Genetic variation of heat tolerance in a model ectotherm: an approach using thermal death time curves

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Abstract
The assessment of thermal tolerance holds significant importance in predicting the physiological responses of ectotherms, particularly in elucidating their capacity for evolutionary adaptation in the context of global warming. Current approaches to assessing thermal tolerance have limitations that can lead to misleading results, especially with regard to the heritability of thermal limits. In this study, we examined twenty isogenic lines of Drosophila melanogaster from the DGRP panel to characterize their thermal death time (TDT) curves, which account for the duration and intensity of heat stress. Furthermore, we examined the extent of genetic variation in the two parameters that characterize TDT curves, namely CTmax and thermal sensitivity z. Our analysis revealed evidence of genetic (co)variation for both parameters. Results from simulations of the evolutionary consequences of selection on CTmax and z suggest that directional selection to increase CTmax will also increase z as a correlated response. However, directional selection to increase z may have the opposite effect. We conclude that the evolution of thermosensitive or thermotolerant strategies is better achieved by directional selection to decrease or increase CTmax, which may aid in mitigating the effects of global warming on ectotherms.

Keywords: global warming, heritability, isogenic lines, thermal death time curves
1. Introduction

There is substantial evidence pointing to an unprecedented rise in the temperature of our planet. According to climate models, if the present warming trends persist, the surface temperature of the Earth's surface will surpass the average at the end of the 19th century by 1.5°C [1]. It is not surprising that this rise will have repercussions on the biota present on our planet, particularly for animals such as ectotherms, whose physiological processes are closely linked to ambient temperature [2,3]. Climate change is already having an impact at the demographic level. Many species are shifting their ranges, often towards cooler regions [4], while others are threatened with extinction [5]. There is ample evidence that some species might be able to adapt to rising temperatures through the evolution of heritable stress-tolerant phenotypes [6]. Some lineages are already able to withstand high temperatures [7], while others show phenotypic plasticity [8], and some species may evolve in response to warming [6,9]. Assessing the adaptive capacity and heritability of heat tolerance is challenging, as inaccuracies in models can lead to either under- or overestimation of vulnerability to climate change [10,11].

The current debate about the evolutionary potential of heat tolerance in ectotherms can be summarized in two main points. First, there is the notion that ectotherms possess limited plasticity for heat tolerance, suggesting that heat tolerance is both evolutionary and physiologically fixed. Natural selection appears to affect physiological responses to lower temperatures more than to higher temperatures [12]. Interspecific studies - i.e., among species - have failed to detect genetic variability in heat tolerance, variation between species and populations, and a lack of latitudinal diversity [13,14]. Likewise, selection and heritability experiments on single species suggest limited increases in upper thermal limits. Second, a complicating factor in understanding the genetic basis of upper thermal limits is that these are to some extent affected by methodological issues [for a review, see 12]. Heat tolerance is often estimated using ramping assays to assess upper critical thermal limits (CTmax), which are the maximum ambient temperatures that an ectotherm can tolerate under a given experimental condition before succumbing to heat [15]. However, this approach is difficult to replicate at the individual level, especially when mortality is the measured endpoint [11]. The conclusion of these studies is clear and suggests a limited evolutionary potential for ectotherms to increase their ability to tolerate high temperatures. Despite the possibility of methodological issues underestimating the actual evolutionary potential to withstand increasing temperatures [10], it is imperative to acknowledge that CTmax is merely a component of the more intricate trait “thermal tolerance” [16].

An increasing number of studies have employed thermal death time (TDT) curves to measure heat tolerance [17–22]. However, the extent to which TDT curves reflect evolutionary changes within a species remains unclear. TDT curves imply that an individual’s survival probability is influenced by both temperature and exposure time [23], and may provide a more nuanced view of the two reasons mentioned above for the ongoing debate. In this context, we take advantage of recent research using the Drosophila Genetic Reference Panel (DGRP) [16,24] to investigate the genetic variation of heat tolerance in Drosophila melanogaster using TDT curves. The wild-type DGRP lines of this panel are derived from a single natural population and have been inbred to homozygosity, providing extensive information on genetic variation at multiple levels [25] and offering unique opportunities to quantify the genetic basis of physiological traits such as heat tolerance.
2. Material and methods

(a) Experimental flies and rearing conditions

Twenty inbred, isogenic wild-type *Drosophila melanogaster* (Meigen 1830) lines from the *Drosophila* Genetic Reference Panel (DGRP) were used as the study model. These lines were previously employed in a study with a different objective [19], and we assume that this subset represents an approximately random collection with respect to the focus of interest described here.

All twenty selected lines were obtained from the Bloomington *Drosophila* Stock Center in March 2018. They were maintained in quarantine on standard cornmeal-agar-yeast media at room temperature (approximately 22°C) for four generations until May 2018. The rearing conditions were identical to those detailed in Leiva et al. [19].

(b) Thermal death time (TDT) curves

We measured the heat tolerance of individual female and male flies using a similar experimental protocol as outlined in Verspagen et al. [22]. This involved using a heating circulating bath and a wireless thermometer to measure temperature consistently throughout each trial. Individual virgin flies were placed in sealed 4-mL glass vials, arranged on a Plexiglas™ rack, and submerged in a 9.5-L glass aquarium filled with water set to a constant temperature of 36, 37, 38, or 39°C. During each trial, a Nikon D5300 with the time-lapse feature captured images at 10-second intervals. Subsequently, the compiled images were transformed into reversed videos using the open-source software Blender. Before initiating the thermal tolerance experiments, the flies were allowed approximately 30 minutes in the vials at room temperature for recovery following light CO$_2$ anaesthesia. This recovery period proved effective, as the flies exhibited active flying or walking inside the vial.

A total of 1,686 flies underwent survival time measurements, and the parameters of the thermal death time (TDT) curves (CTmax and $z$) were calculated for each DGRP line and sex. The calculation utilized the equation outlined in Rezende et al. [23]:

$$\log_{10} t = \frac{C_{\text{max}} - T}{z}$$

(1)

where $t$ represents the survival time in minutes, CTmax is the temperature ($°C$) where the survival time is $\log_{10} t = 0$ after 1 min of exposure to assay temperature, $T$ is the assay temperature ($°C$), and the thermal sensitivity $z$ is the temperature change ($°C$) required for a ten-fold difference in survival time. We maintained control over assay temperature ($T$) and measured time ($t$) as the dependent variable. The estimation of CTmax and $z$ for each DGRP line and sex involved regression analysis of $\log_{10}$-transformed survival times against the four temperature treatments.

(c) Estimation of variance components and broad-sense heritability of CTmax and $z$

For each DGRP genotype, we first assessed the effect of stressful temperatures (covariate) on survival time (in $\log_{10}$ min) by using for each sex the following general mixed ANCOVA model that allows separating slopes and intercepts:
Survival time = $\beta_0 + \beta_1 \cdot \text{stress } T^a + \mu_0 Z + \mu_1 Z \cdot \text{stress } T^a + \epsilon$ (2)

where

$\beta_0$ and $\beta_1$: intercept and slope of the fixed effect of stress temperature $\text{stress } T^a$,

$\mu_0 \sim N(0, G_{11})$: vector of random coefficients representing the effect of each genotype on $\beta_0$,

$\mu_1 \sim N(0, G_{22})$: vector of random coefficients representing the effect of each genotype on $\beta_1$,

$Z$: Design matrix (array of dummy variables) representing the genotypes,

$G$: variance-covariance matrix for the random effects,

$\epsilon$: vector of random errors

We fitted linear mixed-effects models [26] and obtained various estimates of the variance components $\hat{\sigma}^2_{\beta_0i}$ ($i = 1, \ldots, 20$), $\hat{\sigma}^2_{\beta_1i}$ and $\hat{\sigma}^2_\epsilon$ (caret denotes "estimate") that refer, respectively, to the variation in the intercepts and slopes of the TDT curves for the DGRP genotypes, and the residual variation. Model coefficients and variance components were then used to estimate $CT_{max} = \beta_1/\beta_0$ and $z = -1/\beta_1$, whose Taylor expanded variances [27, p. 240], became,

$$\text{Var}(CT_{maxi}) = \text{Var}\left(\frac{-\hat{\beta}_{0i}}{\hat{\beta}_{1i}}\right) \approx \left(\frac{\mu_{-\hat{\beta}_{0i}}}{\mu_{\hat{\beta}_{1i}}}\right)^2 \left[\hat{\sigma}^2_{-\beta_0i} + \hat{\sigma}^2_{\beta_1i} - 2 \frac{\text{Cov}(\hat{\beta}_{0i}, \hat{\beta}_{1i})}{\mu_{-\hat{\beta}_{0i}} \mu_{\hat{\beta}_{1i}}}\right]$$

$$\text{Var}(z_i) \approx \frac{\hat{\sigma}^2_{\beta_1i}}{(\mu_{z_i})^4}.$$  \hspace{1cm} (3)

We then estimated broad-sense heritability as:

$$H^2_{CT_{max}} = \frac{\text{Var}(CT_{maxi})}{\text{Var}(CT_{maxi}) + \text{Var}(z_i) + \frac{\hat{\sigma}^2_\epsilon}{n_0}}$$

$$H^2_z = \frac{\text{Var}(z_i)}{\text{Var}(CT_{maxi}) + \text{Var}(z_i) + \frac{\hat{\sigma}^2_\epsilon}{n_0}}$$

where
\[ n_0 = \frac{1}{a - 1} \left( \sum_{i=1}^{a} n_i - \frac{\sum_{i=1}^{a} n_i^2}{\sum_{i=1}^{a} n_i} \right) \]  

(5)

is the appropriate mean value of the number of flies from each sex and DGRP line used at each stressful temperature to estimate the TDT curves [28, p. 212] The reason we divided the residual variance by \( n_0 \) is because we are using line means [29]. In our case \( n_0 = 10.2480 \) for females and \( n_0 = 10.8055 \) for males.

Delete-one-DRGP genotype at a time data resampling was also carried out to estimate the genetic components of variance and their standard errors [30]. A total of 20 pseudovalues for each sex were obtained by dropping, in turn, each DGRP line and calculating:

\[ \phi_i = N\hat{\theta}_N - (N - 1)\hat{\theta}_{N-1,i}, \]

(6)

where \( \phi_i \) is the \( i \)th pseudovalue, \( \hat{\theta}_N \) is the corresponding variance estimate using all \( N = 20 \) DGRP genotypes, and \( \hat{\theta}_{N-1,i} \) is that estimate by dropping the \( i \)th DGRP genotype alone. The jackknife estimate is the average of \( \phi_i \), and its standard error is given by

\[ SE = \frac{\sum_{i=1}^{N} (\phi_i - \hat{\phi})^2}{N(N - 1)}. \]

(7)

Approximate 95% jackknife confidence intervals were obtained as \( \hat{\phi} \pm 2 \text{ SE} \). Initially, the analyses for computing variance components and heritability were implemented by MS in MATLAB. To enhance reproducibility, FPL and EJN replicated the analyses and implemented them in R version 4.3.2 [31]. The data used in these analyses were based on a recently reported study [19,32].

(d) Hypothetical selection on the TDT curves

Appropriate estimates of the additive-genetic G and phenotypic P (co)variance matrices in an outbred Drosophila population are needed to explore the hypothetical evolutionary consequences of selection on CTmax and \( z \) from the multivariate breeder’s equation \( \Delta \mu = G\beta = GP^{-1}s \). Here, the term \( \Delta \mu \) is the vector of changes in trait means, \( \beta \) is the vector of selection gradients, and \( s \) is the vector of selection differentials [33,34].

The estimates of genetic variance-covariance components and broad-sense heritability in the highly inbred DGRP lines yield only the relative contributions of CTmax and \( z \) to the total genetic variance in the TDT curves [see 29]. To our knowledge, there are no estimates of the narrow-sense heritability of TDT curves. Current evidence suggests that CTmax (estimated by different methodologies) is moderately heritable, and it seems reasonable to assume that its narrow-sense heritability is \( h_{\text{CTmax}}^2 \approx 0.25 \) [35–37]. Based on
this information and the relative contribution of CTmax and \( z \) to the total genetic variance of TDT curves, we approached the hypothetical consequences of several selective scenarios on the evolution of thermal tolerance (represented by these two parameters in the TDT curves) in an outbred population. Note that this will be “what-if” scenarios and more accurate estimates of G and P would be needed for a more satisfactory answer.

3. Results

(a) Determination of TDT curves and variance components

We observed substantial variation in thermal death time (TDT) curves across genetic lines for both females and males (Figure 1). At 36°C, the average survival times (± SD) were 150 ± 47 minutes for females and 111 ± 41.6 minutes for males. These durations decreased significantly at 39°C, with females surviving for 12.7 ± 5.13 minutes and males for 14.7 ± 5.6 minutes on average. Notably, this variation in thermal tolerance across genetic lines was consistently observed for both sexes (Figure 1).

Table 1 provides estimates of the variance-covariance components and broad-sense heritability using different methods for estimating parameters in the linear mixed-effects model. These estimates were highly consistent across the various methods of estimation. As indicated by the jackknife 95% confidence intervals, all variance components were significantly different from zero. Furthermore, the genetic covariance between \( \beta_0 \) and \( \beta_1 \) was negative. Broad-sense heritability was around 0.75 for CTmax and around 0.25 for \( z \). In other words, CTmax accounts for approximately 75% and \( z \) accounts for approximately 25% of the total genetic variance in the TDT curves.

![Figure 1. Thermal death time (y-axis in log\(_{10}\)-scale) curves for females (left) and males (top). Dots represent the individual survival time for females (green, left plot) and males (brown, right plot).](image-url)
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Table 1. Estimates of variance-covariance components and broad-sense heritability using various methods for estimating parameters in the linear mixed-effects model.
(b) Hypothetical selection on the TDT curves

We can employ the raw phenotypic (co)variance matrix of the females from the DGRP lines as a representative of the $P$ matrix in an outbred population, omitting any changes in phenotypic variance caused by inbreeding for the sake of simplicity [38]. Assuming absence of epistasis, the genetic variation in the DGRP lines for CTmax and $z$ (REML estimates for females in Table 1) has been rescaled so that $h^2_{\text{CTmax}} \approx 0.25$ and $h^2_{z} \approx 0.07$. Hence, we presume the subsequent (co)variance genetic, environmental, and phenotypic matrices for CTmax and $z$ in the hypothetical outbred base population:

$$G = \begin{bmatrix} 0.1489 & 0.0423 \\ 0.0423 & 0.0133 \end{bmatrix}; E = \begin{bmatrix} 0.4466 & 0.2618 \\ 0.2618 & 0.1691 \end{bmatrix}; P = \begin{bmatrix} 0.5955 & 0.3041 \\ 0.3041 & 0.1823 \end{bmatrix}$$

The vector of phenotypic means in this base population is $\bar{\mu} = [41.9924 \ 2.7592]^T$ ($T$ stands for transpose), where the first value is for CTmax and the second for $z$ (female means from the DGRP lines). We assume that these values are representative of the outbred population, which is strictly true if allelic effects are additive [39].

We simulated three scenarios for increasing thermoresistance (Figure 2): directional selection for CTmax ($s = [1 \ 0]^T$), directional selection for both CTmax and $z$ ($s = [1 \ 0.5]^T$), and directional selection for $z$ ($s = [0 \ 0.5]^T$). These selection differentials correspond to intensities of selection $i_{\text{CTmax}} = 1.3$ and $i_{z} = 1.2$. The hypothetical selection regimes used to illustrate the effects of selection are much stronger than we would expect in nature because directional selection tends to be weak and rarely shifts mean by more than half of a phenotypic standard deviation [40,41]. However, under extreme climatic events, such as heat waves, which are considered to be major triggers of evolution [42], these intensities of selection may not be unrealistic [43].
Figure 2. The thermal death time (TDT) curve linearly describes the relationship between test temperature ($T$) and survival time ($t$, log$_{10}$ scale) under heat stress conditions in *D. melanogaster*. Thermal sensitivity ($z$) is the reciprocal of the slope ($\beta_1$), representing the increase in temperature required to reduce survival time by one order of magnitude (10-fold). CTmax is the intersection at log$_{10} = 0$, corresponding to the knockdown or death temperature after 1 minute of exposure. The blue line represents a thermosensitive genotype, which exhibits improved tolerance to acute, intense heat stress but reduced tolerance to chronic, less intense heat. Conversely, the red line depicts a thermoresistant genotype, exhibiting better tolerance to chronic stress but lower tolerance to acute stress.

The simulated scenarios are shown in Figure 3. It is striking that a directional selection gradient always has a different sign than its selection differential, and the evolutionary response can be against the selection differential (Figure 3C). The inference from these hypothetical selection regimes is that directional selection to increase thermal sensitivity $z$ seems to hinder evolutionary responses to increasing CTmax. In other words, directional selection to increase CTmax also increases $z$ as a correlated response (Figure 3A) and, as expected, drives the population towards a more thermoresistant state (Figure 2). However, directional selection to increase $z$ results in a decrease of CTmax as a correlated response, which, in turn, also decreases $z$, and seemingly paradoxically drives the population towards increasing thermosensitivity (Figure 2).
Figure 3. Hypothetical strong directional selection for increased thermoresistance. In blue the 95% confidence ellipses of a simulated population of $N = 10,000$ flies from a bivariate normal distribution whose phenotypes for CTmax and $z$ are the sum of genetic effects with mean $\mu = [41.9924 \ 2.7592]^T$ and additive-genetic (co)variance $G$, plus environmental effects with mean 0 and (co)variance $E$ (see text). The arrows centred on the bivariate means represent the directions in which the data vary the most (i.e., the eigenvectors of the covariance matrix of the data). In black are the 95% confidence ellipses after an evolutionary shift in the means (we ignore changes in variance and covariance). Panel A plots directional selection for increasing CTmax; the vector of selection gradients is $\beta = [11.3294 \ -18.8959]^T$. Panel B plots directional selection for increasing both CTmax and $z$, where $\beta = [1.8814 \ -0.3958]^T$. Panel C plots directional selection for increasing $z$, where $\beta = [-9.4480 \ 18.5002]^T$. 

\[ \Delta \mu = \begin{bmatrix} 0.8864 \\ 0.2289 \end{bmatrix} \]  
\[ \Delta \mu = \begin{bmatrix} 0.2633 \\ 0.0744 \end{bmatrix} \]  
\[ \Delta \mu = \begin{bmatrix} -0.6231 \\ -0.1545 \end{bmatrix} \]
4. Discussion

Heat tolerance has traditionally been examined from a physiological perspective, with a focus on the mechanisms that cause animals to succumb to heat stress, such as oxygen limitation or excessive water loss [see 44 for a review]. In this study, we present a novel viewpoint by examining the genetic components of heat tolerance through the lens of thermal death time (TDT) curves. This allows partitioning the relative contribution of CTmax and z to the more complex trait “thermotolerance” (Figure 2). Our approach has relied on inbred Drosophila melanogaster lines from the DGRP panel, and it would be highly desirable to extend these analyses to outbred populations.

The limited number of previous studies that have assessed the thermal tolerance of DGRP lines focused primarily on measuring critical thermal limits [45,46]. These, along with other studies [12,47], suggest that heat tolerance is somewhat evolutionary constrained, an idea often invoked to explain the absence of strong latitudinal clines in heat tolerance [13]. Typically, assessments of organisms’ vulnerability to global warming usually compare experimentally derived thermal limits using ramping trials, in which animals are exposed to increasingly higher temperatures [15,48]. During these trials, the intensity and duration of stress increase concurrently. As a result, animals often succumb to heat stress in rapid succession, leading to small variances and small standard deviations in the measurements, making CTmax an attractive endpoint to use in treatment comparisons. However, CTmax is a single point, whereas the trait of interest, namely the ability of an organism to deal with heat stress, is a linear function describing how stress intensity and stress duration impact survival [23]. Thus, ramping trials approach overlooks the cumulative nature of heat injury and the time-dependent effects of thermal tolerance [49–51], potentially underestimating organisms’ vulnerability to global warming [52]. We contend that utilizing TDT curves to evaluate both CTmax and z parameters, as well as their underlying genetic basis, would provide more accurate predictions. Here, we have developed a methodological approach to estimate the variance components and heritability of the relevant parameters in the TDT curves.

The hypothetical selection scenarios allow us to understand how thermosensitive and thermotolerant strategies (Figure 2) can evolve. These scenarios suggest that thermosensitive or thermotolerant strategies are better achieved by directional selection to decrease or increase CTmax. This conclusion holds in more realistic scenarios, where the intensity of selection on CTmax and/or z might be relatively weak. We acknowledge that our approach is only a rough estimate of the problem and that several caveats could be raised. For instance, although there is a high positive correlation across species for parameters CTmax and z (r = 0.92; [23]), the G matrix used to simulate the hypothetical scenarios could overestimate the additive genetic covariance in an outbred Drosophila population due to higher linkage disequilibria in the DGRP lines [29]. However, for the time being, we believe that the present conclusions may be broadly applicable.

In summary, our findings suggest that the genetic correlation between CTmax and z impose constraints on thermal tolerance strategies. The simulations performed here highlight the importance of considering the multivariate nature of thermal tolerance traits and their genetic correlations when predicting evolutionary responses to climate change. Ultimately, this can be achieved by utilizing approaches that measure thermal tolerances considering both the duration and intensity of heat stress and its potential for evolution. Adopting such
integrative approaches will enable more accurate predictions of how species might respond to increasing temperatures in a rapidly changing planet.

**Data accessibility**

Data files and code supporting analyses, figures and tables of this study are publicly available on GitHub ([https://github.com/felixpleiva/Genetic_variation_TDT](https://github.com/felixpleiva/Genetic_variation_TDT)). When using the code from this manuscript, please cite it as: Leiva FP, Santos M, Niklitschek EJ, Rezende EL, & Verberk WCEP. (2024). Paper data and code for: Genetic variation of heat tolerance in a model ectotherm: an approach using thermal death time curves. Zenodo. DOI will be available here.

**Authors' contributions**

Félix P. Leiva: conceptualization, data curation, formal analysis, funding acquisition, investigation, methodology, project administration, resources, software, validation, visualization, writing – original draft preparation, writing – review and editing; Mauro Santos: conceptualization, formal analysis, methodology, software, writing – original draft preparation, writing – review and editing; Edwin J. Niklitschek: formal analysis, software, validation, writing – review and editing; Enrico L. Rezende: conceptualization, investigation, writing – review and editing; Wilco C.E.P. Verberk: conceptualization, formal analysis, funding acquisition, investigation, methodology, resources, supervision, writing – review and editing. All authors gave final approval for publication.

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