Flies developed small bodies and small cells in warm and in thermally fluctuating environments

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Running title: Thermal plasticity of cell size
Summary

Although plasma membranes benefit cells by regulating the flux of materials to and from the environment, these membranes cost energy to maintain. Since smaller cells provide relatively more membrane area for transport, ectotherms that develop in warm environments should consist of small cells despite the energetic cost. Effects of constant temperatures on cell size qualitatively match this prediction, but effects of thermal fluctuations on cell size are unknown. Thermal fluctuations could favour either small or large cells: small cells facilitate transport during peaks in metabolic demand whereas large cells minimize the resources needed for homeoviscous adaptation. To explore this problem, we examined effects of thermal fluctuations during development on the size of epidermal cells in the wings of *Drosophila melanogaster*. Flies derived from a temperate population were raised at two mean temperatures (18°C and 25°C), with either no variation or a daily variation of ± 4°C. Flies developed faster at a mean temperature of 25°C. Thermal fluctuations sped development, but only at 18°C. An increase in the mean and variance of temperature caused flies to develop smaller cells and wings. Thermal fluctuations reduced the size of males at 18°C and the size of females at 25°C. The thorax, the wings, and the cells decreased with an increase in the mean and in the variance of temperature, but the response of cells was the strongest. Based on this pattern, we hypothesize that development of the greater area of membranes under thermal fluctuations provides a metabolic advantage that outweighs the greater energetic cost of remodelling membranes.
Introduction

When one considers the diversity of life, cell size rarely comes to mind as a variable trait that impacts the fitness of multicellular organisms. Indeed, foundations of some ecological theories were built on the assumption that all organisms have cells of the same size (West et al., 1997; discussed in Kozlowski and Konarzewski, 2004). Yet, cell size responds to artificial selection (Trotta et al., 2007) and varies among species (Kozlowski et al., 2010), populations (Goodman and Heah, 2010), and even life stages (Davison, 1955). We have good reasons to suspect that natural selection contributes to this variation in cell size. A given change in the volume of a cell causes a smaller change in the area of its surface. This disproportional scaling determines the costs and benefits of cell size (Szarski, 1983; Woods, 1999; Kozlowski et al., 2003; Atkinson et al., 2006). On one hand, a substantial portion of a cell’s energy goes to maintain ionic gradients across cell membranes (Rolfe and Brown, 1997), which enable cellular communication and trans-membrane transport. In fact, cell size correlates with resting metabolic rate in some animals (Davison, 1955; Starostova et al., 2009). All else being equal, the energetic cost of supporting cell membranes should favour large cells, which possess little surface area relative to cytoplasmic volume (frugal strategy in Szarski’s (1983) cell size model). On the other hand, since cell membranes regulate the exchange of chemicals between cells and their environment, the relatively large surface area of small cells should enhance metabolic performance. Therefore, when organisms must quickly process large quantities of resources, natural selection should favour genotypes that develop small cells despite the elevated energetic cost (wasteful strategy in Szarski’s (1983) cell size model).

Researchers have used this framework to predict the responses of cells to constant temperatures during development. Smaller cells are predicted to develop in warmer environments, where metabolic capacity and demands for resources are high, and oxygen supplies are low (Woods, 1999; Atkinson et al., 2006). Indeed, smaller cells have been observed at higher developmental temperatures in a wide range of ectotherms, including protists (Butler and Rogerson, 1996; Atkinson et al., 2003), rotifers (Stelzer, 2002), planarians (Romero and Baguna, 1991), nematodes (van Voorhies, 1996), fish (van Voorhies, 1996), lizards (Goodman and Heah, 2010), and flies (Partridge et al., 1994; Blanckenhorn and Llaurens, 2005). Moreover, *Drosophila melanogaster* evolved smaller cells at a higher temperature under controlled conditions (Partridge et al., 1994). These data support the idea that optimal cell size depends on the thermal environment, but tell us little about the responses of organisms to natural environments that fluctuate in temperature.
We use the theoretical framework described above to evaluate the plasticity of cell size in fluctuating thermal environments. The balance of opposing selective pressures should determine the optimal cell size in environments where temperatures fluctuate. As the temperature changes, cells rebuild their membranes to maintain the fluidity that enables normal functions, a process referred to as homeoviscous adaptation (Hazel, 1995). When the temperature drops, the distribution of fatty acids in phospholipids shifts from saturated to unsaturated. This process requires dietary fatty acids as substrates, molecular oxygen for processes governed by desaturases, and ATP to fuel enzymatic activity (Stanley-Samuelson, 1988; Shanklin and Cahoon, 1998; Martin-Creuzburg et al., 2012). Since other biochemical reactions compete for these resources, homeoviscous adaptation imposes some cost that likely depends on the area of cell membranes. Thus, an organism composed of large cells, which sum to less cell membrane area, would save resources at fluctuating temperatures. Alternatively, cells that spend some time at high temperatures require sufficient resources to meet extreme metabolic demands (Pörtner, 2002; Angilletta, 2009). Small cells would provide the relatively large surface areas and short diffusion distances needed to transport resources (Szarski, 1983; Woods, 1999; Kozlowski et al., 2003). Depending on the relative importance of these processes, selection could favour genotypes that either increase or decrease cell sizes when thermal environments fluctuate.

To explore the impact of thermal fluctuations during development on cell size, we raised isofemale lines of *Drosophila melanogaster* (Meigen) under constant and fluctuating temperatures and compared the sizes of epidermal cells in their wings. The genotypes used in this experiment were derived from a temperate population that experiences strong diel and seasonal fluctuations in temperature; such fluctuations induce changes in membrane composition of *D. melanogaster* (Overgaard et al., 2006; Cooper et al., 2012). The sizes of cells in wings have been used frequently as a proxy for the sizes of other cells throughout the body (reviewed by Arendt, 2007). In these cases, one assumes that the mean size of wing cells correlates with the mean sizes of other types of cells. In species of *Drosophila*, this assumption has gained empirical support (Stevenson et al., 1995). More importantly, the sizes of cells in different organs of *D. melanogaster* were observed to respond similarly to developmental conditions. Cells in both the wings and two other organs were smaller at higher temperatures (Azevedo et al., 2002), and a concerted reduction of cell sizes in wings and the abdomen occurred during hypoxia (Heinrich et al., 2011). Thus, sizes of epidermal cells and their responses to temperature should provide a proxy of these traits in other tissues of flies.
Methods

Design of the experiment

In June of 2008, we used banana-bait traps to collect females of *D. melanogaster* at three sites in Terre Haute, Indiana (USA). Thirteen isofemale lines were formed by placing inseminated females into freshly yeasted vials containing moist instant medium (Formula 424, Carolina Biological Supply, Burlington, NC, USA). These vials were maintained at 21°C with a 12:12 light cycle. In the following generation, emerging males were examined to verify that the species was *D. melanogaster*. To form replicates within isofemale lines, four pairs of virgin males and virgin females from each line were transferred to new vials containing the same medium. After mating, females were transferred to new vials and were allowed to lay eggs for 24 h. After this period, each female was transferred to another new vial for a period of 24 h, and this process was repeated until we had four vials of eggs from each female.

At each transfer, the vial of eggs from each isofemale line was placed into one of four thermal treatments: 1) a constant environment of 25°C, 2) a fluctuating environment with a mean of 25°C (21° and 29°C during scotophase and photophase, respectively), 3) a constant environment of 18°C, and 4) a fluctuating environment with a mean of 18°C (14° and 22°C during scotophase and photophase, respectively). Lines were allocated to these thermal treatments using a Latin square design (Bradley, 1958). The treatments were maintained by programmable incubators (Model 818, Precision Scientific, Chicago, USA). To ensure adequate level of humidity, we placed containers of water in each incubator. The accuracy of each thermal treatment was verified to the nearest 0.5°C using data loggers (iButton Thermochron, Dallas Semiconductors, Dallas, TX, USA); mean temperatures were nearly identical between the fluctuating treatments (18.2° and 25.1°C) and constant treatments (18.1° and 25.2°C). The light cycle of each thermal treatment was 12L:12D.

Measuring sizes of thorax, wings and cells

We measured development time and morphological traits of flies in each thermal treatment. Developmental time was estimated as the number of days between oviposition and the first sign of emergence. The length of the thorax and the dimensions of the left wing were used as proxies for body size and wing size, respectively. The mean size of epidermal cells in the wing was a convenient estimate of cell size.

For measurements of morphology, we sampled up to four males and four females from each vial at seven days after eclosion. After anaesthesia with CO₂ and chloroform, flies were
placed on their sides under a dissecting microscope (Model M28, Leica Microsystems, Buffalo Grove, IL, USA). An ocular micrometer was used to measure thorax length to the nearest 0.025 mm. Following Partridge et al. (1994) we measured the distance between the base of the most anterior humeral bristle to the posterior tip of the scutellum. The left wing of each fly was removed with surgical micro-scissors. Each wing was then flattened with a drop of xylene on a microscopic slide and mounted with Permount medium (FisherScientific, Fair Lawn, NJ, USA). The dorsal surface of each wing was digitized under a microscope (Model DC5-163, National Optical, San Antonio, TX, USA). The images were used to calculate the dimensions of the wing and the density of trichomes. Since each epidermal cell supports a single trichome, the density of trichomes reflects the mean size of epidermal cells.

Following Gilchrist et al. (2001), we used imaging software (ImageJ, National Institutes of Health, USA) to measure two dimensions of each wing (Fig. 1): 1) the width of the wing, from the intersection of vein V and the trailing edge to the leading edge along a trajectory perpendicular to vein III, and 2) the length of the distal segment of vein III. A Principal Component Analysis (PCA) was used to generate an index of wing size from these two dimensions. To calculate trichome density, we counted trichomes in a circle (0.01 mm²) between the distal segments of wing veins IV and V (Fig. 1). Following Dobzhansky (1929), the reciprocal of trichome density was considered an estimate of the mean area of epidermal cells in the wing.

**Statistical modelling**

We used General Linear Models to estimate the effects of mean temperature (18°C vs. 25°C), thermal variation (present vs. absent), and sex (male vs. female) on developmental time, thorax length, wing size, and cell size. Line and family (nested within line) were included as random factors when they improved the fit of the model. For the analysis of wing size (score for PC1), thorax length was included as a continuous factor. For the analysis of cell size, wing size was included as a continuous factor. Initially, we modelled all main effects and interactions. Then, we removed terms from the model, starting with the highest order term, until we arrived at the model with the lowest value of AIC (Burnham and Anderson, 2002). Following the procedure described by Zuur and colleagues (2009), we fitted models using the nlme library (Pinheiro et al., 2011) of the R Statistical Package (R Development Core Team, 2011).

**Results**
As in previous studies, both developmental time and body size were affected by temperature. An increase in the mean or variance of temperature accelerated development (Table 1). The effect of thermal fluctuations was more pronounced at low mean temperature (Fig. 2A). The mean and variance of temperature interacted to determine thorax sizes of males and females (Table 2; Fig. 2B). A higher mean temperature during development yielded flies with smaller thoraxes. In constant environments, the thermal dependence of thorax size was more pronounced for males than it was for females. However, in thermally fluctuating environments, thorax sizes of males and females decreased by a similar magnitude with increasing mean temperature. Thermal fluctuations caused a decrease in thorax size, but this response was much smaller in magnitude than was the response to an increase in mean temperature; moreover, this response was observed only among males that developed at a mean of 18°C and among females that developed at a mean of 25°C (Fig. 2B).

Wing size, independent of thorax size, also depended on sex and developmental temperatures (Table 3, Fig. 3). The first principal component of wing dimensions described 98% of the variation. The two dimensions contributed equally to this principal component, loading at a value of 0.99. Thus, greater scores for this principal component reflected larger wings. The score for this principal component generally increased with increasing thorax size, but females developed larger wings than males if both sexes were compared at the same thorax size. Disproportionately larger wings were also produced when mean temperature or thermal variance was low. As with thorax size, wing size decreased more with increasing mean temperature than it did with increasing thermal variance. The scaling of wing size with thorax size was steeper for flies that developed at 18°C than it was for flies that developed at 25°C. This interaction was particularly pronounced in males (Fig. 3).

The sizes of cells within wings were also affected by thermal conditions during development (Table 4, Fig. 4). Cell size scaled positively with wing size in both males and females, but the scaling was steeper for males. Females had larger cells than did males for a given wing size. Although large wings consisted of large cells, flies that developed at high or fluctuating temperatures had disproportionally smaller cells for their wing size. The effect of thermal fluctuations on cell size was particularly visible for females, but was still weaker in magnitude than the effect of mean temperature.

**Discussion**

Either an increase in the mean or variance of temperature caused flies to develop smaller epidermal cells in their wings, suggesting that flies which experience higher temperatures...
produce organs from smaller cells. In a previous experiment, flies consistently produced smaller cells in wings and in two other organs when developing at a higher constant temperature (Azevedo et al., 2002). A correlated thermal plasticity of cell sizes in different cell types occurred also in dung flies (Blanckenhorn and Llaurens, 2005) and planarians (Romero and Baguna, 1991). In our experiment, small flies produced small wings composed of small cells. Since higher temperatures caused smaller body sizes, the thermal plasticity of cell size was partly coupled with the thermal plasticity of body size. Thermal conditions also directly affected cell size. Either a higher mean or variance of temperature caused flies to develop small wings relative to thorax size and small cells relative to wing size.

Consequently, flies exposed to a higher maximal temperature developed the smallest epidermal cells relative to their body sizes. Thus, our results suggest that the thorax, the wings and the cells respond similarly to temperature, but the response of cells is the strongest of these responses. Although thermal effects on cell size have been observed previously in various ectotherms (Partridge et al., 1994; Butler and Rogerson, 1996; Blanckenhorn and Llaurens, 2005; Goodman and Heah, 2010), these effects were not dissected to separate the independent response to temperature from the response related to body and organ sizes.

The observation that flies produced smaller cells in thermally fluctuating environments accords with the hypothesis that the area of cell membranes, and thus cell size, should depend on metabolic demands. Given the nonlinear relationship between temperature and performance, ectotherms that experience fluctuating temperatures can perform equal to, better, or worse than conspecifics from constant environments with the same mean temperature (Ruel and Ayres, 1999; Angilletta, 2009; Bozinovic et al., 2011). Consistent with this idea, our flies developed faster in the thermally fluctuating environment when the mean temperature was 18°C but at similar rates when the mean temperature was 25°C. In a previous study (Bozinovic et al., 2011), thermal fluctuations sped population growth of *D. melanogaster* at a mean of 17 °C but slowed population growth at a mean of 24 °C. The greater performance of flies at fluctuating temperatures requires resource delivery to meet metabolic demand. Small cells might have enabled fast acquisition of resources during brief exposures to peak temperatures. Such conditions open “windows of opportunity” for cells to grow and divide, which could ultimately enhance the fitness of the organism.

Cell membranes serve as points of exchange between the cytoplasm and its surroundings. Thus, the larger area of exchange and the shorter distance of diffusion associated with smaller cells could speed transport and enhance performance (Szarski, 1983; Woods, 1999; Kozlowski et al., 2003). Still other factors could favour larger surface areas of
cell membranes at higher temperatures. For example, oxygen permeates more efficiently
through the hydrocarbon phase of a membrane than through the aqueous phase of cytoplasm;
thus, cell membranes form pathways, rather than barriers, along which oxygen can penetrate
tissue (Subczynski et al., 1989). Since smaller cells provide a greater density of cell
membranes, tissues should become perfused with more oxygen, speeding its delivery to
mitochondria. Consistent with this idea, flies that developed in a hypoxic environment
produced small cells in two types of tissues (Heinrich et al., 2011). Furthermore, experimental
evolution of D. melanogaster in hypoxic conditions led to small cells that consumed oxygen
faster than did large cells (Zhou et al., 2007). Importantly, the superior diffusion of oxygen
through the hydrocarbon phase of membranes is more pronounced at high temperatures,
suggesting that small cells especially benefit a warm organism (Subczynski et al., 1989).

Our findings should help to refine the theory of optimal cell size and metabolic scaling
and direct researchers toward new hypotheses. Since flies that experience temporal changes in
body temperature develop smaller cells, two scenarios remain possible: either membrane
remodelling during thermal change requires little energy, or the metabolic advantages of small
cells outweigh the energetic savings of large cells. We favour the latter hypothesis for two
reasons. First, flies rapidly remodel their membranes during thermal change (Hazel 1995;
Overgaard et al., 2006). Although no one knows the specific cost of membrane remodelling
during thermal change, phospholipid metabolism can require substantial amounts of ATP
(Purdon et al., 2002). Second, females in our experiment shrank their cells at fluctuating
temperatures more than did males, both in absolute terms and in relative terms (3.2 % vs. 3.1
% at 18 °C, and 3.5 % vs. 1.9 % at 25 °C). This pattern makes sense when one considers that
the balance between the cost of remodelling and the potential to acquire resources depends on
the state of a cell. Females produced larger cells with a relatively small area of membrane
compared to males, which should make them more prone to limitation imposed by a trans-
membrane transport during expositions to higher temperatures. Therefore, females should
gain a greater metabolic advantage from shrinking their cells than do males. At the same time,
females would not gain a significant energetic advantage from enlarging their cells, given
their already small area of membranes. At least one other study has generated patterns that
accord with this hypothesis; bryozoans developed smaller cells at a higher temperature in a
tissue that acquired oxygen through passive diffusion, but did not do so in a tissue that
received active ventilation (Atkinson et al., 2006).

To fully understand the impact of thermal fluctuations on cell size, we need to answer
several questions. First, do cells of all tissues grow or shrink in concert, as we have assumed
here? Although some evidence supports our assumption (Stevenson et al., 1995; Azevedo et al., 2002; Kozlowski et al., 2010), future research should address plastic responses and evolutionary changes of cell size in diverse tissues. Second, how much does the area of cell membranes impact the capacity of cells to acquire resources, especially oxygen? Third, how much energy does a cell spend to remodel a given area of membrane? The answers to these questions will enable biologists to develop quantitative models of optimal cell size.

Hypotheses related to the fitness consequences of membrane maintenance and efficient resource delivery can be directly tested to explicitly determine the selective advantage of different cellular strategies among environments. For example, genotypes that differ in the degree of membrane plasticity and in cell size can be competed to see whether thermal fluctuations favour large cells with a greater capacity for remodelling membranes. Genotypes that differ in rates of membrane transport and in cell size could be competed to see whether resource pulses favour small cells with greater rates of transport. These types of experiments will be needed to understand how the plasticity of cell size evolves to enhance the ability to acquire resources while reducing the costs of membrane maintenance in changing thermal environments.

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References


Figure captions

Figure 1. Two dimensions of wings were measured and they were described with a principal component analysis. The density of trichomes in the circle (0.01 mm²) was used to estimate the mean area of cells in each wing blade.

Figure 2. Generally, flies in warmer environments developed faster. When the mean temperature was low, fluctuations in temperature also sped development (A). Although females were generally larger than males, both sexes developed smaller bodies in warmer environments (B). Data are means and standard deviations estimated from the most likely statistical model.

Figure 3. Flies at a low, constant temperature developed the largest wings for a given body size, whereas flies in a warm, thermally fluctuating environment developed the smallest wings. Wing size was indexed as the score for the first principal component of two wing dimensions (Fig. 1). Lines display the relationships estimated from the most likely statistical model.

Figure 4. Flies at a low, constant temperature developed the largest cells for a given wing size, whereas flies in a warm, thermally fluctuating environment developed the smallest cells. Wing size was indexed as the score for the first principal component of two wing dimensions; cell area was calculated from the density of trichomes on each wing blade (Fig. 1). Lines display the relationships estimated from the most likely statistical model.
Figure 1
Figure 2

A) Mean developmental temperature (°C)

- Thorax size (mm)
  - Females: 0.75, 0.80, 0.85, 0.90, 0.95, 1.00, 1.05
  - Males: 0.75, 0.80, 0.85, 0.90, 0.95, 1.00, 1.05

Developmental time (d)
- Constant temperature
- Fluctuating temperature

B) Thorax size (mm)
- Females: 1.00, 0.95, 0.90, 0.85, 0.80
- Males: 1.00, 0.95, 0.90, 0.85, 0.80

Mean developmental temperature (°C)
Figure 3

The graph shows the relationship between thorax length (mm) and index of wing size for males and females at two different temperatures: 18 °C and 25 °C. The data points are color-coded to represent constant and fluctuating conditions. The graphs indicate a positive correlation between thorax length and index of wing size.
Figure 4

The figure shows scatter plots for males and females, comparing index of wing size with cell area (mm$^2$ x 10,000$^{-1}$). Two temperatures are considered: 18 °C and 25 °C. The plots indicate a positive correlation between the index of wing size and cell area, with different symbols and line styles representing constant and fluctuating conditions.
Table 1. Inferential statistics for the most likely General Linear Model of developmental time in *Drosophila melanogaster*, as determined by the Akaike information criterion.

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Table 2. Inferential statistics for the most likely General Linear Mixed Model of thorax length in *Drosophila melanogaster*, as determined by Akaike information criterion.

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Table 3. Inferential statistics for the most likely General Linear Mixed Model of wing size in *Drosophila melanogaster* developed at two average temperatures, as determined by Akaike information criterion. Wing size was the first principal component derived from analysis of two wing dimensions (Fig. 1).

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Table 4. Inferential statistics for the most likely General Linear Mixed Model of cell size in *Drosophila melanogaster*, as determined by Akaike information criterion. Wing size was the first principal component derived from analysis of two wing dimensions; cell size was the mean area of epidermal cells in the wing blade (Fig. 1).

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</tbody>
</table>