Developmental model of static allometry in holometabolous insects

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The regulation of static allometry is a fundamental developmental process, yet little is understood of the mechanisms that ensure organs scale correctly across a range of body sizes. Recent studies have revealed the physiological and genetic mechanisms that control nutritional variation in the final body and organ size in holometabolous insects. The implications these mechanisms have for the regulation of static allometry is, however, unknown. Here, we formulate a mathematical description of the nutritional control of body and organ size in \textit{Drosophila melanogaster} and use it to explore how the developmental regulators of size influence static allometry. The model suggests that the slope of nutritional static allometries, the ‘allometric coefficient’, is controlled by the relative sensitivity of an organ’s growth rate to changes in nutrition, and the relative duration of development when nutrition affects an organ’s final size. The model also predicts that, in order to maintain correct scaling, sensitivity to changes in nutrition varies among organs, and within organs through time. We present experimental data that support these predictions. By revealing how specific physiological and genetic regulators of size influence allometry, the model serves to identify developmental processes upon which evolution may act to alter scaling relationships.

Keywords: allometry; insulin; body-size regulation

\textbf{1. INTRODUCTION}

Biological diversity is dominated by variation in shape. This is perhaps no more apparent than in one of the most successful metazoan taxa, the insects. From a very simple body plan (head, thorax, abdomen, a pair of antenna, two pairs of wings and three pairs of legs) come myriad forms, from bees to beetles and fleas to flies. Much of this diversity is a result of variation in the relative, rather than the absolute, size of the insect body parts. The same is true for almost all other metazoan phyla. The question of how biological diversity is a result of variation in shape in insects. From a very simple body plan (head, thorax, abdomen, a pair of antenna, two pairs of wings and three pairs of legs) come myriad forms, from bees to beetles and fleas to flies. Much of this diversity is a result of variation in the relative, rather than the absolute, size of the insect body parts. The same is true for almost all other metazoan phyla. The question of how biological diversity is a result of variation in shape in holometabolous insects.

Allometric relationships, the relationship between the size of one trait and the size of another trait or the body as a whole, are traditionally modelled using the allometric equation

\begin{equation}
    y = bx^a,
\end{equation}

where \( x \) and \( y \) are the size of two given traits, respectively (Dubois 1897; Huxley & Tessier 1936). When \( x \) and \( y \) are the traits in conspecific individuals at the same developmental stage, the relationship is a static allometry (Schlichting & Pigliucci 1998; Shingleton et al. 2007). A log transformation of equation (1.1) produces a simple linear equation, \( \log(y) = \log(b) + a \log(x) \) and log–log plots of the size of different traits among individuals of the same species typically reveal linear allometries with an intercept of \( \log(b) \) and a slope of \( \alpha \), called the ‘allometric coefficient’ (Huxley & Tessier 1936). For morphological traits, as long as \( x \) and \( y \) are the same dimension, \( \alpha \) is often close to 1, such that the relative size of the two traits is constant irrespective of absolute size. This condition is called isometry (figure 1a). However, \( \alpha \) may also be less than or greater than 1, such that as the size of \( y \) becomes smaller or larger relative to \( x \) as both traits get absolutely larger. These conditions are called hypo- and hyperallometry, respectively (figure 1bc). The allometric equation therefore summarizes how the relative sizes of traits scale with each other and with the overall body size, and hence capture the relationship between size and shape in complex organisms. Not all static allometries are linear, however. They may be sigmoidal or discontinuous depending on the trait, the species or the unit of measurement (Emlen & Nijhout 2000). Nevertheless, in this paper, we concentrate on those static allometries that are linear on a log–log scale.

Despite the extensive application of the allometric equation to the traits of organisms from almost every phylum, very little is understood of the developmental processes that produce linear allometries, and how these processes can be modified to generate the diversity of scaling relationships we see both within and between species (Stern & Emlen 1999; Shingleton et al. 2007). Huxley recognized that the allometric coefficient \( \alpha \) could represent the differential growth rate of two traits growing simultaneously. That is, \( \alpha = m/n \), where \( m \) and \( n \) are the growth rates of the two traits (Huxley 1924; Shea 1985). In such cases, the static allometry follows the growth trajectories of the two structures and different points on the static allometry represent individuals that grow for different periods of time. However, the growth of different structures is not always synchronous. Some structures, for example, the antlers on deer, grow only after the rest of
body has stopped growing. Others, for example, the adult organs of holometabolous insects, have growth periods that only partially overlap with each other and with the growth period of the body (Williams 1980). Under such circumstances, it is unclear what developmental mechanisms regulate the slope and intercept of allometric relationships between traits (Nijhout & Wheeler 1996).

Recent elucidations of the developmental regulation of body and organ size in holometabolous insects provide the first clues as to how static allometries may be controlled (for review see Shingleton et al. 2007). These mechanisms concern the nutritional regulation of growth and so are particularly applicable to our understanding of nutritional static allometries; that is, static allometries where variation in trait size is a consequence of variation in developmental nutrition. Nutritional status is a major regulator of body and organ size in animals (Oldham et al. 2000a) and so the mechanisms that control the developmental response to nutrition probably underlie many of the static allometries observed in nature.

Here, we develop a mathematical model of nutritional static allometry regulation in holometabolous insects. The physiological and genetic mechanisms that underlie the nutritional regulation of final body and organ size in holometabolous insects have previously been proposed by Shingleton et al. (2005, 2007) and Nijhout et al. (2006). Because static allometry describes the scaling relationship between final body and organ size, the mechanisms that regulate final body and organ size necessarily regulate allometry and, if correct, should be able to explain the observed allometric relationships. However, the impact that specific aspects of these mechanisms have on allometry expression are difficult to evaluate using verbal arguments and thought experiments alone. Mathematical modelling allows us to investigate how the physiological and genetic regulators of body size interact with the physiological and genetic regulators of organ size, and whether the hypothesized interaction can generate biologically reasonable allometries. In this paper, we first summarize the features of the mechanisms regulating final body and organ size before we use them to formulate a quantitative model of allometry regulation. The timing and dynamics of organ growth are perhaps best understood in the fruit fly Drosophila, and so we will develop our model using Drosophila as an example holometabolous insect.

### 2. Nutritional Regulation of Body Size in Drosophila

The mechanism of the nutritional regulation of body size in Drosophila is illustrated in figure 1. In fruit flies, as in all holometabolous insects, growth is restricted to the larval stages (Bakker 1959; Ashburner 1989). A developing larva feeds and grows exponentially until it reaches a critical size at the beginning of the third larval instar (figure 2a(i)) (Nijhout & Williams 1974b; Nijhout 2003; Mirth et al. 2005). By an incompletely known mechanism, attainment of critical size initiates a hormonal cascade (figure 2a(ii)) that ultimately results in the release of ecdysteroids (figure 2a(iii)) (Baehrecke 1996). The subsequent rise in ecdysteroids causes the larva to stop feeding and wander away from its food to find a place to metamorphose, called the wandering stage (Berreur et al. 1979). The cessation of feeding marks the end of body growth and, because adults possess an exoskeleton that prohibits further growth, fixes the maximum size of the adult fly (figure 2a(iv)). There is temporal separation between the attainment of critical size and the cessation of feeding and growth, called the terminal growth period (TGP) (Shingleton et al. 2007). The final size of the fly is thus determined by (i) critical size, plus the amount of growth achieved in the TGP, which is in turn determined by (ii) the duration of the TGP and (iii) the rate of growth during the TGP (Shingleton et al. 2007). Variation in developmental nutrition potentially affects final body size by influencing any or all of these three factors. However, nutrition does not appear to regulate critical size in Drosophila (De Moed et al. 1999; Shingleton et al. 2005; A. W. Shingleton, unpublished data), and has only a small effect on the duration of the body’s TGP, with complete starvation causing slightly precocious pupariation (Beadle et al. 1938; Mirth et al. 2005). Rather, nutrition primarily influences final body size by regulating the body’s rate of growth during its TGP (figure 2b).

It is the insulin/TOR-signaling (IIS) pathway that coordinates growth rate with nutritional condition in Drosophila and most metazoans (for review see Conlon & Raff 1999; Oldham et al. 2000a; Edgar 2006). In Drosophila nutrition regulates the release of insulin-like peptides (dILPs) from various tissues around the body (Brogiolo et al. 2001; Britton et al. 2002; Rulifson et al. 2002). These dILPs bind to the insulin receptor (Inr) of dividing cells and, through a well-elucidated signal transduction pathway, regulate the rate of cell growth and proliferation (Edgar 2006). Starvation down-regulates the IIS-pathway through several mechanisms. A reduction in circulating nutrients can be sensed directly by dividing cells, leading to suppression of the IIS-pathway via the target-of-rapamycin (TOR) (Oldham et al. 2000b; Neufeld 2003). Starvation also reduces the release of dILPs, down-regulating the IIS-pathway via Inr (Ikeya et al. 2002). Finally, other hormones, for example

![Figure 1](image-url)
juvenile hormone (JH) (Truman et al. 2006), ecdysone (Caldwell et al. 2005; Colombani et al. 2005; Mirth et al. 2005) and imaginal disc growth factors (IDGF) (Kawamura et al. 1999), also regulate the IIS-pathway, and may themselves be nutritionally regulated. Collectively, these various humoral factors are likely dispersed uniformly throughout the developing larva, and so the IIS-pathway can coordinate growth throughout the body in response to changing nutrition (Goberdhan & Wilson 2002).

3. NUTRITIONAL REGULATION OF ORGAN SIZE IN DROSOPHILA

The mechanism of the nutritional regulation of organ size, also illustrated in figure 2, is less well elucidated than for body size. In holometabolous insects like Drosophila, the adult organs develop as imaginal discs within the growing larva. Although the cells that will become the discs are specified during embryogenesis, the point in development when discs initiate growth varies among discs, with most discs starting growth in the first or second instar (Madhavan & Schneiderman 1977; Cohen 1993). Like the body, growth of the discs is approximately exponential (Martin 1982; Bryant & Levinson 1985) and ends with cell differentiation. Growth cessation appears to be regulated by the same hormonal cascade that is initiated upon attainment of critical size (Emlen & Allen 2003; Mirth 2005). Differentiation in the eye discs is regulated by the rise in ecdysteroids that causes larvae to stop feeding (Brennan et al. 1998), and differentiation in the wing and leg is also tied to changes in the levels of circulating ecdysteroids and JH (Martin & Shearn 1980; Fristrom & Fristrom 1993; Riddiford 1993). Like the body, therefore, the imaginal discs also have TGPs that begin when a larva reaches critical size (figure 2a(i)) and end when hormone levels rise above a certain threshold (figure 2a(v),(vi)). However, the discs have different sensitivities to the hormone cascade initiated at critical size, and so their TGPs are typically longer than the TGP of the body (Garcia-Bellido & Merriam 1971; Freeman 1997).

Final organ size is therefore determined by (i) the size of the imaginal discs at critical size, (ii) the duration of the disc’s TGP and (iii) the rate of growth during that TGP (Shingleton et al. 2007). Again, nutrition could affect final body size by influencing any or all of these three factors. However, the size of the discs at critical size does not appear to be affected by nutrition or changes in IIS (Shingleton et al. 2005). Further, since nutrition does not appear to substantially influence the body’s TGP, it seems unlikely that it substantially affects the TGPs of the imaginal discs, since all TGPs are controlled by the same hormonal cascade. As with the body as a whole, therefore, nutrition appears to influence final organ size primarily by regulating the rate of disc growth during the discs’ TGP via the IIS-pathway (figure 2b) (Shingleton et al. 2005, 2007).

4. A MODEL OF NUTRITIONAL STATIC ALLOMETRY EXPRESSION IN DROSOPHILA

In order to better understand the implications the mechanistic description of body and organ size regulation has for the control of static allometry (figure 2), we have formalized the mechanism into a mathematical model. The model predicts final body and organ size for a fly experiencing a constant but non-zero level of nutrition throughout development. The model assumes that this constant level of nutrition translates into a constant level of IIS. Setting different levels of IIS alters the growth rate of the body and organs during their respective TGPs and generates a variation in body and organ size. Consequently, the model can be used to explore how the body and organs scale under different nutritional conditions, and how this allometric relationship is influenced by the model’s biological parameters.

Because imaginal disc growth trajectories have been best elucidated in the wing (Garcia-Bellido & Merriam 1971; Madhavan & Schneiderman 1977; Martin 1982; Bryant & Levinson 1985; Milan et al. 1996), we will use the wing as an example imaginal disc. Nevertheless, the
parameters of the model can be changed to explore the allometric regulation of other discs.

(a) Exponential growth

The growth of the body (Bakker 1959) and imaginal discs (Martin 1982; Bryant & Levinson 1985) is approximately exponential in Drosophila, as is typical for insects (Nijhout et al. 2006). Changes in nutrition change the rate of growth, and in our model we will assume that this occurs exclusively through the IIS pathway and the pathways the IIS pathway interacts with. We will not assume a priori that all growth is affected by changes in nutrition. Rather, the growth rate is divided into two components: an insulin-sensitive growth rate and an insulin-insensitive, or 'intrinsic', growth rate. The insulin-sensitive growth rate is modified by the level of IIS, which here refers to a combination of the level of circulating dILPs, nutrients and other humoral factors that regulate the IIS pathway. In contrast, the intrinsic growth rate does not vary with the nutritional level and occurs as long as the larva has sufficient nutrients to sustain life. This does not mean that intrinsic growth is not dependent on nutrition, since a lack of nutrition precludes growth. Rather, the intrinsic growth rate does not vary with the level of nutrition.

Growth of the body is modelled as

\[ dB/dt = (s_b + s_i \cdot i) \cdot B_i, \]  

(4.1)

and growth of the disc is modelled as

\[ dD/dt = (s_d + s_i \cdot i) \cdot D_i, \]  

(4.2)

where \( B_i \) and \( D_i \) are the size of the body and disc, respectively, at time \( t \); \( c \) is the intrinsic growth rate; \( s \) is the insulin-sensitive growth rate; and \( i \) is an index of the level of IIS, which varies between 0 and 1. Growth rate \((c + s \cdot i)\) therefore varies between \( c \), when IIS is minimal and \( i = 0 \), and \( c + s \) when IIS is maximal and \( i = 1 \). The parameter \( s \) thus captures the extent to which growth rate changes with the changes in IIS, and can be thought of as a measure of 'insulin sensitivity'. Both \( c \) and \( s \) can be the growth rate in any dimension (length, area and volume) and in any unit (cell number, mm, μg, etc.).

(b) Growth rate and final body size

In Drosophila, the body’s growth rate is not constant throughout development. Growth of the body (dry body mass) is exponential both before and after critical size, but the exponent of growth is lower once critical size is attained (Bakker 1959). Growth rate also appears to show a decline immediately before growth stops at the beginning of the wandering larval stage (Bakker 1959). Because this decline is relatively slight, however, we will approximate body growth as two exponential growth periods, the first from hatching to critical size and the second from critical size to the beginning of the wandering stage (figure 3). We will model growth rate of the body before critical size as in equation (4.1) and growth rate after critical size as

\[ dB/dt = (c_d + s_d \cdot i) \cdot B_i, \]  

(4.3)

where \( c' \) and \( s' \) are the intrinsic and insulin-sensitive growth rates after critical size.

Final body size is the critical size plus the amount of growth achieved during the body’s TGP (figure 2). Neither critical size nor TGP are substantially affected by changes in nutrition. Final body size can therefore be modelled by solving equation (4.3) for \( t = T_B \), where \( T_B \) is the duration of the body’s TGP, i.e. the time between the attainment of critical size and the beginning of the wandering stage. Thus,

\[ B_F = L \cdot e^{(c_c + s_c \cdot i) \cdot T_B}, \]  

(4.4)

where \( B_F \) is final body size under level of IIS \( i \) and \( L \) is the critical size.

(c) Growth rate and final disc size

Growth of the wing imaginal discs is also not constant throughout development. The wing discs begin growth midway through the first larval instar but the growth slows after the attainment of critical size and stops temporarily at pupariation (Bryant & Levinson 1985). Cell division resumes after the wing discs have everted (Garcia-Bellido & Merriam 1971; Milan et al. 1996) before permanently stopping when the wing discs differentiate, 24 hours after pupariation at 25°C (Milan et al. 1996). This later stage of growth has not, however, been well characterized. Further, the size of the wing disc at pupariation is thought to be the major determinant of final wing size (Day & Lawrence 2000). We will therefore only model wing disc growth up to the point of pupariation.

Like the body, wing disc growth can be considered as two exponential growth periods. The first is from midway through the first larval instar to critical size, the duration of which depends on the rate of the body’s growth to
critical size, and the second is from after critical size to pupariation (figure 3). We will describe growth rate of the disc before critical size using equation (4.2) and growth rate after critical size as

\[
dD/dt = (c_i + s_i \cdot i) \cdot D_i,
\]

where \(c_i\) and \(s_i\) are the intrinsic and insulin-sensitive growth rates after critical size, respectively. We will model the final wing disc size by analysing each of these growth periods separately.

Discs begin their growth after the body has started growing (Cohen 1993). It is not clear what initiates disc growth but a reasonable hypothesis is that it begins when the body reaches a certain size. In our model, disc growth begins when the body reaches proportion \(a\) of critical size \(L\) at time \(T_2\) after the hatching of the embryo to larva. The value of \(T_2\) therefore depends on the body’s growth rate up to \(aL\). We can model this by solving equation (4.2) for \(t = T_2\)

\[
aL = B_0 \cdot e^{(c_b + s_b \cdot i) \cdot T_2},
\]

where \(B_0\) is the size of the body at hatching.

It follows that

\[
T_2 = \frac{1}{(c_b + s_b \cdot i)} \cdot \ln \left( \frac{aL}{B_0} \right).
\]

By the same argument, critical size \((L)\) is reached at time \(T_i\) after the larva hatches, such that

\[
T_i = \frac{1}{(c_b + s_b \cdot i)} \cdot \ln \left( \frac{L}{B_0} \right).
\]

Again, \(T_i\) depends on the body’s growth rate up to \(L\).

We can model the size of the disc at critical size at a given level of insulin signalling by solving equation (4.2) for \(t = T_1 - T_2\), that is the amount of time between the beginning of disc growth and the attainment of critical size, such that

\[
D_L = D_0 \cdot e^{(c_d + s_d \cdot i) \cdot (T_1 - T_2)},
\]

where \(D_L\) is the size of the disc at critical size and \(D_0\) is the size of the disc at the beginning of growth. Inserting the values for \(T_1\) and \(T_2\) (equations (4.7) and (4.8), respectively) into equation (4.9), the size of the disc at critical size is

\[
D_L = D_0 \cdot e^{-(c_d + s_d \cdot i) \cdot (aL + s_b \cdot i)}.
\]

Once critical size is reached the disc continues to grow through its TGP. Because the disc’s TGP is probably regulated by the same hormone fluctuations that regulate the body’s TGP \((T_0)\), we can hypothesize that the former is some multiple of the latter. Final disc size can be modelled by solving equation (4.5) for \(t = T_0\)

\[
D_F = D_L \cdot e^{(c_d + s_d \cdot i) \cdot K \cdot T_0},
\]

where \(D_F\) is the final size of the disc and \(K\) is the factor relating the disc’s TGP to the body’s TGP. Inserting the values of \(D_L\) gives the final disc size as

\[
D_F = D_0 \cdot e^{-(c_d + s_d \cdot i) \cdot (aL + s_b \cdot i)} \cdot e^{(c_d + s_d \cdot i) \cdot K \cdot T_0}.
\]

(d) Fitting parameter values: growth under optimal conditions

The parameters of the model are real biological parameters and their values can be determined empirically. Several authors have published data for the growth of the body and wing imaginal discs of wild-type Drosophila reared on 100% food at 25°C (Bakker 1959; Madhavan & Schneiderman 1977; Martin 1982; Bryant & Levinson 1985). We have used these data to fit approximate parameter values to our model (table 1). Because the data were collected from flies reared under optimal conditions, we will assume that for these data the level of insulin signalling is maximal and that \(i = 1\).

Newly hatched larvae weigh 0.0022 mg (dry weight) and grow exponentially up to the critical size (Bakker 1959). The exponent of growth before critical size \((c_b + s_b) = 0.088\) (Bakker 1959). Critical size \((L)\) is 0.21 mg and is attained 53 hours after hatching (Bakker 1959). After critical size, the growth remains exponential but the rate is approximately halved, such that the exponent \((c_d + s_d) = 0.041\) (Bakker 1959). Growth continues until larval stop feeding, 74 hours after hatching (Bakker 1959). The TGP of the body \((T_0)\) is therefore 21 hours.

The wing imaginal discs initially constitute approximately 38 cells at hatching, and start growing 15 hours later (Madhavan & Schneiderman 1977). At this point, the body size is approximately 0.008 mg (Bakker 1959), hence \(a = 0.035\). Like the body, the growth of the wing discs appears to slow down slightly after critical size. The exponent of growth before critical size \((c_d + s_d)\) is 0.11, and the exponent after critical size \((c_d + s_d)\) is 0.075 (Bryant & Levinson 1985). Cell proliferation pauses at pupariation (93 hours, \(K = 1.9\)) when the wing discs constitute approximately 45–50 000 cells (Bryant & Levinson 1985).

### Table 1. Parameters used in the model and their values, if known, from published data.

<table>
<thead>
<tr>
<th>parameter</th>
<th>meaning</th>
<th>parameter value</th>
</tr>
</thead>
<tbody>
<tr>
<td>(B_0/D_0)</td>
<td>initial body/disc size (mg and cell number, respectively)</td>
<td>0.0022 38</td>
</tr>
<tr>
<td>(c)</td>
<td>intrinsic (insulin-insensitive) growth rate</td>
<td></td>
</tr>
<tr>
<td>(s)</td>
<td>insulin-sensitive growth rate</td>
<td></td>
</tr>
<tr>
<td>(i)</td>
<td>level of insulin signalling (ranges from 0 to 1)</td>
<td></td>
</tr>
<tr>
<td>(c + s)</td>
<td>growth rate under optimal conditions ((i = 1)), before critical size</td>
<td>0.088 0.11</td>
</tr>
<tr>
<td>(c' + s)</td>
<td>growth rate under optimal conditions ((i = 1)), after critical size</td>
<td>0.041 0.075</td>
</tr>
<tr>
<td>(L)</td>
<td>critical size (mg)</td>
<td>0.21</td>
</tr>
<tr>
<td>(T_0)</td>
<td>body’s TGP</td>
<td>21</td>
</tr>
<tr>
<td>(T_1)</td>
<td>time from hatch to attainment of critical size</td>
<td></td>
</tr>
<tr>
<td>(T_2)</td>
<td>time from hatch to initiation of disc growth</td>
<td></td>
</tr>
<tr>
<td>(a)</td>
<td>body size (as proportion of (L)) at which disc starts growing</td>
<td>0.035</td>
</tr>
<tr>
<td>(K)</td>
<td>disc’s TGP (as multiple of (T_0))</td>
<td>1.9</td>
</tr>
</tbody>
</table>
activity (electronic supplementary material, figure S1). This suggests that the body has no intrinsic growth rate and that \(c_{\text{b}} = 0.088\) and \(c_{\text{b}}' = 0.041\).

We can extend these assumptions to the wing. Under maximal nutritional conditions \((i = 1)\), the wing’s growth rate is 0.11 before critical size \((c_{\text{d}}' + s_{\text{d}}')\) and 0.075 after critical size \((c_{\text{d}}' + s_{\text{d}}')\). If the wing also has no intrinsic growth rate, that is \(c_{\text{d}} = c_{\text{d}}' = 0\), it follows that \(s_{\text{d}} = 0.11\) and \(s_{\text{d}}' = 0.075\) (Bryant & Levinson 1985). Using these parameter values, the model predicts a highly hyperallometric relationship between wing-cell number and body size as \(i\) varies, with an allometric coefficient \((\alpha)\) of 3.5 (figure 4a). However, the observed nutritional static allometry between adult wing cell number and body size has an allometric coefficient of 0.43 (figure 4b). The incompatibility of our model with the observed allometry suggests that our assumption that growth of the wings is entirely insulin-sensitive is incorrect. Consequently, the wings may have an intrinsic growth rate, either before or after the attainment of critical size.

The model suggests that the wings do not have intrinsic growth before attainment of critical size. When \(c_{\text{d}} > 0\) the body:wing allometry is nonlinear (figure 5c). This is because at very low levels of IIS, the time to critical size is long (electronic supplementary material, figure S2). If wing discs grow even when IIS is low, this developmental delay will lead to an increase in the disc size at critical size. This compensates for slow growth of the discs during their TGP and results in larger wings at low compared with intermediate levels of IIS (figure 5c). However, the allometric relationship between wing and body size created by rearing flies under different nutritional conditions is positive and linear (figure 4b); flies reared on low-quality diets, and those experiencing low levels of IIS, do not have larger wings than those reared on intermediate and high-quality diets. This suggests that the wings do not have intrinsic growth before the attainment of critical size.

By contrast, the model does generate a linear static allometry when the wings have an intrinsic growth rate after critical size; that is \(c_{\text{d}}' > 0\). It can be shown (electronic supplementary material) that when \(c_{\text{b}}\) and \(c_{\text{d}} = 0\) the allometric coefficient \((\alpha)\) between body and organ size can be modelled as
\[
\alpha = \frac{s_{\text{d}}'}{s_{\text{b}}'} \cdot K.
\] (4.13)

For an allometric coefficient of 0.43, if \(K = 1.9\) (table 1) then \(s_{\text{d}}' = 0.226 \times s_{\text{b}}'.\) Thus to produce the observed hypoallometry between wing cell number and body size, the insulin-sensitive growth rate of the discs must be considerably less than that of the body: if \(s_{\text{b}}' = 0.041\), then \(s_{\text{d}}' = 0.009.\) However, published data indicate that under optimal nutritional conditions \((i = 1)\), the exponent of wing growth \((c_{\text{d}}' + s_{\text{d}}') = 0.075\) (Bryant & Levinson 1985). This suggests that \(c_{\text{d}}' = 0.066\) and that the wing does have an intrinsic growth rate after the attainment of critical size.

Equation (4.13) can be generalized to apply to any two organs, neither of which is the body, again assuming that neither the organs nor the body have an intrinsic growth rate prior to critical size. The more general equation is
\[
\alpha_p = \frac{s_{\text{d}}'}{s_{\text{b}}'} \cdot \frac{K_p}{K}\] (4.14)
where $a_{pq}$ is the allometric coefficient for traits $p$ and $q$. For example, the relationship between the size of the maxillary palp and the male genitals of *Drosophila* is highly hypoallometric (figure 1), such that $\alpha = 0.35–0.5$, depending on the genotype. This suggests that either the genitals have a growth rate that is relatively insensitive to changes in IIS compared with other organs, or that the genitals’ TGP is relatively short compared with other organs.

Equation (4.13) suggests that the parameters that primarily influence an organ’s allometric coefficient with body size are its post-critical size insulin-sensitive growth rate ($s'_d$) relative to that of the body ($s'_b$) and the relative duration of its TGP ($K$). Using different values of $s'_b$, $s'_d$ and $K$ in the model generates a range of allometries with different slopes (figure 5a). Changes in $K$, while slightly affecting the slope of the allometry, also affect the minimal disc size and so have a more dramatic effect on the allometry’s intercept. By contrast, changes in $s'_d$ and $s'_b$ have a substantial effect on the allometry’s slope, but do not affect the minimal body or disc size and so have little effect on the allometry’s intercept. Changes in when the imaginal discs start growing prior to critical size ($a$), the insulin-sensitive growth rate of the disc and the body before critical size ($s_d$ and $s_b$ respectively), critical size ($L$), the disc’s intrinsic growth rate after critical size ($c'_d$) all influence the intercept but not the slope of the allometry (figure 5b–d).

**5. DISCUSSION**

Simple allometry, modelled as $y = bx^\alpha$, is widely applicable to scaling relationships in myriad animals. Nevertheless, the biological meaning of the equation’s parameters, in particular the allometric coefficient $\alpha$, is unclear for any animal except those whose body parts grow synchronously (Nijhout & Wheeler 1996). Many organisms do not grow synchronously, including holometabolous insects. Rather, different structures grow at different periods. In such situations, it is not immediately obvious what the biological regulators of $\alpha$ are.

Our simple model of nutritional static allometry in holometabolous insects suggest that $\alpha$ is regulated by the relative sensitivity of an organ’s growth rate to changes in IIS and the duration of development when IIS affects the final organ size (equation (4.13)). This has an intuitive appeal. The slope of a static allometry captures the extent to which factors that affect body size also affect organ size. For example, an organ will be hyperallometric to body size if the conditions that alter body size have a greater affect on organ size. For both the organs and the body, their insulin-sensitive growth rates and their TGP will affect how much their final size is affected by the changes in IIS. Thus, the greater an organ’s insulin-sensitive growth rate is compared with the body’s ($s'_d/s'_b$), or the longer its relative TGP ($K$), the more the organ’s final size will be affected by changes in IIS relative to the body, and the more hyperallometric will be its relationship to body size.

**Allometry and insulin sensitivity**

The possibility that different discs differ in their growth response to IIS has been hypothesized before (Nijhout 2003; Emlen et al. 2007; Shingleton et al. 2007). Reduced
nutrition and IIS produce smaller individuals, yet with ostensibly normal proportions (except for certain organs such as the male genitals (Shingleton et al. 2005)). Proportionality is maintained if each organ grows at a particular rate relative to each other (Nijhout 2003). However, all organs are probably exposed to the same level of circulating growth factors. Consequently, for different organs to maintain the correct relative growth rate, they must grow at different rates in response to a particular level of circulating insulin (Nijhout 2003). Our model suggests an additional nuance to this observation. Two organs may have very different growth rates under the same level of IIS, but will only maintain proportionality as IIS varies if they share the same level of insulin-sensitive growth and the same TGP. Thus, there are two aspects to a disc’s response to IIS: the rate at which a disc grows at a particular level of circulating insulin and how this rate changes with changes in IIS. This latter aspect is captured by the insulin-sensitive growth rate $s$ in our model, and we restrict the term insulin sensitivity to the magnitude of $s$.

The model suggests that if two discs differ in either their insulin sensitivity or in their TGP then, all other things being equal, their scaling relationship will not be isometric. Conversely, if two organs are isometric, but differ in their TGP, then they should also differ in their insulin sensitivity. The point in development when a disc attains critical size almost eliminates body growth (electronic supplementary material, figure S1). By contrast, the wing discs continue to grow until final disc size is attained after attainment of critical size exhibit no body growth but their discs continue to grow until final disc size is appropriate for the much reduced body size.

(b) Allometry and intrinsic growth

The model predicts that the effect of IIS on disc growth varies through development, for wing discs at least: before attainment of critical size, and when changes in nutrition can delay development, the model suggests that the growth rates of the body and the wing discs are entirely insulin-sensitive. After attainment of critical size, when the final duration of growth is fixed, the model suggests that there is a decline in the wing discs’ insulin-sensitive growth rate, with the discs taking on an intrinsic growth rate in order to sustain their overall rate of growth. Changing a disc’s post-critical size intrinsic growth rate alters the intercept but not the slope of its allometry (figure 5c), while changing the disc’s insulin sensitivity alters the slope but not the intercept (figure 5a). Consequently, separating intrinsic from insulin-sensitive growth may allow discs to regulate the slope of their allometry independent of the intercept of their allometry. We might therefore expect intrinsic growth to be a common feature of imaginal discs in holometabolous insects.

There is good experimental evidence that discs in holometabolous insects have an intrinsic growth rate. It is possible to manipulate the IIS pathway in developing larvae using a temperature-sensitive mutation of the Inr (Shingleton et al. 2005). Switching off the IIS pathway in all tissues after attainment of critical size almost eliminates body growth (electronic supplementary material, figure S1). By contrast, the wing discs continue to grow (electronic supplementary material figure S1, figure 6), suggesting that they have intrinsic growth even when their cells have severely reduced IIS.

Figure 6. The wings of pre- and post-critical size larvae differ in their response to starvation. Third-instar larvae were starved (open bars) or fed (filled bars) for 24 hours, either before or after attainment of critical size (approx. 8 hours after the moult from second to third instar), and the increase in their wing area $(\mu m^2)$ over that period was measured. Each bar represents 10–18 discs. See electronic supplementary material for methods.

Further evidence for intrinsic growth comes from work on the tobacco hornworm Manduca sexta. In this insect, disc growth responds only to changes in IIS in the presence of JH (Truman et al. 2006). In animals that have had their corpora allata excised (the gland that produces JH), starvation reduces but does not prevent disc growth. This remaining growth, which the authors term ‘morphogenetic growth’, is presumably an intrinsic growth that does not require nutritional input. The authors argue that morphogenetic growth sets the lower limit of disc size. In our model, intrinsic growth also sets the lower limit of disc size. Hypothetical larva in which IIS is eliminated immediately after attainment of critical size exhibit no body growth but their discs continue to grow until final disc size is appropriate for the much reduced body size.

An important aspect of intrinsic growth is that it is not present throughout development. In particular, our model suggests that, in order to express a linear static allometry, discs should not have intrinsic growth prior to attainment of critical size, when the time to the cessation of growth is not yet fixed. This appears to be the case in the buckeye butterfly Precis coenia, where starvation prevents growth of both the body and the wing discs in larvae that are still feeding (Miner et al. 2000). Further, the wing size at critical size is not influenced by changes in IIS in Drosophila (Shingleton et al. 2007), providing indirect evidence that the wing discs do not have intrinsic growth early in development. However, to test this directly, we starved pre- and post-critical size larvae and measured the extent of wing-disc growth over the subsequent 24 hours. Starvation prevents body growth and severely inhibits insulin signalling in a developing larva. The wing discs showed almost no additional growth in larvae starved before critical size, but grew considerably in larvae starved after critical size (figure 6). The fact that the wings grew slightly in larvae starved before critical size probably reflects a lag in the cessation of disc growth after a larva is removed from food rather than an intrinsic growth rate, although a direct manipulation of the insulin-signalling pathway in developing wing discs would confirm this. Nevertheless, our model and our data suggest that there is
reprogramming of the imaginal discs’ response to nutrition at or around the attainment of critical size.

The finding that JH is responsible for linking disc growth with nutrition in M. sexta makes it an excellent candidate for reprogramming the imaginal discs. In M. sexta, and possibly all holometabolous insects, the attainment of critical size is followed by a decline in JH levels (Nijhout & Williams 1974a). In Coleoptera and Lepidoptera, this decline is essential in ensuring that the succeeding moult is larval–pupal rather than larval–larval (Nijhout & Williams 1974a; Safranek et al. 1980; Gilbert et al. 1996). In Drosophila, there is also a decline in JH after critical size (Buhlen et al. 1984; Bownes & Rembold 1987). The application of JH throughout larval life does not prevent pupariation, but does prevent differentiation of the abdomen during the pupal–adult moult (Postlethwait 1974). Our model suggests a second role for the decline in JH in holometabolous insects, that of regulating the switch from wholly insulin-dependent growth to partially insulin-independent growth, essential in the regulation of allometry.

(c) Expanding the model
Although the model has been developed to explore the primary developmental factors that influence allometry in Drosophila, it can easily be applied to other holometabolous insects. Like Drosophila, all holometabolous insects have organs that develop as imaginal discs and are thought to have a critical size for metamorphosis. Unlike Drosophila, however, this critical size may be influenced by nutrition, for example in M. sexta. The effect of nutrition on critical size can be built into our model by making \( L \), critical size, a function of \( i \), the level of IIS. Similarly, while the duration of the body’s TGP in Drosophila appears to be only slightly affected by nutrition (Beadle et al. 1938; Mirth et al. 2005), this effect may be more substantial in other insects. This can be built into our model by making \( T_0 \), the body’s TGP, a function of \( i \). Finally, in many holometabolous insects, the imaginal discs do not begin growth until after the attainment of critical size. This is thought to be the ancestral state (Truman et al. 2006). The TGP for such discs is effectively their entire growth period. This can easily be accommodated in our model by replacing \( D_{12} \), disc size at critical size, with \( D_0 \), disc size at the beginning of growth, in equation (4.11).

6. CONCLUSIONS
The model suggests that two factors, insulin sensitivity and TGP, are fundamental regulators of the slope of allometries in holometabolous insects. Data concerning the insulin sensitivity and TGP of different imaginal discs in Drosophila, or any holometabolous insect, are lacking. Consequently, it is not yet possible to test whether the model can accurately predict the nutritional static allometries of different Drosophila organs. Nevertheless, the model makes a number of testable hypotheses concerning the mechanisms that regulate allometry in animals with asynchronous growth. First, even among organs that are isometric to each other and to body size, there should be differences in insulin sensitivity if these organs also differ in the period in which IIS influence their final size. In holometabolous insects, this is the TGP. The organs of many animals, including humans, begin and stop growth at different points in development, and so will have different periods during which their growth can be influenced by IIS. We might expect that these organs also differ in their insulin sensitivity. Second, the model suggests that there is reprogramming of imaginal discs after the attainment of critical size so that they take on an intrinsic growth rate. Our experimental data confirm this reprogramming in Drosophila, while data from M. sexta suggest that the reprogramming may be regulated by changes in the level of JH. Thus, the model provides a developmental reason for the finding that JH interacts with IIS to influence disc growth in M. sexta.

The model clarifies how altering specific developmental parameters may alter the expressed allometric relationship between organ size and body size. Changing key parameters, for example, insulin-sensitivity (\( s_i \)), duration of TGP (\( K \)) and critical size (\( L \)), all influence aspects of the an organ’s static allometry (figure 5). Variation in static allometry underlies variation in animal shape, and so the evolution of allometric relationships is a major factor in the generation of phenotypic diversity. Hitherto, there has been very little understanding of the developmental processes that regulate static allometries, essential if we are to identify candidate developmental mechanisms, pathways and genes upon which selection may act. Our model provides such candidates. The next stage is to explore whether the variation in these candidates correlates with that in expressed allometries in holometabolous insects, both intra- and interspecifically.

More than 100 years after the problem of allometry was first recognized and formulated by those studying the brain:body ratios in humans (Lapicque 1898), we are finally beginning to identify the developmental mechanisms that are involved in the regulation of nutritional static allometry. The relevance of these mechanisms to other forms of environmental static allometry created by other regulators of size, for example, temperature and oxygen levels, and to genetic static allometry and evolutionary allometry has yet to be determined. Nevertheless, this model and models similar to it (Nijhout et al. 2006) provide a firm developmental context for continued research.

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REFERENCES
Beadle, G., Tatum, E. & Clancy, C. 1938 Food level in relation to rate of development and eye pigmentation in


