuclei and their associated cytoplasmic islands are called energids. Figure 9.2 shows the nuclei and their essential microfilament and microtubule domains in prophase of the twelfth mitotic division.
Following cycle 13, the oocyte plasma membrane folds inward between the nuclei, eventually partitioning off each somatic nucleus into a single cell (Figure 9.3). This process creates the cellular blastoderm, in which all the cells are arranged in a single-layered jacket around the yolk core of the egg (Turner and Mahowald 1977; Foe and Alberts 1983). Like any other cell formation, the formation of the cellular blastoderm involves a delicate interplay between microtubules and microfilaments. The first phase of blastoderm cellularization is characterized by the invagination of cell membranes and their underlying actin microfilament network into the regions between the nuclei to form furrow canals.

This process can be inhibited by drugs that block microtubules. After the furrow canals have passed the level of the nuclei, the second phase of cellularization occurs. Here, the rate of invagination increases, and the actin-membrane complex begins to constrict at what will be the basal end of the cell (Schejter and Wieschaus 1993; Foe et al. 1993). In Drosophila, the cellular blastoderm consists of approximately 6000 cells and is formed within 4 hours of fertilization.

The midblastula transition

After the nuclei reach the periphery, the time required to complete each of the next four divisions becomes progressively longer. While cycles 1–10 are each 8 minutes long, cycle 13, the last cycle in the syncytial blastoderm, takes 25 minutes to complete. Cycle 14, in which the Drosophila embryo forms cells (i.e., after 13 divisions), is asynchronous. Some groups of cells complete this cycle in 75 minutes, whereas other groups of cells take 175 minutes (Figure 9.4; Foe 1989).
Transcription from the nuclei (which begins around the eleventh cycle) is greatly enhanced at this stage. This slowdown of nuclear division and the concomitant increase in RNA transcription is often referred to as the midblastula transition (see Chapter 8). Such a transition is also seen in the embryos of numerous vertebrate and invertebrate phyla. The control of this mitotic slowdown (in *Xenopus*, sea urchin, starfish, and *Drosophila* embryos) appears to be effected by the ratio of chromatin to cytoplasm (Newport and Kirschner 1982; Edgar et al. 1986a). Edgar and his colleagues compared the early development of wild-type *Drosophila* embryos with that of a haploid mutant. These haploid *Drosophila* embryos have half the wild-type quantity of chromatin at each cell division. Hence a haploid embryo at the eighth cell cycle has the same amount of chromatin that a wild-type embryo has at cell cycle 7. The investigators found that whereas wild-type embryos formed their cellular blastoderm immediately after the thirteenth division, the haploid embryos underwent an extra, fourteenth, division before cellularization. Moreover, the lengths of cycles 11–14 in wild-type embryos corresponded to those of cycles 12–15 in the haploid embryos. Thus, the haploid embryos follow a pattern similar to that of the wild-type embryos—only they lag by one cell division.

**Gastrulation**

At the time of midblastula transition, gastrulation begins. The first movements of *Drosophila* gastrulation segregate the presumptive mesoderm, endoderm, and ectoderm. The prospective mesoderm—about 1000 cells constituting the ventral midline of the embryo—folds inward to produce the **ventral furrow** (Figure 9.5). This furrow eventually pinches off from the surface to become a ventral tube within the embryo. It then flattens to form a layer of mesodermal tissue beneath the ventral ectoderm. The prospective endoderm invaginates as two pockets at the anterior and posterior ends of the ventral furrow. The pole cells are internalized along with the endoderm. At this time, the embryo bends to form the cephalic furrow.

The ectodermal cells on the surface and the mesoderm undergo convergence and extension, migrating toward the ventral midline to form the **germ band**, a collection of cells along the ventral midline that includes all the cells that will form the trunk of the embryo. The germ band extends posteriorly and, perhaps because of the egg case, wraps around the top (dorsal) surface of the embryo (Figure 9.5D).
Thus, at the end of germ band formation, the cells destined to form the most posterior larval structures are located immediately behind the future head region. At this time, the body segments begin to appear, dividing the ectoderm and mesoderm. The germ band then retracts, placing the presumptive posterior segments into the posterior tip of the embryo (Figure 9.5E).

While the germ band is in its extended position, several key morphogenetic processes occur: organogenesis, segmentation, and the segregation of the imaginal discs* (Figure 9.5e). In addition, the nervous system forms from two regions of ventral ectoderm. As described in Chapter 6, neuroblasts differentiate from this neurogenic ectoderm within each segment (and also from the nonsegmented region of the head ectoderm). Therefore, in insects like Drosophila, the nervous system is located ventrally, rather than being derived from a dorsal neural tube as in vertebrates.

The general body plan of Drosophila is the same in the embryo, the larva, and the adult, each of which has a distinct head end and a distinct tail end, between which are repeating segmental units (Figure 9.7). Three of these segments form the thorax, while another eight segments form the abdomen. Each segment of the adult fly has its own identity. The first thoracic segment, for example, has only legs; the second thoracic segment has legs and wings; and the third thoracic segment has legs and halteres (balancers). Thoracic and abdominal segments can also be distinguished from each other by differences in the cuticle. How does this pattern arise?

During the past decade, the combined approaches of molecular biology, genetics, and embryology have led to a detailed model describing how a segmented pattern is generated along the anterior-posterior axis and how each segment is differentiated from the others.

The anterior-posterior and dorsal-ventral axes of Drosophila form at right angles to one another, and they are both determined by the position of the oocyte within the follicle cells of the ovary. The rest of this chapter is divided into three main parts. The first part concerns how the anterior-posterior axis is specified and how it determines the identity of each segment. The second part concerns how the dorsal-ventral axis is specified by the interactions between the oocyte and its surrounding follicle cells. The third part concerns how embryonic tissues are specified to become particular organs by their placement along these two axes.
*Imaginal discs are those cells set aside to produce the adult structures. The details of imaginal disc differentiation will be discussed in Chapter 18. For more information on Drosophila developmental anatomy, see Bate and Martinez-Arias 1993; Tyler and Schetzer 1996; and Schwalm 1997.

The Origins of Anterior-Posterior Polarity

The anterior-posterior polarity of the embryo, larva, and adult has its origin in the anterior-posterior polarity of the egg (Figure 9.8). The maternal effect genes expressed in the mother's ovaries produce messenger RNAs that are placed in different regions of the egg. These messages encode transcriptional and translational regulatory proteins that diffuse through the syncytial blastoderm and activate or repress the expression of certain zygotic genes. Two of these proteins, Bicoid and Hunchback, regulate the production of anterior structures, while another pair of maternally specified proteins, Nanos and Caudal, regulates the formation of the posterior parts of the embryo. Next, the zygotic genes regulated by these maternal factors are expressed in certain broad (about three segments wide), partially overlapping domains. These genes are called gap genes (because mutations in them cause gaps in the segmentation pattern), and they are among the first genes transcribed in the embryo.

Differing concentrations of the gap gene proteins cause the transcription of pair-rule genes, which divide the embryo into periodic units. The transcription of the different pair-rule genes results in a striped pattern of seven vertical bands perpendicular to the anterior-posterior axis. The pair-rule gene proteins activate the transcription of the segment polarity genes, whose mRNA and protein products divide the embryo into 14 segment-wide units, establishing the periodicity of the embryo. At the same time, the protein products of the gap, pair-rule, and segment polarity genes interact to regulate another class of genes, the homeotic selector genes, whose transcription determines the developmental fate of each segment.

The Maternal Effect Genes

*Embryological evidence of polarity regulation by oocyte cytoplasm*
Classic embryological experiments demonstrated that there are at least two "organizing centers" in the insect egg, one in the anterior of the egg and one in the posterior. For instance, Klaus Sander (1975) found that if he ligated the egg early in development, separating the anterior from the posterior region, one half developed into an anterior embryo and one half developed into a posterior embryo, but neither half contained the middle segments of the embryo. The later in development the ligature was made, the fewer middle segments were missing. Thus, it appeared that there were indeed gradients emanating from the two poles during cleavage, and that these gradients interacted to produce the positional information determining the identity of each segment. Moreover, when the RNA of the anterior of insect eggs was destroyed (by either ultraviolet light or RNase), the resulting embryos lacked a head and thorax. Instead, these embryos developed two abdomens and telsons (tails) with mirror-image symmetry: telson-abdomen-abdomen-telson (Figure 9.9; Kalthoff and Sander 1968; Kandler-Singer and Kalthoff 1976). Thus, Sander's laboratory postulated the existence of a gradient at both ends of the egg, and hypothesized that the egg sequesters an RNA that generates a gradient of anterior-forming material.

The molecular model: protein gradients in the early embryo

In the late 1980s, the gradient hypothesis was united with a genetic approach to the study of *Drosophila* embryogenesis. If there were gradients, what were the morphogens whose concentrations changed over space? What were the genes that shaped these gradients? And did these morphogens act by activating or inhibiting certain genes in the areas where they were...
concentrated? Christiane Nüsslein-Volhard led a research program that addressed these questions. The researchers found that one set of genes encoded gradient morphogens for the anterior part of the embryo, another set of genes encoded morphogens responsible for organizing the posterior region of the embryo, and a third set of genes encoded proteins that produced the terminal regions at both ends of the embryo (Figure 9.10; Table 9.1). This work resulted in a Nobel Prize for Nüsslein-Volhard and her colleague, Eric Wieschaus, in 1995.

Table 9.1. Maternal effect genes that effect the anterior-posterior polarity of the Drosophila embryo

<table>
<thead>
<tr>
<th>Gene</th>
<th>Mutant phenotype</th>
<th>Proposed function and structure</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>AnterioGroup</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>bicoid (bcd)</td>
<td>Head and thorax deleted, replaced by Graded anterior morphogen; contains homeodomain; inverted telson</td>
<td></td>
</tr>
<tr>
<td>exuperantia (exu)</td>
<td>Anterior head structures deleted</td>
<td>Anchors bicoid mRNA</td>
</tr>
<tr>
<td>swallow (swa)</td>
<td>Anterior head structures deleted</td>
<td>Anchors bicoid mRNP</td>
</tr>
<tr>
<td><strong>PosterioGroup</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>nanos (nos)</td>
<td>No abdomen</td>
<td>Posterior morphogen; represses hunchback</td>
</tr>
<tr>
<td>tudor (tud)</td>
<td>No abdomen, no pole cells</td>
<td>Localization of Nanos</td>
</tr>
<tr>
<td>oskar (osk)</td>
<td>No abdomen, no pole cells</td>
<td>Localization of Nanos</td>
</tr>
<tr>
<td>vasa (vas)</td>
<td>No abdomen, no pole cells; oogenesis defective</td>
<td></td>
</tr>
<tr>
<td>valois (val)</td>
<td>No abdomen, no pole cells; Stabilization of the Nanos localization complex</td>
<td></td>
</tr>
<tr>
<td>pumilio (pum)</td>
<td>No abdomen</td>
<td>Helps Nanos protein bind hunchback message</td>
</tr>
<tr>
<td>caudal (cad)</td>
<td>No abdomen</td>
<td>Activates posterior terminal genes</td>
</tr>
<tr>
<td><strong>TerminalGroup</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>torso (tor)</td>
<td>No termini</td>
<td>Possible morphogen for termini</td>
</tr>
<tr>
<td>trunk (trk)</td>
<td>No termini</td>
<td>Transmits Torsolike signal to Torso</td>
</tr>
<tr>
<td>fs(1)Nasrat[U]</td>
<td>No termini; collapsed eggs</td>
<td>Transmits Torsolike signal to Torso</td>
</tr>
<tr>
<td>fs(1)polehole[U]</td>
<td>No termini; collapsed eggs</td>
<td>Transmits Torsolike signal to Torso</td>
</tr>
</tbody>
</table>

*Source: After Anderson 1989.*

The anterior-posterior axis of the Drosophila embryo appears to be patterned before the nuclei even begin to function. The nurse cells of the ovary deposit mRNAs in the developing oocyte, and these mRNAs are apportioned to different regions of the cell. In particular, four maternal messenger RNAs are critical to the formation of the anterior-posterior axis:

• **bicoid** and **hunchback** mRNAs, whose protein products are critical for head and thorax formation

• **nanos** and **caudal** mRNAs, whose protein products are critical for the formation of the abdominal segments

The **bicoid** mRNAs are located in the anterior portion of the unfertilized egg, and are tethered to the anterior microtubules. The **nanos** messages are bound to the cytoskeleton in the
posterior region of the unfertilized egg. The hunchback and caudal mRNAs are distributed throughout the oocyte. Upon fertilization, these mRNAs can be translated into proteins. At the anterior pole, the bicoid RNA is translated into Bicoid protein, which forms a gradient highest at the anterior. At the posterior pole, the nanos message is translated into Nanos protein, which forms a gradient highest at the posterior. Bicoid protein inhibits the translation of the caudal RNA, allowing Caudal protein to be synthesized only in the posterior of the cell. Conversely, Nanos protein, in conjunction with Pumilio protein, binds to hunchback RNA, preventing its translation in the posterior portion of the embryo. Bicoid also elevates the level of Hunchback protein in the anterior of the embryo by binding to the enhancers of the hunchback gene and stimulating its transcription. The result of these interactions is the creation of four protein gradients in the early embryo (Figure 9.11):
• An anterior-to-posterior gradient of Bicoid protein
• An anterior-to-posterior gradient of Hunchback protein
• A posterior-to-anterior gradient of Nanos protein
• A posterior-to-anterior gradient of Caudal protein

The Bicoid, Hunchback, and Caudal proteins are transcription factors whose relative concentrations can activate or repress particular zygotic genes. The stage is now set for the activation of zygotic genes in those nuclei that were busily dividing while this gradient was being established.

Evidence that the bicoid gradient constitutes the anterior organizing center

In *Drosophila*, the phenotype of the bicoid mutant provides valuable information about the function of gradients. Instead of having anterior structures (acron, head, and thorax) followed by abdominal structures and a telson, the structure of the bicoid mutant is telson-abdomen-abdomen-telson (Figure 9.12).

It would appear that these embryos lack whatever substances are needed for the formation of anterior structures. Moreover, one could hypothesize that the substance that these mutants lack is the one postulated by Sander and Kalthoff to turn on genes for the anterior structures and turn off genes for the telson structures (compare Figures 9.9 and 9.12).
Further studies have strengthened the view that the product of the wild-type \textit{bicoid} gene is the morphogen that controls anterior development. First, \textit{bicoid} is a maternal effect gene. Messenger RNA from the mother's \textit{bicoid} genes is placed in the embryo by the mother's ovarian cells (Figure 9.13; Frigerio et al. 1986; ABerleth et al. 1988).

The \textit{bicoid} RNA is strictly localized in the anterior portion of the oocyte (Figure 9.13B), where the anterior cytoskeleton anchors it through the message’s 3’ untranslated region (Ferrandon et al. 1997; Macdonald and Kerr 1998). This mRNA is dormant until fertilization, at which time it receives a longer polyadenylate tail and can be translated. Driever and Nüsslein-Volhard (1988b) have shown that when Bicoid protein is translated from this RNA during early cleavage, it forms a gradient, with the highest concentration in the anterior of the egg and the lowest in the posterior third of the egg. Moreover, this protein soon becomes concentrated in the embryonic nuclei in the anterior portion of the embryo (Figure 9.13C-D; see also Figure 5.35).

Further evidence that Bicoid protein is the anterior morphogen came from experiments that altered the steepness of the gradient. Two genes, \textit{exuperantia} and \textit{swallow}, are responsible for keeping the \textit{bicoid} message at the anterior pole of the egg. In their absence, the \textit{bicoid} message diffuses farther into the posterior of the egg, and the gradient of Bicoid protein is less steep (Driever and Nüsslein-Volhard 1988a). The phenotype produced by these two mutants is similar to that of \textit{bicoid}-deficient embryos, but less severe. These embryos lack their most anterior structures and have an extended mouth and thoracic region. Thus, by altering the gradient of Bicoid protein, one correspondingly alters the fate of the embryonic regions.
Confirmation that the Bicoid protein is crucial for initiating head and thorax formation came from experiments in which purified bicoid RNA was injected into early-cleavage embryos (Figure 9.14; Driever et al. 1990). When injected into the anterior of bicoid-deficient embryos (whose mothers lacked bicoid genes), the bicoid RNA rescued the embryos and caused them to have normal anterior-posterior polarity. Moreover, any location in an embryo where the bicoid message was injected became the head. If bicoid RNA was injected into the center of an embryo, that middle region became the head, and the regions on either side of it became thorax structures. If a large amount of bicoid RNA was injected into the posterior end of a wild-type embryo (with its own endogenous bicoid message in its anterior pole), two heads emerged, one at either end.

The next question then emerged: How might a gradient in Bicoid protein control the determination of the anterior-posterior axis? Recent evidence suggests that Bicoid acts in two ways to specify the anterior of the Drosophila embryo. First, it acts as a repressor of posterior formation. It does this by binding to and suppressing the translation of caudal RNA, which is found throughout the egg and early embryo. The Caudal protein is critical in specifying the posterior domains of the embryo, and it activates the genes responsible for the invagination of the hindgut (Wu and Lengyel 1998). The Bicoid protein binds to a specific region of the caudal message's 3′ untranslated region, thereby preventing the translation of this message in the anterior section of the embryo (Figure 9.15; Dubnau and Struhl 1996; Rivera-Pomar et al. 1996). This suppression is necessary, for if Caudal protein is made in the anterior, the head and thorax are not properly formed.

Second, the Bicoid protein functions as a transcription factor. Bicoid protein enters the nuclei of early-cleavage embryos, where it activates the hunchback gene. The transcription of hunchback is seen only in the anterior half of the embryo—the region where Bicoid protein is found. Mutants deficient in maternal and zygotic Hunchback protein lack mouthparts and thorax structures. In the late 1980s, two laboratories independently demonstrated that Bicoid protein binds to and activates the hunchback gene (Driever and Nüsslein-Volhard 1989; Struhl et al. 1989). The Hunchback protein derived from the synthesis of new hunchback mRNA joins the Hunchback protein synthesized by the translation of maternal messages in the anterior of the embryo. The Hunchback protein, also a transcription factor, represses abdominal-specific genes, thereby allowing the region of hunchback expression to form the head and thorax.
The Hunchback protein also works with Bicoid in generating the anterior pattern of the embryo. Based on two pieces of evidence, Driever and co-workers (1989) predicted that at least one other anterior gene besides hunchback must be activated by Bicoid. First, deletions of hunchback produce only some of the defects seen in the bicoid mutant phenotype. Second, as we saw in the swallow and exuperantia experiments, only moderate levels of Bicoid protein are needed to activate thorax formation (i.e., hunchback gene expression), but head formation requires higher concentrations. Driever and co-workers (1989) predicted that the promoters of such a head-specific gap gene would have low-affinity binding sites for Bicoid protein. This gene would be activated only at extremely high concentrations of Bicoid protein—that is, near the anterior tip of the embryo. Since then, three gap genes of the head have been discovered that are dependent on very high concentrations of Bicoid protein for their expression (Cohen and Jürgens 1990; Finkelstein and Perrimon 1990; Grossniklaus et al. 1994). The buttonhead, empty spiracles, and orthodenticle genes are needed to specify the progressively anterior regions of the head. In addition to needing high Bicoid levels for activation, these genes also require the presence of Hunchback protein to be transcribed (Simpson-Brose et al. 1994; Reinitz et al. 1995). The Bicoid and Hunchback proteins act synergistically at the enhancers of these "head genes" to promote their transcription.

**The posterior organizing center: localizing and activating nanos**

The posterior organizing center is defined by the activities of the nanos gene (Lehmann and Nüsslein-Volhard 1991; Wang and Lehmann 1991; Wharton and Struhl 1991). The nanos RNA is produced by the ovarian nurse cells and is transported into the posterior region of the egg (farthest away from the nurse cells). The nanos message is bound to the cytoskeleton in the posterior region of the egg through its 3′ UTR and its association with the products of several other genes (oskar, valois, vasa, staufen, and tudor).* If nanos or any other of these maternal effect genes are absent in the mother, no embryonic abdomen forms (Lehmann and Nüsslein-Volhard 1986; Schüpbach and Wieschaus 1986).

The nanos message is dormant in the unfertilized egg, as it is repressed by the binding of the Smaug protein to its 3′ UTR (Smibert et al. 1996). At fertilization, this repression is removed, and Nanos protein can be synthesized. The Nanos protein forms a gradient that is highest at the posterior end. Nanos functions by inactivating hunchback mRNA translation (Figure 9.16, see also Figure 9.11; Tautz 1988).

In the anterior of the cleavage-stage embryo, the hunchback message is bound in its 3′ UTR by the Pumilio protein, and the message can be translated into Hunchback protein. In the posterior of the early embryo, however, the bound Pumilio can be joined by the Nanos protein. Nanos binds to Pumilio and deadenylates the hunchback mRNA, preventing its translation (Barker et al. 1992; Wreden et al. 1997). The hunchback RNA is initially present throughout the embryo, although more can be made from zygotic nuclei if they are activated by Bicoid protein. Thus, the combination of Bicoid and Nanos proteins causes a gradient of Hunchback protein across the egg. The Bicoid protein activates hunchback gene transcription in the anterior part of the embryo, while the Nanos protein inhibits the translation of hunchback RNA in the posterior part of the embryo.
The terminal gene group

In addition to the anterior and posterior morphogens, there is third set of maternal genes whose proteins generate the extremes of the anterior-posterior axis. Mutations in these terminal genes result in the loss of the unsegmented extremities of the organism: the acron and the most anterior head segments and the telson (tail) and the most posterior abdominal segments (Degelmann et al. 1986; Klingler et al. 1988). A critical gene here appears to be torso, a gene encoding a receptor tyrosine kinase. The embryos of mothers with mutations of the torso gene have neither acron nor telson, suggesting that the two termini of the embryo are formed through the same pathway. The torso RNA is synthesized by the ovarian cells, deposited in the oocyte, and translated after fertilization. The transmembrane Torso protein is not spatially restricted to the ends of the egg, but is evenly distributed throughout the plasma membrane (Casanova and Struhl 1989). Indeed, a dominant mutation of torso, which imparts constitutive activity to the receptor, converts the entire anterior half of the embryo into an acron and the entire posterior half into a telson. Thus, Torso must normally be activated only at the ends of the egg.

Stevens and her colleagues (1990) have shown that this is the case. Torso protein is activated by the follicle cells only at the two poles of the oocyte. Two pieces of evidence suggest that the activator of the Torso protein is probably the Torso-like protein: first, loss-of-function mutations in the torso-like gene create a phenotype almost identical to that produced by torso mutants, and second, ectopic expression of Torso-like causes the activation of the Torso protein in the new location. The torso-like gene is usually expressed only in the anterior and posterior follicle cells, and the secreted Torso-like protein can cross the perivitelline space to activate the Torso protein in the egg membrane (Martin et al. 1994; Furriols et al. 1998). In this manner, the Torso-like protein activates the Torso protein in the anterior and posterior regions of the oocyte membrane. The end products of the RTK-kinase cascade activated by the Torso protein diffuse into the cytoplasm at both ends of the embryo (Figure 9.17; Gabay et al. 1997; see Chapter 6).

These kinases are thought to inactivate a transcriptional inhibitor of the tailless and huckebein gap genes (Paroush et al. 1997). These two genes then specify the termini of the embryo. The distinction between the anterior and posterior termini depends on the presence of Bicoid. If the terminal genes act alone, the terminal regions differentiate into telsons. However, if Bicoid is also present, the region forms an acron (Pignoni et al. 1992).

The anterior-posterior axis of the embryo is therefore specified by three sets of genes: those that define the anterior organizing center, those that define the posterior organizing center, and those that define the terminal boundary region. The anterior organizing center is located at the anterior end of the embryo and acts through a gradient of Bicoid protein that functions as a
transcription factor to activate anterior-specific gap genes and as a translational repressor to suppresses posterior-specific gap genes. The posterior organizing center is located at the posterior pole and acts translationally through the Nanos protein to inhibit anterior formation and transcriptionally through the Caudal protein to activate those genes that form the abdomen. The boundaries of the acron and telson are defined by the product of the torso gene, which is activated at the tips of the embryo. The activation of those genes responsible for constructing the posterior is performed by Caudal, a protein whose synthesis (as we have seen above) is inhibited in the anterior portion of the embryo. The next step in development will be to use these gradients of transcription factors to activate specific genes along the anterior-posterior axis.

The Segmentation Genes

The process of cell fate commitment in Drosophila appears to have two steps: specification and determination (Slack 1983). Early in development, the fate of a cell depends on environmental cues, such as those provided by the protein gradients mentioned above. This specification of cell fate is flexible and can still be altered in response to signals from other cells. Eventually, the cells undergo a transition from this loose type of commitment to an irreversible determination. At this point, the fate of a cell becomes cell-intrinsic. The transition from specification to determination in Drosophila is mediated by the segmentation genes. These genes divide the early embryo into a repeating series of segmental primordia along the anterior-posterior axis. Mutations in segmentation genes cause the embryo to lack certain segments or parts of segments. Often these mutations affect parasegments, regions of the embryo that are separated by mesodermal thickenings and ectodermal grooves. The segmentation genes divide the embryo into 14 parasegments (Martinez-Arias and Lawrence 1985). The parasegments of the embryo do not become the segments of the larva or adult; rather, they include the posterior part of an anterior segment and the anterior portion of the segment behind it (Figure 9.18). While the segments are the major anatomical divisions of the larval and adult body plan, they are built according to rules that use the parasegment as the basic unit of construction.

There are three classes of segmentation genes, which are expressed sequentially (see Figure 9.8). The transition from an embryo characterized by gradients of morphogens to an embryo with distinct units is accomplished by the products of the gap genes. The gap genes are activated or repressed by the maternal effect genes, and they divide the embryo into broad regions, each containing several parasegment primordia.

TheKrüppel gene, for example, is expressed primarily in parasegments 4–6, in the center of the Drosophila embryo (Figures 9.19A; 9.8B); the absence of the Krüppel protein causes the embryo to lack these regions. The protein products of the gap genes interact with neighboring gap gene proteins to activate the transcription of the pair-rule genes.
The products of these genes subdivide the broad gap gene regions into parasegments. Mutations of pair-rule genes, such as *fushi tarazu* (Figures 9.8C, 9.19B, 9.20), usually delete portions of alternate segments. Finally, the **segment polarity** genes are responsible for maintaining certain repeated structures within each segment. Mutations in these genes cause a portion of each segment to be deleted and replaced by a mirror-image structure of another portion of the segment. For instance, in *engrailed* mutants, portions of the posterior part of each segment are replaced by duplications of the anterior region of the subsequent segment (Figures 9.19C, 9.8D). Thus, the segmentation genes are transcription factors that use the gradients of the early-cleavage embryo to transform the embryo into a periodic, parasegmental structure.

After the parasegmental boundaries are set, the pair-rule and gap genes interact to regulate the homeotic selector genes, which determine the identity of each segment. By the end of the cellular blastoderm stage, each segment primordium has been given an individual identity by its unique constellation of gap, pair-rule, and homeotic gene products (*Levine and Harding 1989*).
The gap genes

The gap genes were originally discovered through a series of mutant embryos that lacked groups of consecutive segments (Figure 9.21; Nüsslein-Volhard and Wieschaus 1980). Deletions caused by mutations of the hunchback, Krüppel, and knirps genes span the entire segmented region of the Drosophila embryo. The giant gap gene overlaps with these three, and mutations of the tailless and huckebein genes delete portions of the unsegmented termini of the embryo.

The expression of the gap genes is dynamic. There is usually a low level of transcriptional activity across the entire embryo that becomes defined into discrete regions of high activity as cleavage continues (Jäckle et al. 1986). The critical element appears to be the expression of the Hunchback protein, which by the end of nuclear division cycle 12 is found at high levels across the anterior part of the embryo, and then forms a steep gradient through about 15 nuclei. The last third of the embryo has undetectable Hunchback levels. The transcription patterns of the anterior gap genes are initiated by the different concentrations of the Hunchback and Bicoid proteins.

High levels of Hunchback protein induce the expression of giant, while the Krüppel transcript appears over the region where Hunchback begins to decline. High levels of Hunchback protein also prevent the transcription of the posterior gap genes (such as knirps) in the anterior part of the embryo (Struhl et al. 1992). It is thought that a gradient of the Caudal protein, highest at the posterior pole, is responsible for activating the abdominal gap genes knirps and giant.

The giant gene has two methods for its activation, one for its anterior expression band and one for its posterior expression band (Rivera-Pomar 1995; Schulz and Tautz 1995).
After the establishment of these patterns by the maternal effect genes and Hunchback, the expression of each gap gene becomes stabilized and maintained by interactions between the different gap gene products themselves (Figure 9.21B). For instance, Krüppel gene expression is negatively regulated on its anterior boundary by the Hunchback and Giant proteins and on its posterior boundary by the Knirps and Tailless proteins (Jäckle et al. 1986; Harding and Levine 1988; Hoch et al. 1992). If Hunchback activity is lacking, the domain of Krüppel expression extends anteriorly. If Knirps activity is lacking, Krüppel gene expression extends more posteriorly. The boundaries between the regions of gap gene transcription are probably created by mutual repression. Just as the Giant and Hunchback proteins can control the anterior boundary of Krüppel transcription, so Krüppel protein can determine the posterior boundaries of giant and hunchback transcription. If an embryo lacks the Krüppel gene, hunchback transcription continues into the area usually allotted to Krüppel (Jäckle et al. 1986; Kraut and Levine 1991). These boundary-forming inhibitions are thought to be directly mediated by the gap gene products, because all four major gap genes (hunchback, giant, Krüppel, and knirps) encode DNA-binding proteins that can activate or repress the transcription of other gap genes (Knipple et al. 1985; Gaul and Jäckle 1990; Capovilla et al. 1992).

**The pair-rule genes**

The first indication of segmentation in the fly embryo comes when the pair-rule genes are expressed during the thirteenth division cycle. The transcription patterns of these genes are striking in that they divide the embryo into the areas that are the precursors of the segmental body plan. As can be seen in Figure 9.22B-E and Figure 9.8C, one vertical band of nuclei (the cells are just beginning to form) expresses a pair-rule gene, then another band of nuclei does not express it, and then another band of nuclei expresses it again. The result is a "zebra stripe" pattern along the anterior-posterior axis, dividing the embryo into 15 subunits (Hafen et al. 1984). Eight genes are currently known to be capable of dividing the early embryo in this fashion; they are listed in Table 9.2. It is important to note that not all nuclei express the same pair-rule genes. In fact, within each parasegment, each row of nuclei has its own constellation of pair-rule gene expression that distinguishes it from any other row.

How are some nuclei of the *Drosophila* embryo told to transcribe a particular gene while their neighbors are told not to transcribe it? The answer appears to come from the distribution of the protein products of the gap genes. Whereas the RNA of each of the gap genes has a very discrete distribution that defines abutting or slightly overlapping regions of expression, the protein products of these genes extend more broadly.
Table 9.2. Major genes affecting segmentation pattern in *Drosophila*

<table>
<thead>
<tr>
<th>Category</th>
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<tr>
<td>Gap genes</td>
<td>Pair-rule genes</td>
</tr>
<tr>
<td>Krüppel (Kr)</td>
<td>fushi tarazu (ftz)</td>
</tr>
<tr>
<td>knirps (kni)</td>
<td>odd-paired (opa)</td>
</tr>
<tr>
<td>hunchback (hh)</td>
<td>odd-skipped (slp)</td>
</tr>
<tr>
<td>giant (gt)</td>
<td>sloppy-paired (slp)</td>
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<tr>
<td>tailless (tll)</td>
<td>paired (prd)</td>
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<tr>
<td>huckeindein (hkb)</td>
<td>段</td>
</tr>
<tr>
<td>buttonhead (btd)</td>
<td>Segment</td>
</tr>
<tr>
<td>empty spiracles (ems)</td>
<td>polarity genes</td>
</tr>
<tr>
<td>orthodenticle (otd)</td>
<td></td>
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<tr>
<td>Pair-rule genes</td>
<td>Primary</td>
</tr>
<tr>
<td>hairy (h)</td>
<td>hairy, even-skipped, runt</td>
</tr>
<tr>
<td>even-skipped (eve)</td>
<td>even-skipped</td>
</tr>
<tr>
<td>runt (run)</td>
<td>runt</td>
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</tbody>
</table>

In fact, they overlap by at least 8–10 nuclei (which at this stage accounts for about two to three segment primordia). This was demonstrated in a striking manner by Štanojevic and co-workers (1989). They fixed cellularizing blastoderms (i.e., the stage when cells are beginning to form at the rim of the syncytial embryo), stained the Hunchback protein with an antibody carrying a red dye, and simultaneously stained the Krüppel protein with an antibody carrying a green dye. Cellularizing regions that contained both proteins bound both antibodies and were stained bright yellow (see Figure 9.8B). Krüppel protein overlaps with Knirps protein in a similar manner in the posterior region of the embryo (Pankratz et al. 1990).

Three genes are known to be the primary pair-rule genes. These genes—hairy, even-skipped, and runt—are essential for the formation of the periodic pattern, and they are directly controlled by the gap gene proteins. The enhancers of the primary pair-rule genes are recognized by gap gene proteins, and it is thought that the different concentrations of gap gene proteins determine whether a pair-rule gene is transcribed or not. The enhancers of the primary pair-rule genes are often modular: the control over each stripe is located in a discrete region of the DNA. One of the best-studied enhancers is that for the even-skipped gene.

The structure of this enhancer is shown in Figure 9.22A. It is composed of modular units arranged in such a way that each unit regulates a separate stripe. For instance, the second even-skipped stripe is repressed by both Giant and Krüppel proteins and is activated by Hunchback protein and low concentrations of Bicoid (Figure 9.23; Small et al. 1991, 1992; Štanojevic et al. 1991).
DNase I footprinting (see Chapter 5) showed that the enhancer region for this stripe contains six binding sites for Krüppel protein, three for Hunchback protein, three for Giant protein, and five for Bicoid protein. Similarly, even-skipped stripe 5 is regulated negatively by Krüppel protein (on its anterior border) and by Giant protein (on its posterior border) (Small et al. 1996; Fujioka 1999).

The importance of these enhancers can be shown by both genetic and biochemical means. First, a mutation in a particular enhancer can delete its particular stripe and no other. Second, if a reporter gene such as lacZ (encoding β-galactosidase) is fused to one of these enhancer elements, the lacZ gene is expressed only in that particular stripe (see Figure 9.22; Fujioka et al. 1999). Third, the placement of the stripes can be altered by deleting the gap genes that regulate them. Thus, the placement of the stripes of pair-rule gene expression is a result of (1) the modular cis-regulatory enhancer elements of the pair-rule genes and (2) the trans-regulatory gap gene proteins that bind to these enhancer sites.

Once initiated by the gap gene proteins, the transcription pattern of the primary pair-rule genes becomes stabilized by their interactions among themselves (Levine and Harding 1989). The primary pair-rule genes also form the context that allows or inhibits the expression of the later-acting secondary pair-rule genes. One such secondary pair-rule gene is fushi tarazu (ftz; Japanese, "too few segments;" Figures 9.8, 9.19, 9.20). Early in cycle 14, ftz RNA and protein are seen throughout the segmented portion of the embryo. However, as the proteins from the primary pair-rule genes begin to interact with the ftz enhancer, the ftz gene is repressed in certain bands of nuclei to create interstripe regions. Meanwhile, the Ftz protein interacts with its own promoter to stimulate more transcription of the ftz gene (Figure 9.24; Edgar et al. 1986b; Karr and Kornberg 1989; Schier and Gehring 1992).

The expression of the each pair-rule gene in seven stripes divides the embryo into fourteen parasegments, with each pair-rule gene being expressed in alternate parasegments. Moreover, each row of nuclei within each parasegment expresses a particular and unique combination of pair-rule products. These products will activate the next level of segmentation genes, the segment polarity genes.

The segment polarity genes

So far, our discussion has described interactions between molecules within the syncytial embryo. But once cells form, interactions take place between the cells. These intercellular interactions are mediated by the segment polarity genes, and they accomplish two important tasks. First, they reinforce the parasegmental periodicity established by the earlier transcription factors. Second, through this cell-to-cell signaling, cell fates are established within each parasegment.

The segment polarity genes encode proteins that are constituents of the Wingless and Hedgehog signal transduction pathways (see Chapter 6). Mutations in these genes lead to defects in segmentation and in gene expression pattern across each parasegment.
The development of the normal pattern relies on the fact only one row of cells in each parasegment is permitted to express the Hedgehog protein, and only one row of cells in each parasegment is permitted to express the Wingless protein.

The key to this pattern is the activation of the engrailed gene in those cells that are going to express the Hedgehog protein. The engrailed gene is activated when cells have high levels of the Even-skipped, Fushi tarazu, or Paired transcription factors. Moreover, it is repressed in those cells that receive high levels of Odd-skipped, Runt, or Sloppy-paired proteins.

As a result, Engrailed is expressed in fourteen stripes across the anterior-posterior axis of the embryo (see Figure 9.8D). (Indeed, in mutations that cause the embryo to be deficient in Fushi tarazu, only seven bands of Engrailed are expressed.)

These stripes of engrailed transcription mark the anterior boundary of each parasegment (and the posterior border of each segment). The wingless gene is activated in those bands of cells that receive little or no Even-skipped or Fushi tarazu proteins, but which do contain the Sloppy-paired protein. This causes wingless to be transcribed solely in the row of cells directly anterior to the cells where engrailed is transcribed (Figure 9.25).

Once wingless and engrailed expression is established in adjacent cells, this pattern must be maintained to retain the parasegmental periodicity of the body plan established by the pair-rule genes. It should be remembered that the mRNAs and proteins involved in initiating these patterns are short-lived, and that the patterns must be maintained after their initiators are no longer being
synthesized. The maintenance of these patterns is regulated by interactions between cells expressing wingless and those expressing engrailed. The Wingless protein, secreted from the wingless-expressing cells, diffuses to adjacent cells. The cells expressing engrailed can bind this protein because they contain the Drosophila membrane receptor protein for Wingless, D-Frizzled-2 (see Figure 6.23; Bhanot et al. 1996). This receptor activates the Wnt signal transduction pathway, resulting in the continued expression of engrailed (Siegfried et al. 1994).

Moreover, this activation starts another portion of this reciprocal pathway. The Engrailed protein activates the transcription of the hedgehog gene in the engrailed-expressing cells. The Hedgehog protein can bind to the Hedgehog receptor (the Patched protein) on neighboring cells. When it binds to the adjacent posterior cells, it stimulates the expression of the wingless gene. The result is a reciprocal loop wherein the Engrailed-synthesizing cells secrete the Hedgehog protein, which maintains the expression of the wingless gene in the neighboring cells, while the Wingless-secreting cells maintain the expression of the engrailed and hedgehog genes in their neighbors in turn (Heemskerk et al. 1991; Ingham et al. 1991; Mohler and Vani 1992). In this way, the transcription pattern of these two types of cells is stabilized. This interaction creates a stable boundary, as well as a signaling center from which Hedgehog and Wingless proteins diffuse across the parasegment.

The diffusion of these proteins is thought to provide the gradients by which the cells of the parasegment acquire their identities. This process can be seen in the dorsal epidermis, where the rows of larval cells produce different cuticular structures depending on their position within the segment. The 1° row consists of large, pigmented spikes called denticles. Posterior to these cells, the 2° row produces a smooth epidermal cuticle. The next two cell rows have a 3° fate, making small, thick hairs, and these are followed by several rows of cells that adopt the 4° fate, producing fine hairs (Figure 9.26).

The wingless-expressing cells lie within the region producing the fine hairs, while the hedgehog-expressing cells are near the 1° row of cells. The fates of the cells can be altered by experimentally increasing or decreasing the levels of Hedgehog or Wingless protein.

For example, if the hedgehog gene is fused to a heat shock promoter and the embryos are grown at a temperature that activates the gene, more Hedgehog protein is made, and the cells normally showing 3° fates will become 2° cells. The rows of 4° cells farthest from the Wingless-secreting cells may also become 3° or 2° cells. It seems that the cells closest to the Wingless secreters cannot respond to Hedgehog, and Hedgehog cannot, by itself, specify the 1° fate (which may require the expression of certain pair-rule gene products). Thus, Hedgehog and Wingless appear necessary for elaborating the entire pattern of cell types across the parasegment.
However, the mechanism by which they accomplish this specification is not clear. Either these signals act in a graded fashion, as morphogens, or they act locally to initiate a cascade of local signaling events, in which each interaction uses a different ligand and receptor (Figure 9.26). The resulting pattern of cell fates also changes the focus of patterning from parasegment to segment. There are now external markers, as the engrailed-expressing cells become the most posterior cells of each segment.

The Homeotic Selector Genes

Patterns of homeotic gene expression

After the segmental boundaries have been established, the characteristic structures of each segment are specified. This specification is accomplished by the homeotic selector genes (Lewis 1978).

There are two regions of Drosophila chromosome 3 that contain most of these homeotic genes (Figure 9.27). One region, the Antennapedia complex, contains the homeotic genes labial (lab), Antennapedia (Antp), sex combs reduced (scr), deformed (dfd), and proboscipedia (pb). The labial and deformed genes specify the head segments, while sex combs reduced and Antennapedia contribute to giving the thoracic segments their identities. The proboscipedia gene appears to act only in adults, but in its absence, the labial palps of the mouth are transformed into legs (Wakimoto et al. 1984; Kaufman et al. 1990). The second region of homeotic genes is the bithorax complex (Lewis 1978). There are three protein-coding genes found in this complex: ultrabithorax (ubx), which is required for the identity of the third thoracic segment; and the abdominal A (abdA) and Abdominal B (AbdB) genes, which are responsible for the segmental identities of the abdominal segments (Sánchez-Herrero et al. 1985). The lethal phenotype of the triple-point mutant Ubx-, abdA-, AbdB- is identical to that resulting from a deletion of the entire bithorax complex (Casanova et al. 1987). The chromosome region containing both the Antennapedia complex and the bithorax complex is often referred to as the homeotic complex (Hom-C).

Because these genes are responsible for the specification of fly body parts, mutations in them lead to bizarre phenotypes. In 1894, William Bateson called these organisms "homeotic mutants," and they have fascinated developmental biologists for decades. For example, the body of the normal adult fly contains three thoracic segments, all of which produce a pair of legs. The first thoracic segment does not produce any further appendages, but the second thoracic segment produces both a set of legs and a set of wings. The third thoracic segment produces a set of wings and a set of balancers known as halteres. In homeotic mutants, these specific segmental identities
can be changed. When the ultrabithorax gene is deleted, the third thoracic segment (which is characterized by halteres) becomes transformed into another second thoracic segment. The result (Figure 9.28) is a fly with four wings—an embarrassing situation for a classic dipteran. Similarly, the Antennapedia protein is usually used to specify the second thoracic segment of the fly. But when flies have a mutation wherein the Antennapedia gene is expressed in the head (as well as in the thorax), legs rather than antennae grow out of the head sockets (Figure 9.29). In the recessive mutant of Antennapedia, the gene fails to be expressed in the second thoracic segment, and antennae sprout out of the leg positions (Struhl 1981; Frischer et al. 1986; Schneuwly et al. 1987).

These major homeotic selector genes have been cloned and their expression analyzed by in situ hybridization (Harding et al. 1985; Akam 1987). Transcripts from each gene can be detected in specific regions of the embryo and are especially prominent in the central nervous system (see Figure 9.27).

**Initiating the patterns of homeotic gene expression**

The initial domains of homeotic gene expression are influenced by the gap genes and pair-rule genes. For instance, the expression of the abdA and AbdB genes is repressed by the gap gene proteins Hunchback and Krüppel. This inhibition prevents these abdomen-specifying genes from being expressed in the head and thorax (Casares and Sánchez-Herrero 1995). Conversely, the Ultrabithorax gene is activated by certain levels of the Hunchback protein, so that it is originally transcribed in a broad band in the middle of the embryo, and the transcription of Antennapedia is activated by Krüppel (Harding and Levine 1988; Struhl et al. 1992). The boundaries of homeotic gene expression are soon confined to the parasegments defined by the Fushi tarazu and Even-skipped proteins (Ingham and Martinez-Arias 1986; Müller and Bienz 1992).

**Maintaining the patterns of homeotic gene expression**

The expression of homeotic genes is a dynamic process. The Antennapedia gene (Antp), for instance, although initially expressed in presumptive parasegment 4, soon appears in parasegment 5. As the germ band expands, Antp expression is seen in the presumptive neural tube as far posterior as parasegment 12. During further development, the domain of Antp expression contracts again, and Antp transcripts are localized strongly to parasegments 4 and 5. Like that of other homeotic genes, Antp expression is negatively regulated by all the homeotic gene products expressed posterior to it (Harding and Levine 1989; González-Reyes and Morata 1990). In other words, each of the bithorax complex genes represses the expression of Antennapedia. If the Ultrabithorax gene is deleted, Antp activity extends through the region that would normally have expressed Ubx and stops where the Abd region begins. (This allows the third thoracic segment to form wings like the second thoracic segment, as in Figure 9.29.) If the entire bithorax complex is deleted, Antp expression extends throughout the abdomen. (Such a larva does not survive, but the cuticle pattern throughout the abdomen is that of the second thoracic segment.)
As we saw above, the gap gene and pair-rule gene proteins are transient, but the identities of the segments must be stabilized so that differentiation can occur. Thus, once the transcription patterns of the homeotic genes have become stabilized, they are "locked" into place by alteration of the chromatin conformation in these genes. The repression of homeotic genes appears to be maintained by the Polycomb family of proteins, while the active chromatin conformation appears to be maintained by the Trithorax proteins (Ingham and Whittle 1980; McKeon and Brock 1991; Simon et al. 1992).

Realisator genes

The search is now on for "realisator genes," those genes that are the targets of the homeotic gene proteins and which function to form the specified tissue or organ primordia. In the formation of the second thoracic segment, for example, Antennapedia is expressed. Casares and Mann (1998) have shown that Antennapedia protein binds to the enhancer of the homothorax gene and prevent its expression. Homothorax is necessary for producing a transcription factor critical for antenna formation. Therefore, one of Antennapedia's functions is to suppress those genes necessary for antenna development.

The Ultrabithorax protein is able to repress the expression of the Wingless gene in those cells that will become the halteres of the fly. One of the major differences between the appendage-forming cells of the second and the third thoracic segments is that Wingless expression occurs in the appendage-forming cells of the second thoracic segment, but not in those of the third thoracic segment. Wingless acts as a growth promoter and morphogen in these tissues. In the third thoracic segment, Ubx protein is found in these cells, and it prevents the expression of the Wingless gene (Figure 9.30; Weatherbee et al. 1998). Thus, one of the ways in which Ubx protein specifies the third thoracic segment is by preventing the expression of those genes that would generate the wing tissue.

Another target of the homeotic proteins, the distal-less gene (itself a homeobox-containing gene: see Sidelights and Speculations), is necessary for limb development and is active solely in the thorax. Distal-less expression is repressed in the abdomen, probably by a combination of Ubx and AbdA proteins that can bind to its enhancer and block transcription (Vachon et al. 1992; Castelli-Gair and Akam 1995). This presents a paradox, since parasegment 5 (entirely thoracic and leg-producing) and parasegment 6 (which includes most of the legless first abdominal segment) both expressUbx. How can these two very different segments be specified by the same gene? Castelli-Gair and Akam (1995) have shown that the mere presence of Ubx protein in a group of cells is not sufficient for specification. Rather, the time and place of its expression within the parasegment can be critical. Before Ubx expression, parasegments 4–6 have similar potentials. At division cycle 10, Ubx expression in the anterior parts of parasegments 5 and 6 prevents those parasegments from forming structures (such as the anterior spiracle) characteristic of parasegment 4. Moreover, in the posterior compartment of parasegment 6 (but not parasegment 5), Ubx protein blocks the formation of the limb primordium by repressing the distal-less gene. At division cycle 11, by which time Ubx has pervaded all of parasegment 6, the distal-less gene has become self-regulatory and cannot be repressed by Ubx (Figure 9.31).
*Like the placement of the bicoid message, the location of the nanos message is determined by its 3′ untranslated region. If the bicoid 3′ UTR is experimentally placed on the protein-encoding region of nanos RNA, the nanos message gets placed in the anterior of the egg. When the RNA is translated, the Nanos protein inhibits the translation of hunchback and bicoid mRNAs, and the embryo forms two abdomens— one in the anterior of the embryo and one in the posterior (Gavis and Lehmann 1992). The localization of nanos RNA is ultimately dependent on interactions between the oocyte and the neighboring follicle cells that localize the oskar message to the posterior pole.

Aficionados of information theory will recognize that the process by which the anterior-posterior information in morphogenetic gradients is transferred to discrete and different parasegments represents a transition from analog to digital specification. Specification is analog, determination digital. This process enables the transient information of the gradients in the syncytial blastoderm to be stabilized so that it can be utilized much later in development (Baumgartner and Noll 1990).

† The interactions between genes and gene products are facilitated by the fact that these reactions occur within a syncytium, in which the cell membranes have not yet formed.

§ Homeo means "similar." Homeotic mutants are mutants in which one structure is replaced by another (as where an antenna is replaced by a leg). Homeotic genes are genes whose mutation can cause such transformations; thus, they are genes that specify the identity of a particular body segment. The homeobox is a conserved DNA sequence of about 180 base pairs that is shared by many homeotic genes. This sequence encodes the 60-amino acid homeodomain, which recognizes specific DNA sequences. The homeodomain is an important region of the transcription factors encoded by homeotic genes (see Sidelights & Speculations). Not all genes with homeoboxes are homeotic genes, however.

¶ Dipterans (two-winged insects such as flies) are thought to have evolved from four-winged insects; it is possible that this change arose via alterations in the bithorax complex. Chapter 22 includes more speculation on the relationship between the homeotic complex and evolution.

The Homeodomain Proteins

Homeodomain proteins are a family of transcription factors characterized by a 60-amino acid domain (the homeodomain) that binds to certain regions of DNA. The homeodomain was first discovered in those proteins whose absence or misregulation caused homeotic transformations of Drosophila segments. It is thought that homeodomain proteins activate batteries of genes that specify the particular properties of each segment. The homeodomain proteins include the products of the eight genes of the homeotic complex, as well as other proteins such as Fushi tarazu, Caudal, Distal-less and Bicoid. Homeodomain proteins are important in determining the anterior-posterior axes of both invertebrates and vertebrates.

In Drosophila, the presence of certain homeodomain proteins is also necessary for the determination of specific neurons. Without these transcription factors, the fates of these neuronal cells are altered (Doe et al. 1988).

The homeodomain is encoded by a 180-base-pair DNA sequence known as the homeobox. The homeodomains appear to be the sites of these proteins that bind DNA, and they are critical in specifying cell fates. For instance, if a chimeric protein is constructed mostly of Antennapedia but with the carboxyl terminus (including the homeodomain) of Ultrabithorax, it
can substitute for Ultrabithorax and specify the appropriate cells as parasegment 6 (Mann and Hogness 1990). The isolated homeodomain of Antennapedia will bind to the same promoters as the entire Antennapedia protein, indicating that the binding of this protein is dependent on its homeodomain (Müller et al. 1988).

The homeodomain folds into three helices, the latter two folding into a helix-turn-helix conformation that is characteristic of transcription factors that bind DNA in the major groove of the double helix (Otting et al. 1990; Percival-Smith et al. 1990). The third helix is the recognition helix, and it is here that the amino acids make contact with the bases of the DNA. A four-base motif, TAAT, is conserved in nearly all sites recognized by homeodomains; it probably distinguishes those sites to which homeodomain proteins can bind. The 5′ terminal T appears to be critical in this recognition, as mutating it destroys all homeodomain binding. The base pairs following the TAAT motif are important in distinguishing between similar recognition sites. For instance, the next base pair is recognized by amino acid 9 of the recognition helix. Mutation studies have shown that the Bicoid and Antennapedia homeodomain proteins use lysine and glutamine, respectively, at position 9 to distinguish related recognition sites. The lysine of the Bicoid homeodomain recognizes the G of CG pairs, while the glutamine of the Antennapedia homeodomain recognizes the A of AT pairs (Figure 9.32; Hanes and Brent 1991). If the lysine in Bicoid is replaced by glutamine, the resulting protein will recognize Antennapedia-binding sites (Hanes and Brent 1989, 1991). Other homeodomain proteins show a similar pattern, in which one portion of the homeodomain recognizes the common TAAT sequence, while another portion recognizes a specific structure adjacent to it.

Cofactors for the Hom-C Genes

The genes of the Drosophila homeotic complex specify segmental fates, but they may need some help in doing it. The DNA-binding sites recognized by the homeodomains of the Hom-C proteins are very similar, and there is some overlap in their binding specificity. In 1990, Peifer and Wieschaus discovered that the product of the Extradenticle (Exd) gene interacts with several Hom-C proteins and may help specify the segmental identities. For instance, the Ubx protein is responsible for specifying the identity of the first abdominal segment (A1). Without Extradenticle protein, it will transform this segment into A3. Moreover, the Exd and Ubx proteins are both needed for the regulation of the decapentaplegic gene, and the structure of the...
decapentaplegic promoter suggests that the Extradenticle protein may dimerize with the Ubx protein on the enhancer of this target gene (Raskolb and Wieschaus 1994; van Dyke and Murre 1994). The Extradenticle protein includes a homeodomain, and the human protein PBX1, which resembles the Extradenticle protein, may play a similar role as a cofactor for human homeotic genes.

The product of the teashirt gene may also be an important cofactor. This zinc finger transcription factor is necessary for the functioning of the Sex combs reduced protein, which distinguishes between the labial and first thoracic segments. It is critical for the specification of the anterior prothoracic (parasegment 3) identity, and it may be the gene that specifies the "groundstate condition" of the homeotic complex. If the bithorax complex and the Antennapedia gene are removed, all the segments become anterior prothorax. The product of the teashirt gene appears to work with the Ser protein to distinguish thorax from head and to work throughout the trunk to prevent head structures from forming (Roder et al. 1992).

**The Generation of Dorsal-Ventral Polarity**

In 1936, embryologist E. E. Just criticized those geneticists who sought to explain Drosophila development by looking at specific mutations affecting eye color, bristle number, and wing shape. He said that he wasn't interested in the development of the bristles of a fly's back; rather, he wanted to know how the fly embryo makes the back itself. Fifty years later, embryologists and geneticists are finally answering that question.*

**The Morphogenetic Agent for Dorsal-Ventral Polarity**

Dorsal-ventral polarity is established by the gradient of a transcription factor called Dorsal. Unlike Bicoid, whose gradient is established within a syncytium, Dorsal forms a gradient over a field of cells that is established as a consequence of cell-to-cell signaling events.

The specification of the dorsal-ventral axis takes place in several steps. The critical step is the translocation of the Dorsal protein from the cytoplasm into the nuclei of the ventral cells during the fourteenth division cycle. Anderson and Nüsslein-Volhard 1984 isolated 11 maternal effect genes, each of whose absence is associated with a lack of ventral structures (Figure 9.33). The absence of another maternal effect gene, cactus, causes the ventralization of all cells. The proteins encoded by these maternal genes are critical for making certain that the Dorsal protein gets into only those nuclei on the ventral surface of the embryo. After its translocation, the Dorsal protein acts on cell nuclei to specify the different regions of the embryo. Different concentrations of Dorsal protein in the nuclei appear to specify different fates in those cells.
The Translocation of Dorsal Protein

The protein that actually distinguishes dorsum (back) from ventrum (belly) is the product of the *dorsal* gene. The RNA transcript of the mother's *dorsal* gene is placed in the oocyte by her ovarian cells. However, Dorsal protein is not synthesized from this maternal message until about 90 minutes after fertilization. When this protein is translated, it is found throughout the embryo, not just on the ventral or dorsal side. How, then, can this protein act as a morphogen if it is located everywhere in the embryo?

In 1989, the surprising answer was found (Roth et al. 1989; Rushlow et al. 1989; Steward 1989). While Dorsal protein can be found throughout the syncytial blastoderm of the early *Drosophila* embryo, it is translocated into nuclei only in the ventral part of the embryo (Figure 9.34A, B). In the nucleus, Dorsal binds to certain genes to activate or suppress their transcription. If Dorsal does not enter the nucleus, the genes responsible for specifying ventral cell types (*snail* and *twist*) are not transcribed, the genes responsible for specifying dorsal cell types (*decapentaplegic* and *zerknüllt*) are not repressed, and all the cells of the embryo become specified as dorsal cells.

This model of dorsal-ventral axis formation in *Drosophila* is supported by analyses of mutations that give rise to an entirely dorsalized or an entirely ventralized phenotype (see Figures 9.33A and 9.34). In those mutants in which all the cells are dorsalized (as is evident by their dorsal cuticle), Dorsal protein does not enter the nucleus in any cell. Conversely, in those mutants in which all cells have a ventral phenotype, Dorsal protein is found in every cell nucleus.

*The signal cascade*

Signal from the oocyte nucleus to the follicle cells.

If Dorsal protein is found throughout the embryo, but gets translocated into the nuclei of only ventral cells, then something else must be providing asymmetrical cues (Figure 9.35). It appears that this signal is mediated through a complex interaction between the oocyte and its surrounding follicle cells.

The follicular epithelium surrounding the developing oocyte is initially symmetrical, but this symmetry is broken by a signal from the oocyte nucleus. The oocyte nucleus is originally located at the posterior end of the oocyte, away from the nurse cells. It then moves to an anterior dorsal position and signals the overlying follicle cells to become the more columnar dorsal follicle cells (Montell et al. 1991; Schüpbach et al. 1991). The dorsalizing signal from the oocyte nucleus appears to be the product of the *gurken* gene (Schüpbach 1987; Forlani et al. 1993). The *gurken* message becomes localized in a crescent between the oocyte nucleus and the oocyte plasma membrane, and its protein product forms an anterior-posterior gradient along the dorsal surface of the oocyte (Neuman-Silberberg and Schüpbach 1993; Figure 9.36).