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## Coordinating Growth and Review Maturation — Insights from *Drosophila*

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### Abstract

Adult body size in higher animals is dependent on the amount of growth that occurs during the juvenile stage. The duration of juvenile development, therefore, must be flexible and responsive to environmental conditions. When immature animals experience environmental stresses such as malnutrition or disease, maturation can be delayed until conditions improve and normal growth can resume. In contrast, when animals are raised under ideal conditions that promote rapid growth, internal checkpoints ensure that maturation does not occur until juvenile development is complete. Although the mechanisms that regulate growth and gate the onset of maturation have been investigated for decades, the emerging links between childhood obesity, early onset puberty, and adult metabolic disease have placed a new emphasis on this field. Remarkably, genetic studies in the fruit fly *Drosophila melanogaster* have shown that the central regulatory pathways that control growth and the timing of sexual maturation are conserved through evolution, and suggest that this aspect of animal life history is regulated by a common genetic architecture. This review focuses on these conserved mechanisms and highlights recent studies that explore how *Drosophila* coordinates developmental growth with environmental conditions.

### Introduction

The life history of insects is similar to that of other animals, with discrete stages representing embryonic development, a juvenile growth phase, sexual maturation, and reproductive adulthood. In *Drosophila*, these stages correspond to four morphologically distinct developmental states: embryo, larva (three instar stages), pupa, and adult. Embryogenesis, along with the first and second larval instars (L1 and L2), each last one day, followed by two days of third instar larval development (L3). The larval growth stage is terminated by puparium formation and four days of metamorphosis, during which the sexually active adult fly is formed [1]. Progression through all of these stages is dictated by pulses of the steroid hormone 20-hydroxyecdysone (20E) [2,3]. A series of enzymatic steps within the endocrine organ of the insect, the prothoracic gland (PG), converts cholesterol into ecdysone, which is released into the circulatory system and modified by peripheral tissues into the active form of the hormone, 20E [4,5]. This steroid acts through a

heterodimer of the ecdysone receptor (EcR) and Ultraspiracle (USP) nuclear receptors to trigger stage-specific transcriptional cascades that direct progression through each stage in the fly life cycle, determining the timing of developmental progression [2,3].

All growth in *Drosophila* normally occurs during the juvenile larval stages, resulting in a remarkable ~200-fold increase in body mass [6]. Thus, the 20E signaling events that determine the duration of larval development are critical for dictating final body size. The pulses of 20E during L1 and L2 trigger molting of the larval cuticle, accommodating the increase in animal size [2,3]. Additional low-titer hormone pulses during the L3 stage prepare the animal for metamorphosis [7], while a high-titer 20E pulse at the end of L3 terminates larval development, arrests growth, and signals the onset of adult maturation [2,3]. This critical role for 20E in determining the duration of larval development implies that key growth regulators must feed into the timing of these events. Recent genetic studies in *Drosophila* have identified these pathways, laying the groundwork for understanding how environmental factors can regulate growth and determine the timing of maturation.

### Linking Body Size to the Timing of Maturation

Both developmental and nutritional signals feed into the timing of the 20E pulses that dictate the duration of larval growth. The manner in which animals respond to these signals, however, changes during larval development, and is centered on an important but poorly understood transition that occurs near the L2-to-L3 molt in laboratory strains of *Drosophila* [8,9]. At this time, larvae first surpass ‘minimal viable weight’, where they achieve sufficient body mass to successfully complete larval and pupal development in the absence of nutrients [10,11]. This event is followed by a key life history event that commits larvae to enter metamorphosis within a definite period of time — the attainment of ‘critical weight’ [9–14]. If animals encounter poor nutrient conditions prior to the onset of critical weight, larval development will stall and subsequent 20E pulses will be delayed until growth conditions improve (Figure 1). This link between nutrition and maturation makes sense, insofar as it provides a means of storing sufficient nutrients for survival during the non-feeding pupal stage of development as well as an opportunity for the animal to achieve an appropriate size for adult reproductive fitness. The time an animal spends developing as a pre-critical weight larva can vary greatly, as progression past the critical weight checkpoint is determined by body size and not time. Although physiological studies in other insects, such as the tobacco hookworm *Manduca sexta*, have demonstrated an important role for juvenile hormone in determining the attainment of critical weight, no studies have shown that this function is conserved in *Drosophila* [12,15].

Larvae that have achieved critical weight have sufficient stored energy to successfully complete metamorphosis, and the high-titer, late-L3 pulse of 20E will occur after a definite period of time, without regard to nutrient availability (Figure 1). With this strategy in effect, environmental conditions will dictate final body size. Larvae that develop in a favorable environment will continue to grow and can significantly increase their body size before entering metamorphosis [9,14,16]. In contrast, post-critical weight larvae that experience starvation will stop growing in size but will continue to mature into smaller fertile adults [9,14,16] (Figure 1). Ultimately, both fed and starved post-critical weight animals enter

metamorphosis within a similar time frame, but animals that continue to feed are larger than animals that experience starvation. Larval development, therefore, is regulated by genetic mechanisms that coordinate developmental progression and growth with nutrient availability, uptake, and utilization. While many of these pathways control cell-intrinsic processes, successful development requires that growth is coordinated among all tissues within the larva. This level of systemic control is achieved by secreted factors that regulate cellular physiology. Intriguingly, these factors not only arise from the brain and endocrine organs, but also are produced and secreted by the main source of stored energy within the animal: the fat body.

## Nutrient Signaling and the Fat Body

The larval fat body, which functions as a hybrid of the mammalian liver and white adipose tissue, plays a central role in sensing nutritional signals and allowing diverse tissues to coordinately respond to changes in metabolic status. The central role of the fat body in regulating organismal growth was first described nearly 35 years ago, when it was found to promote nonautonomous growth in cultured larval imaginal discs [17]. Similarly, quiescent neuroblasts will reenter the cell cycle *in vitro* when co-cultured with fat body tissue from fed larvae [18]. A key to understanding how the fat body regulates peripheral tissue growth was identified in genetic screens for growth modifiers. Mutations in the putative amino-acid transporter *minidiscs* result in developmental arrest and imaginal disc growth defects even though this gene is expressed primarily within the fat body [19]. Similarly, decreased expression of the gene *slimfast* (*slif*), which encodes a cationic amino-acid transporter and is highly expressed in the fat body, delays growth and produces abnormally small animals [20]. Furthermore, tissue-specific depletion of *slif* in the fat body elicits a whole-body growth defect, demonstrating that the fat body can retard organismal growth in response to decreased amino-acid availability [20] (Figure 2). Further studies revealed that this effect is mediated by the TOR signaling pathway, which is a critical regulator of nutrient signaling in *Drosophila* [21,22]. Mutations that disrupt the TOR pathway phenocopy the loss of *slif* function. These phenotypes can also be seen with fat body specific TOR inactivation, while overexpression of the TOR downstream target S6 kinase can partially rescue the growth defects caused by *slif* depletion [20]. The fat body, therefore, can monitor amino-acid levels via the TOR signaling pathway and can remotely coordinate growth and developmental progression (Figure 2).

A primary focus of fat body regulated growth is the insulin signaling pathway [16,23] (Figure 2). The *Drosophila* genome encodes seven insulin-like peptides (DILP1–7) [24,25], three of which, DILP2, 3, and 5, are expressed in two clusters of neurosecretory cells within the larval brain. These insulin-producing cells (IPCs) are functionally similar to pancreatic  $\beta$ -cells, and can secrete DILPs into the hemolymph [26]. Circulating DILPs bind to the *Drosophila* insulin receptor (dInR) on target cells and activate a highly conserved phosphoinositide 3 kinase (PI3K) signaling cascade that inhibits the dFOXO transcription factor, thereby promoting cell-autonomous growth [27]. When a larva experiences nutrient deprivation there is a decrease in dInR-dependent PI3K activity [28]. As a result, dFOXO translocates from the cytoplasm into the nucleus and inhibits cell growth [29–31].

Since larvae possess an open circulatory system, nutrient-deprived animals can rapidly slow development by regulating DILP activity. The expression of both *dilp3* and *dilp5* are transcriptionally downregulated under low nutrient conditions [24]. In contrast, *dilp2* transcript levels are insensitive to nutrient deprivation, but DILP2 protein secretion and signaling activity are heavily influenced by metabolic status. This regulation can be readily visualized in the IPCs, where DILP2 protein is present at a basal level in well-fed animals but accumulates to relatively high concentrations upon starvation or amino-acid deprivation [32].

DILP2 secretion is a primary target of nutrient-dependent fat body signaling (Figure 2). When *slif* expression or TOR activity is specifically disrupted in the fat body, DILP2 is not secreted and accumulates in the IPCs [32]. Furthermore, in a series of elegant experiments, co-culturing brains isolated from starved L3 larvae with fat body tissue or hemolymph from fed animals was shown to promote DILP2 secretion from the starved animals' IPCs [32]. This response, however, did not occur when the brain co-culture was conducted using tissue from starved larvae. Similarly, quiescent larval neuroblasts re-enter the cell cycle in response to TOR activation within the fat body, promoting brain growth during larval stages [18,33,34]. The fat body of fed animals, therefore, must secrete an as yet unidentified factor that promotes DILP secretion. This approach paves the way for a clearer understanding of the mechanisms by which the fat body can sense nutritional status and relay that information to control developmental growth.

The fat body not only controls DILP secretion, but also releases two proteins, Imp-L2 and dALS (acid labile subunit), that interact with circulating DILP2 [35,36] (Figure 2). These two proteins, however, do not promote insulin-signaling, but rather appear to sequester and inactivate DILP2 in a stable trimeric complex, as fat-body-specific depletion of either *Imp-L2* or *dALS* results in an overgrowth phenotype [20,35,36]. These studies provide a new context for understanding the mechanisms by which ALS contributes to mammalian insulin signaling as well as insights into how the larval fat body can control systemic insulin signaling and coordinate organismal growth.

The fat body also releases stored nutrients to ensure the survival of peripheral tissues during periods of starvation. Under normal growth conditions, nutrient-dependent TOR signaling functions cell autonomously to suppress autophagy — a process by which cells can non-specifically degrade bulk cytoplasm for energy production (Figure 2). When animals become nutrient-deprived, decreased TOR signaling results in autophagic degradation of the fat body, thereby releasing nutrients that help sustain the starving animal [37,38]. Intriguingly, decreased insulin signaling within fat body cells can also promote autophagy [37,38]. The fat body, therefore, both coordinates systemic growth and provides an essential source of energy in response to unfavorable environmental conditions.

## An Evolutionarily Conserved Program to Coordinate Growth and Maturation

Once the larva achieves critical weight, the PG begins to release low-titer pulses of 20E, preparing the animal for the cessation of larval development [7] (Figure 1). This 'mid-third

instar transition' consists of key behavioral and developmental changes, including a cessation of feeding, the onset of wandering behavior, glue protein synthesis in the salivary glands, and the initiation of fat body autophagy [37–39]. The regulation of steroid hormone activity to promote maturation appears to define an ancient regulatory pathway by which many animals control this key life history event. This common genetic architecture was first described in the context of the *Caenorhabditis elegans* life cycle, when the animal makes a decision, based on environmental factors, to either continue development to form a reproductive adult or enter a larval diapause state [40,41]. When juvenile worms encounter poor growth conditions during the first larval stage, they indefinitely arrest development by forming a dauer larva, an alternative to the L3 stage that is ideally suited for survival [42].

The genetic mechanisms that regulate developmental growth were discovered through an elegant and unbiased genetic screen for mutations that affect dauer formation [43,44]. The identification and subsequent characterization of these abnormal dauer formation (*daf*) mutants defined three key signaling pathways: insulin, TGF $\beta$ , and the steroid hormone dafachronic acid (DA) (Figure 3). Mutations that eliminate signaling through either the insulin or TGF $\beta$  pathways cause animals to become dauers independent of culture conditions [43,45,46]. In contrast, *daf* mutations that disrupt negative regulators of either pathway, such as the FOXO homolog [47,48], render animals incapable of forming dauers [43]. These genetic studies demonstrate that both pathways converge on the regulation of DA signaling. DA is a steroid hormone that binds to and modulates the activity of the nuclear receptor DAF-12 [49]. When DA is present, DAF-12 promotes continuous development, while in the absence of DA production DAF-12 induces dauer formation. Similarly, loss-of-function mutations that reduce DA synthesis or prevent DA from binding to the DAF-12 ligand-binding domain lead to dauer formation [50–52]. A commitment to adult maturation, therefore, is dependent on the coordinate activity of these three signaling pathways. Favorable growth conditions stimulate insulin and TGF $\beta$  signaling, which, in turn, promotes DA production, DAF-12 activation, and continued progression through larval development [53,54]. Conversely, poor growth conditions reduce insulin and TGF $\beta$  signaling, resulting in decreased DAF-12 signaling and dauer formation [53,54] (Figure 3).

Remarkably, recent studies suggest that a similar genetic framework controls the assessment of critical weight and maturation in *Drosophila* (Figure 3). These pathways exert their effect, in part, by sensitizing the PG to the activity of prothoracicotropic hormone (PTTH), a brain-derived neuropeptide that promotes proper ecdysone release [55]. The *ptth* gene is cyclically expressed throughout L3 development with a periodicity of approximately 8 hours [55]. Once the animal achieves critical weight, PTTH binds to the receptor tyrosine kinase Torso and activates a canonical mitogen-activated protein kinase (MAPK) signaling cascade [56]. This signaling event is essential for monitoring critical weight and for determining the timing of the onset of metamorphosis, as PTTH-dependent activation of MAPK signaling upregulates the expression of ecdysone biosynthetic genes [56]. If constitutively active forms of Torso, Ras, and Raf are expressed in the PG, animals precociously secrete ecdysone and pupariate early [56,57]. Similarly, both ecdysone release and pupariation are delayed when Torso, Ras, Raf, and Erk function are disrupted in the PG or if PTTH signaling is eliminated [56,57].

Interestingly, as in *C. elegans*, both TGF $\beta$  and insulin signaling feed into this hormone signaling pathway (Figure 3). When TGF $\beta$  signaling is reduced within the PG, *Torso* expression is significantly reduced and MAPK activity is downregulated [58]. As a result, the ecdysone biosynthetic genes are not properly expressed and the animals arrest development as L3. The observation that this phenotype is more severe than the elimination of PTTH signaling suggests that TGF $\beta$  can regulate ecdysone release through other pathways as well. At least one of these pathways appears to be insulin signaling [58]. Reduction of TGF $\beta$  signaling in the PG leads to reduced levels of *dInR* and reduced insulin signaling in this tissue. Moreover, expressing *dInR* or *dAkt* specifically in the PG is sufficient to overcome the block in larval development caused by reduced TGF $\beta$  signaling in this tissue [58]. This observation is consistent with several earlier studies that showed that activation of the insulin signaling pathway in the PG results in elevated ecdysone signaling and precocious initiation of metamorphosis [10,57,59]. Conversely, inhibiting insulin signaling within this organ dampens ecdysone signaling and extends larval growth. These results demonstrate that insulin and TGF $\beta$  signaling play a central role in coordinating growth with developmental progression and suggest that Dilp and TGF $\beta$  production or activity is sensitive to changes in body size. A key direction for future research will be to determine how the mechanisms that assess larval growth are linked to growth factor signaling.

It is interesting to note that a few studies in humans suggest that this pathway is conserved through evolution. Juvenile females diagnosed with type I diabetes mellitus exhibit a significant delay in menarche [60], while individuals with Marfan syndrome, which is likely caused by excessive TGF $\beta$  signaling, experience early onset puberty [61–63]. Thus, TGF $\beta$  and insulin signaling appear to control the timing of maturation in many higher organisms, defining a conserved genetic architecture that modulates steroid hormone signaling and the commitment to adult reproductive growth.

## Regulation of Post-Critical Weight Growth

The mid-L3 pulses of 20E correlate with dramatic changes in the larval growth program that allow development to progress independent of nutrient availability. This developmental transition stems, in part, from a fundamental change in the role of insulin signaling. Prior to mid-L3, insulin controls the rate of developmental progression and, when young larvae that harbor temperature-sensitive *dInR* alleles are raised at a non-permissive temperature, development is significantly delayed [64]. In contrast, when these mutants are shifted to a non-permissive temperature after mid-L3, depletion of insulin signaling produces smaller adults but does not affect larval development [64]. Similarly, ectopic expression of *dFOXO* before mid-L3 elicits a developmental delay, but expression during mid- to late-L3 only affects body size [29].

These fundamental changes in insulin signaling are likely a result of increased 20E activity after the mid-L3 transition. This hormone inhibits larval growth, with body size being significantly reduced in animals that are fed exogenous 20E [59]. Elevated 20E signaling does not, however, affect growth in *dFOXO* mutants, indicating that 20E regulates growth by antagonizing insulin signaling [59] (Figure 4). Intriguingly, fat-body-specific depletion of

*EcR* is sufficient to suppress the growth-inhibitory effects of 20E, while overexpression of the insulin signaling inhibitor dPTEN within the fat body has no effect on body size, demonstrating that ecdysone signaling in the fat body acts remotely to control organismal growth [59,65].

Although the mechanism that links systemic growth and insulin signaling with *EcR* activity in the fat body remains unclear, this interaction is due, in part, to 20E-dependent regulation of the *dMyc* transcription factor, which plays a key role in promoting growth (Figure 4). *EcR* activation leads to a decrease in *dMyc* protein levels while, conversely, *EcR* depletion in the fat body results in cell-autonomous induction of *dMyc* expression [65]. The effects of 20E on growth are, at least in part, dependent on *dMyc* because a reduction in *dMyc* expression can suppress the increased growth that results from decreased *EcR* activity [65]. Intriguingly, TOR also regulates *dMyc* expression [66], hinting at a model whereby nutrient sensing and 20E signaling are integrated to coordinate systemic growth.

The interaction between 20E and insulin signaling is not limited to growth, but also influences maturation. Within the fat body, *EcR* activation antagonizes cell-autonomous insulin signaling by interfering with PI3K activity, thereby causing *dFOXO* to translocate into the nucleus [38,59] (Figure 4). While *dFOXO* controls the expression of many genes, recent studies have demonstrated that the stage-specific upregulation of two of these target genes helps larvae prepare for metamorphosis. One of these factors is *dDOR*, which encodes a transcriptional coactivator that physically interacts with *EcR*, and which is required for proper 20E signaling [67]. The 20E-induced translocation of *dFOXO* into the nucleus promotes *dDOR* expression, further activating *EcR* signaling and initiating a feed-forward loop in fat body cells (Figure 4). In addition, both *EcR* and *dFOXO* transcriptionally upregulate *dilp6*, an insulin-like peptide produced by the fat body [68,69]. Although this finding may seem paradoxical to the growth-inhibitory effects normally associated with these two transcription factors, successful development requires some growth after the cessation of larval feeding and the onset of metamorphosis. Hormone-induced expression of *dilp6* allows animals to complete larval growth in the absence of external nutrients, likely by utilizing substrates that are released from the fat body and other larval tissues. In this manner, 20E and insulin act together to establish a transcriptional program that ensures successful completion of larval growth and development.

## Generating an Adult Body

Progenitors for the structures of the adult fly are carried inside the larva in the form of diploid imaginal tissues (ITs), which are specified during embryogenesis and go through several rounds of cell division during larval development. Perhaps not surprisingly, IT growth is influenced by many of the same humoral factors and cell-intrinsic signaling pathways that regulate growth in larval tissues [16,70]. Moreover, communication is maintained between IT growth and maturation to ensure that these tissues are ready for their terminal differentiation during pupal stages. If these tissues are damaged with X-rays or apoptosis, critical weight is increased and maturation is delayed while the tissue regenerates [71,72]. This delay is a direct result of restricting PTH production and 20E responses, due, at least in part, to a retinoid-dependent signal produced in response to damaged ITs [71].

Animals that are raised on a diet lacking the substrates required for retinoid synthesis or that harbor mutations that disrupt retinoid production are unable to delay larval development in response to IT damage [71]. This finding suggests that a retinoid-dependent mechanism helps synchronize IT growth with the onset of maturation.

The low-titer 20E pulses that correlate with the achievement of critical weight also produce changes in the ITs. Prior to this event, unliganded EcR, in complex with USP, represses gene expression in the ITs, reflecting a well-defined role for unliganded nuclear receptors to function as repressors [73,74]. The low-titer 20E pulses that follow the onset of critical weight relieve this repression. In the wing imaginal discs, for example, this switch is reflected by the expression patterns of the genes *cut* and *senseless*, which are repressed during early L3 stages, and are initially expressed in response to the 20E pulses during mid-L3 [75].

## Metabolism and Maturation

Many of the genetic pathways that control developmental growth in higher organisms — insulin, TOR signaling, and nuclear receptors — are also essential metabolic regulators. This relationship is conserved in flies, as mutations that disrupt insulin, TOR, or 20E signaling not only affect growth, but also produce metabolic phenotypes [26,67,76,77]. Development and metabolism, therefore, are inseparably linked, and current studies are focused on better defining the mechanisms that underlie this interaction.

Recent studies have shown that larvae adopt a unique metabolic program that efficiently converts nutrients into biomass to support growth. This growth program is regulated by a nuclear receptor, the *Drosophila* ortholog of mammalian estrogen-related receptor (dERR), which coordinately upregulates the transcription of genes encoding enzymes involved in glycolysis, the pentose phosphate pathway, and lactate production during mid-embryogenesis [78]. The resultant metabolic program, which lasts throughout larval development, is a form of aerobic glycolysis that has been demonstrated in normal proliferating cells and cancer cells to support their remarkable growth [79–81]. Although only previously considered in the context of cell proliferation, it is interesting that this program has been adapted to facilitate the dramatic increase in body mass that occurs during *Drosophila* larval stages.

Intriguingly, many of the genes that are upregulated by dERR at the onset of larval development are coordinately downregulated as the animal prepares for metamorphosis [82]. Although the mechanism that controls this metabolic transition remains unclear, the downregulation of *lactate dehydrogenase* (*ImpL3*) correlates with the transcriptional changes induced by 20E during mid-L3 [39]. A similar phenomenon occurs in the silkworm *Bombyx mori*, where 20E signaling downregulates expression of many of the genes that encode glycolytic enzymes [83]. Thus, in addition to defining the end of larval growth, 20E arrests the metabolic program that supports this process.

A few studies also suggest that 20E initiates distinct metabolic programs at the onset of metamorphosis that are directed toward utilizing stored forms of energy to allow proper growth and development during the non-feeding pupal stages. A central aspect of this



response is the discovery that 20E signals the onset of fat body autophagy during mid-L3 [37,38]. This is in parallel with the ability of 20E signaling to arrest growth through the fat body, mediated by insulin signaling [59], as well as an arrest of cell division in the ITs [84]. The developmentally programmed increase in fat body autophagy suggests that larvae utilize the nutrients that were stored earlier in development, which may explain how post-critical weight animals can develop independent of nutrient availability. This model is supported by observations in *Manduca sexta*, where the concentration of trehalose (the primary circulating sugar in insects) is depleted by starvation in pre-critical weight animals, but remains unaffected when animals are starved post-critical weight [85]. The metabolic transition that occurs at critical weight therefore allows the animal to develop independent of external nutrients and is due, in part, to the interplay between 20E and insulin.

## Perspectives and Future Directions

The discovery that evolutionarily conserved signaling systems regulate larval growth and development establish *Drosophila* as an ideal platform for exploring the basic principles of animal maturation. These findings, combined with the observation that the fat body plays a central role in sensing nutrients and coordinating organismal growth, have emerged at a time when childhood obesity is increasing at an alarming rate. A number of studies have described correlations between obesity and insulin resistance in children and the premature onset of female puberty, linking nutritional status to sexual maturation [86,87]. The onset of puberty is influenced by leptin, a hormone produced by the adipose tissue in response to fat synthesis [88]. The importance of fat storage in regulating maturation makes the identity of fat-body-derived growth regulators of special interest. In addition, identifying the factor(s) that regulate insulin secretion would represent a significant advance. This signal is dependent on TOR and therefore reflects a point at which nutrient availability can be integrated into the systemic growth program.

The role of nutrient sensing in controlling growth and maturation is not limited to the fat body, however, because TOR functions within the PG to regulate nutrient-dependent ecdysone release, although its interactions with PTTH, TGF $\beta$ , and insulin remain poorly understood. When TOR activity is inhibited in the PG, decreased 20E signaling results in delayed pupariation [89]. This effect appears to be the result of TOR-dependent ecdysone production, as reduced TOR signaling causes delayed induction of key ecdysone biosynthetic genes. These results imply a role for TOR in contributing to the genetic hierarchy that regulates the onset of maturation. This hypothesis is supported in *C. elegans*, where loss-of-function mutations in *let-363* and *daf-15*, the *C. elegans* homologs of TOR and the TOR complex component Raptor, respectively, lead to inappropriate dauer formation [90]. The phenotype of these *C. elegans* mutants, however, is morphologically distinct when compared with dauers produced by mutations in the insulin, TGF $\beta$ , or DA signaling pathways [44,90], suggesting that TOR regulation of dauer development is unique. Similarly, pharmacological manipulation of mTOR in the mouse brain suggests that TOR activity normally promotes maturation, although the mechanisms by which TOR regulates puberty remain largely unknown [88].

Finally, environmental factors other than nutrient availability also influence growth and maturation through poorly understood mechanisms. In particular, day length is likely to play a critical role in this process, as PTTH expression is influenced by circadian rhythms [55], and activation of insulin signaling in the PG renders the onset of metamorphosis sensitive to changes in light signals [10]. Furthermore, the neurons that comprise the central circadian clock in the brain directly innervate the PG and are located within close proximity to the PTTH-producing neurons, providing a way to achieve this control [55,91]. Intriguingly, secretion of mammalian gonadotropin-releasing hormone, a neuropeptide that is synthesized in the hypothalamus and is critical during puberty, appears to be regulated by circadian rhythms [92]. A similar relationship was also recently described in *C. elegans*, where LIN-42, the worm homolog of the circadian rhythm protein Period, genetically interacts with DAF-12 and DA signaling to regulate dauer formation [93]. The intersection between growth factor signaling, circadian rhythms, and maturation therefore appears to be conserved throughout animals. Further studies in simple systems such as *Drosophila* and *C. elegans* provide an opportunity to better define this poorly understood regulatory pathway.

## Conclusion

Even though Beadle and his colleagues [9] described *Drosophila* critical weight nearly 75 years ago, only now are we beginning to understand the genetic pathways that coordinate larval growth and maturation. The discovery of these developmental mechanisms appears to define a conserved genetic architecture that regulates juvenile growth and maturation throughout the animal kingdom and provides a new direction for understanding how human puberty and maturation are controlled.

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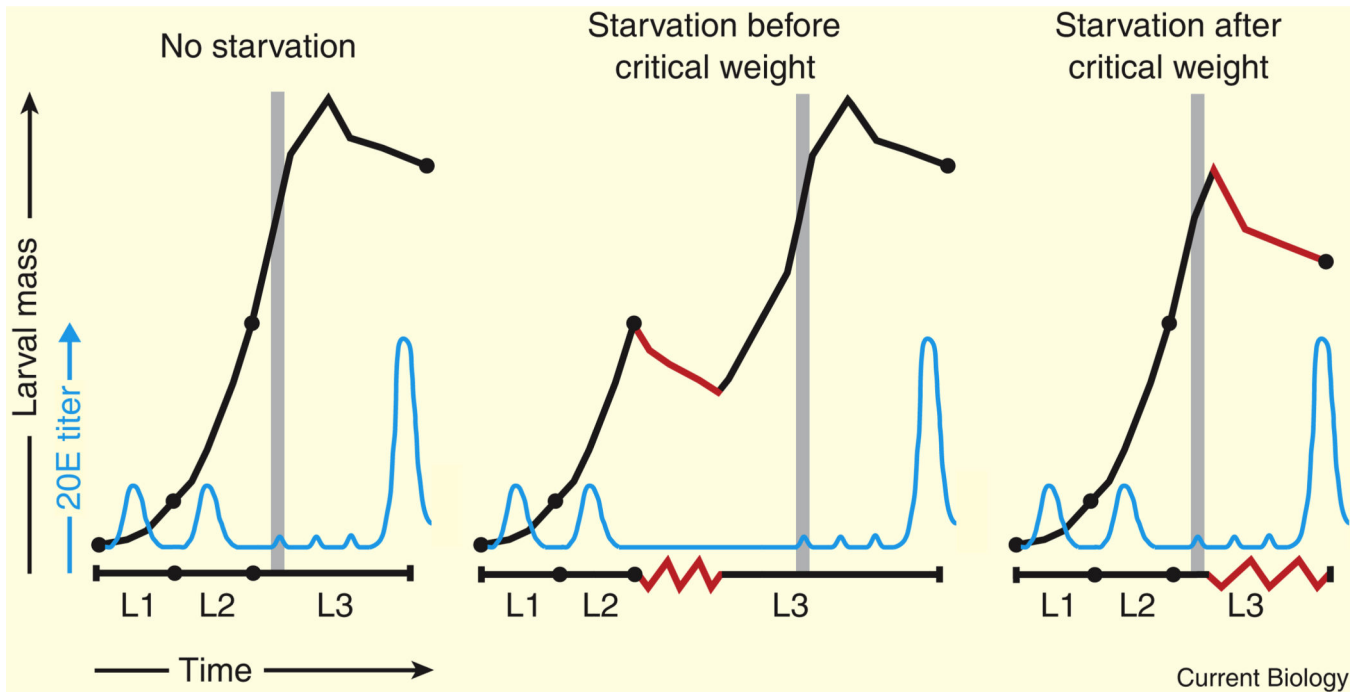
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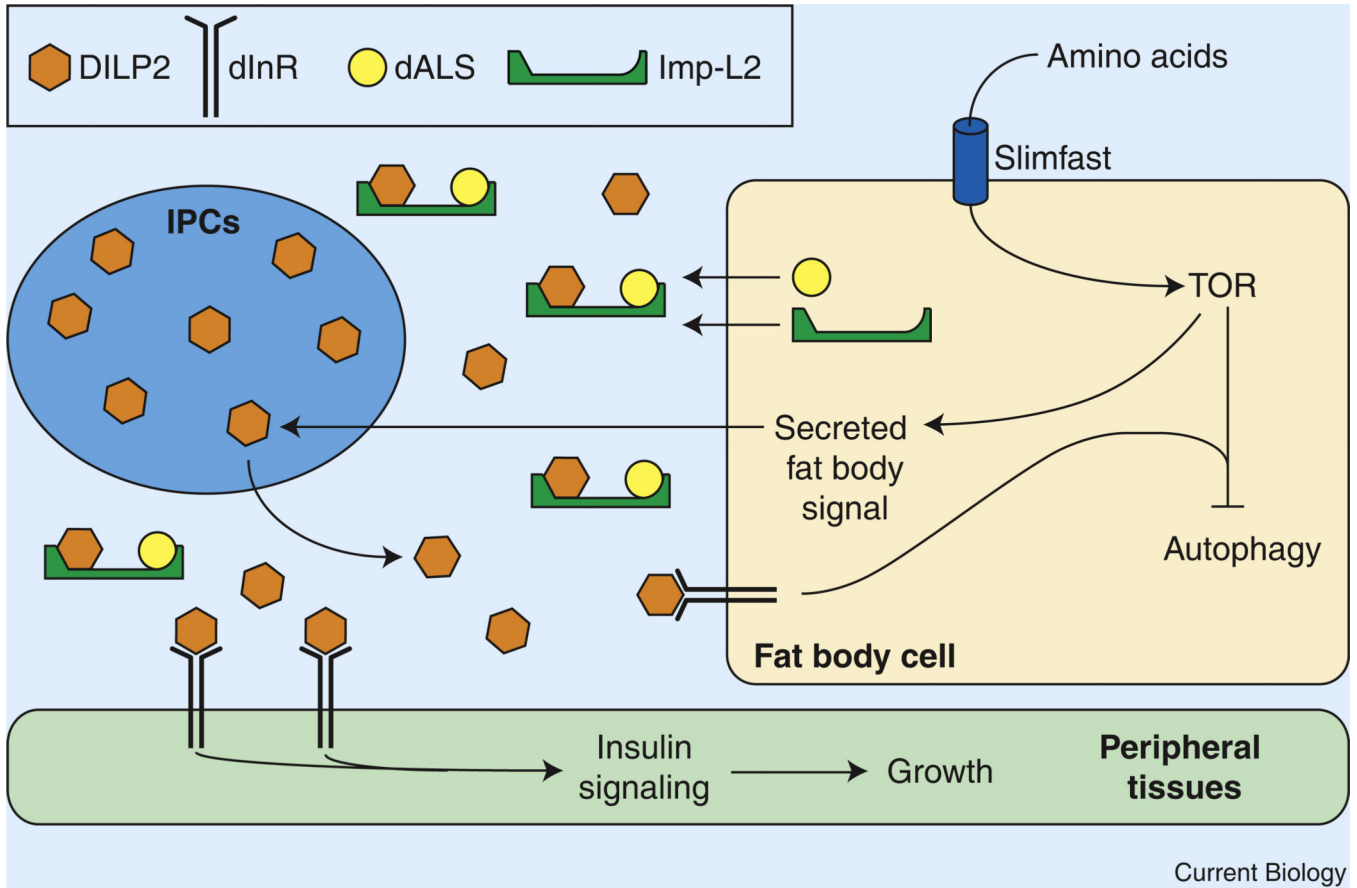
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**Figure 1. A schematic representation of *Drosophila* larval growth and development**

*Drosophila* larvae experience exponential growth (black line) as they develop through three distinct larval instars (L1, L2, and L3). Pulses of 20E (blue line) direct progression through the larval molts. The critical weight checkpoint (grey vertical line) occurs near the L2–L3 molt. A series of low-titer 20E pulses occur at ~8, 20, and 28 h after the L2–L3 molt, followed by a high-titer 20E pulse at the end of L3 that triggers puparium formation [7]. If an animal is starved (red lines) prior to the attainment of critical weight, development stalls until the larva finds a new food source, but final body size is unaffected. After critical weight is achieved, starvation inhibits growth but no longer affects developmental progression, resulting in a significantly smaller final body size.



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**Figure 2. The larval fat body regulates systemic growth**

Pre-critical weight larval growth is regulated by nutrient-dependent signals that emanate from the fat body. Ingested amino acids are sensed by fat body cells and activate TOR kinase, which promotes the release of an unknown factor that stimulates DILP2 secretion from the insulin-producing cells (IPCs). DILP2, in turn, promotes growth and development in peripheral tissues by binding to the insulin receptor (dInR) and activating the insulin signaling pathway. Additionally, the fat body releases ALS and Imp-L2, which form a stable complex with DILP2 and dampen insulin signaling.

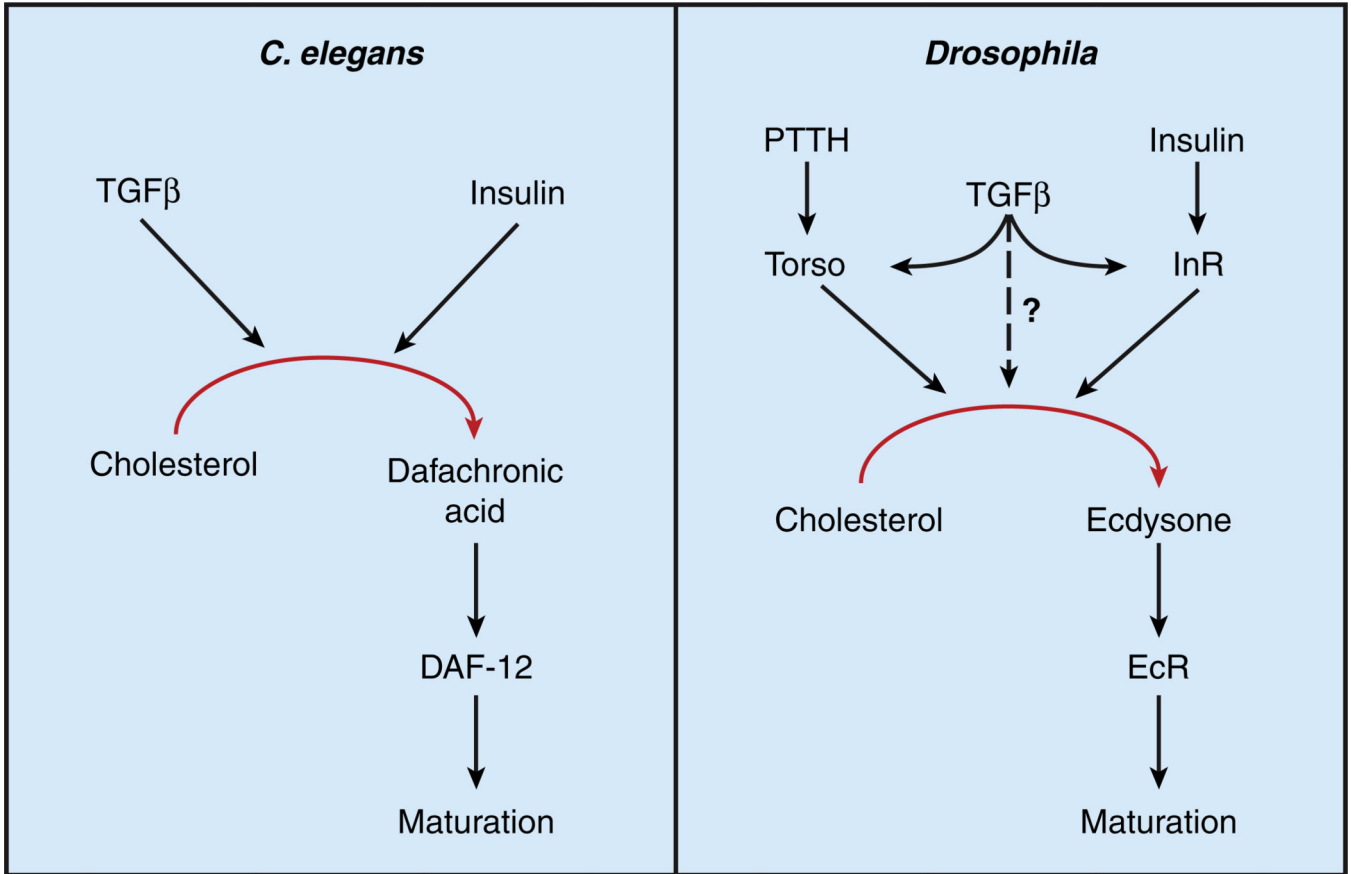
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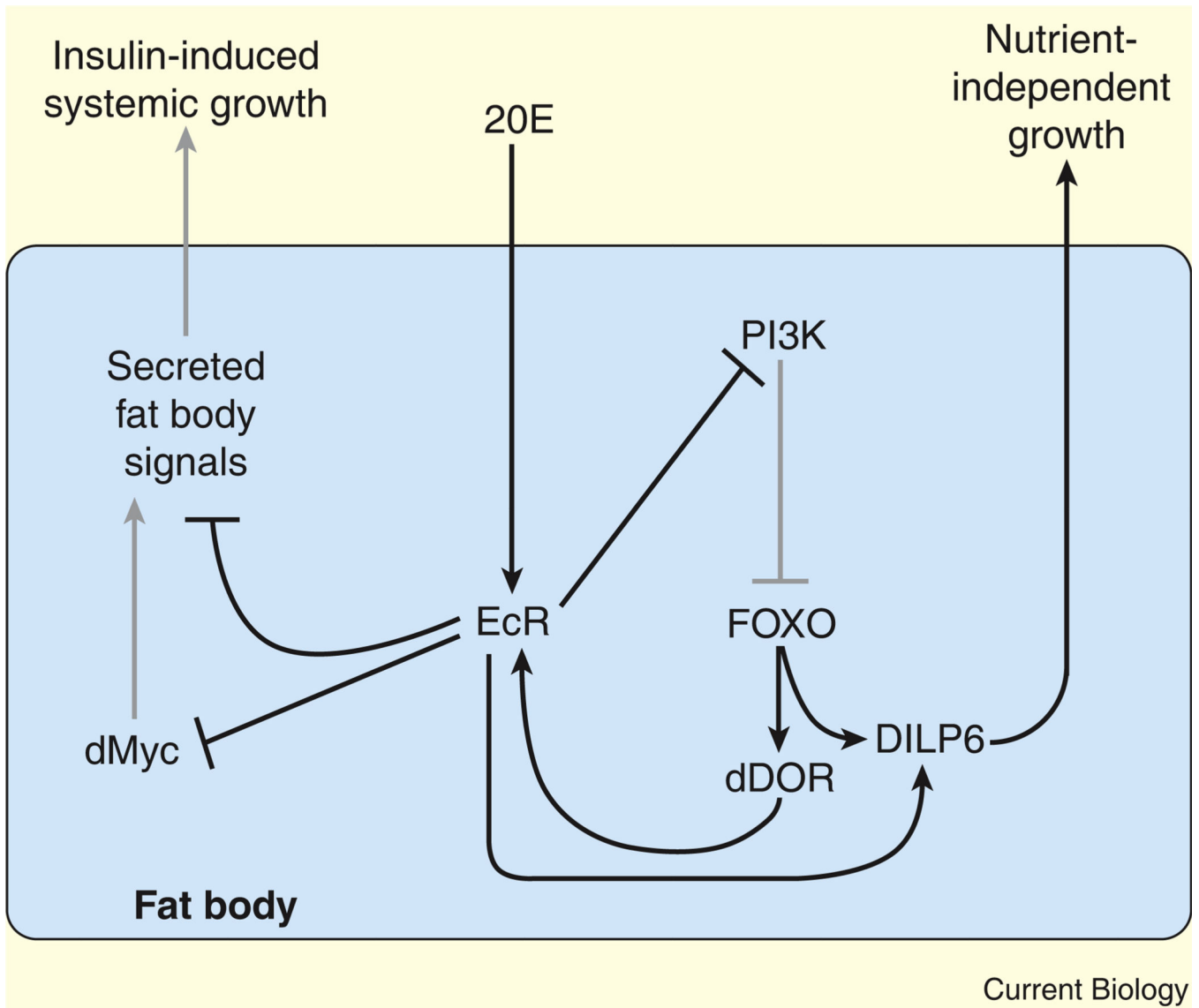




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**Figure 3. A conserved genetic hierarchy regulates animal maturation**

A combination of insulin and TGFβ signaling regulates steroid hormone production and maturation in both *C. elegans* and *Drosophila*. In worms, dietary nutrients and favorable growth conditions increase TGFβ and insulin signaling in endocrine tissues and stimulate dafachronic acid (DA) synthesis. DA systemically activates the nuclear receptor DAF-12, thereby preventing dauer formation and promoting maturation. In *Drosophila*, TGFβ signaling in the prothoracic gland (PG) upregulates expression of Torso and the insulin receptor (dInR), which promote ecdysone synthesis in response to PTTH and insulin, respectively. Ecdysone is then released from the PG, converted into 20E, and promotes maturation by systemically activating the ecdysone receptor (EcR).



**Figure 4. Ecdysone functions in the fat body to regulate systemic growth**

EcR activation by 20E in the fat body inhibits systemic insulin signaling and growth, in part by downregulating dMyc expression. EcR also inhibits PI3K signaling, which allows dFOXO to translocate to the nucleus and activate the expression of target genes. These include *DDOR* and *dilp6*, which is also upregulated by 20E–EcR. *DDOR* further activates EcR signaling, while *dilp6* promotes nutrient-independent growth. Grey lines represent genetic interactions that are downregulated by 20E signaling.