

# The quantitative genetic basis of clinal divergence in phenotypic plasticity

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## ABSTRACT

Phenotypic plasticity is thought to be an important mechanism for adapting to environmental heterogeneity. Nonetheless, the genetic basis of plasticity is still not well understood. In *Drosophila melanogaster* and *D. simulans*, body size and thermal stress resistance show clinal patterns along the east coast of Australia, and exhibit plastic responses to different developmental temperatures. The genetic basis of thermal plasticity, and whether the genetic effects underlying clinal variation in traits and their plasticity are similar, remains unknown. Here we use line-cross analyses between a tropical and temperate population of *D. melanogaster* and *D. simulans* developed at three constant temperatures (18, 25 and 29 °C) to investigate the quantitative genetic basis of clinal divergence in mean thermal response (elevation) and plasticity (slope and curvature) for thermal stress and body size traits. Generally, the genetic effects underlying divergence in mean response and plasticity differed, suggesting that different genetic models may be required to understand the evolution of trait means and plasticity. Furthermore, our results suggest that non-additive genetic effects, in particular epistasis, may commonly underlie plastic responses, indicating that current models that ignore epistasis may be insufficient to understand and predict evolutionary responses to environmental change.

## INTRODUCTION

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Phenotypic plasticity, the ability of a genotype to alter its phenotype under different environmental conditions (Bradshaw 1965), is thought to be an important mechanism for responding to heterogeneous environments (Janzen 1967; Sultan and Spencer 2002; Ghalambor et al. 2004), and may be significant for population persistence in the face of novel environmental change (Chevin et al. 2013). Additionally, while phenotypic plasticity allows an organism to counter environmental changes, canalization, where a phenotype remains constant under different environments, may also be an important mechanism that buffers phenotypes against environmental perturbations (Waddington 1942). As such, phenotypic plasticity and canalization describe different facets of the same phenomenon: the sensitivity of phenotype to the environment (Debat and David 2001). Plastic responses are prevalent in nature, and the large number of studies reporting genotype-by-environmental ( $G \times E$ ) interactions within and/ or between populations suggest that genetic variation for plasticity is widespread (Scheiner 1993; Kruuk et al. 2008; Des Marais et al. 2013). Nonetheless, the genetic basis, and the quantitative genetic architecture of plasticity/ canalization is still not well understood, particularly in natural populations (Scheiner 1993; Via 1993; Via et al. 1995; Flatt 2005).

Two main models have been proposed to describe the genetic mechanisms underlying plastic responses (reviewed in Scheiner 1993; Via et al. 1995). The first model, the allelic sensitivity, or pleiotropic model, suggests that phenotypic plasticity is a function of differential expression of the same genes under different environments (Falconer 1952; Via and Lande 1985; Via 1993). Thus, in different environments similar loci will respond, but

individual alleles may vary in their sensitivity/ expression. The second model, the gene regulation or epistasis model, suggests that plasticity is due to genes that determine the magnitude of responses to environmental effects which interact with genes that determine the average expression of the character (Scheiner and Lyman 1989; Scheiner 1993; Schlichting and Pigliucci 1993). Although there has been some controversy over whether plasticity is underpinned by specific “plasticity genes” or through environmentally sensitive alleles (Via 1993; Scheiner 1993; Schlichting and Pigliucci 1993), mapping, expression and family/ selection studies have found empirical evidence for both models (e.g. allelic sensitivity: Barnes et al. 1989; Weber and Scheiner 1992; Wu 1998; Ungerer et al. 2003; Lacaze et al. 2009; e.g. gene regulation: Li et al. 2010; Zhou et al. 2012; Mendez-Vigo et al. 2016). Furthermore, de Jong (1995) suggested that both models are mathematically equivalent, and may be modelled using a reaction norm approach (i.e. the expression of a character as a function of an environmental variable). Using this approach, de Jong and Gavrillets (2000) found that changes in the genetic variance and covariance of reaction norm parameters under different levels of environmental variance depend on the number of pleiotropic loci, suggesting that the gene regulation and allelic sensitivity models will have different consequences for the evolution of the genetic variance in reaction norm parameters (de Jong and Gavrillets 2000) and thus on the rate of evolution of plasticity.

The relative importance of additive and non-additive genetic variation in adaptive evolution has been the subject of much controversy, with some researchers suggesting that the evolution of quantitative traits will occur predominately via additive gene action (Fisher 1930; Coyne et al. 1997, 2000; Hill et al. 2008), while others propose that non-additive effects may be important (Wright 1931; Fenster et al. 1997; Wade and Goodnight 1998;

Paixao and Barton 2016). The extent to which adaptive evolution occurs predominately via additive or non-additive effects has direct implications for speciation models and the evolution of reproductive isolation (Dobzhansky 1937; Mayr 1954; Carson and Templeton 1984; Orr 1995), the evolution of sex and recombination (Maynard Smith 1978; Barton 1995), the maintenance of genetic variation (Gimelfarb 1989; Hermisson et al. 2003), animal and plant breeding (Lee and Kim 2009; Fethi et al. 2011) and conservation genetics (Fenster et al. 1997). Significantly, epistasis plays an important role in evolutionary models of canalization and genetic robustness (Wagner et al. 1997; Rice 1998; Flatt 2005). Furthermore, a study on *Escherichia coli* found that mutations that show epistasis were disproportionately likely to also show phenotypic plasticity (Remold and Lenski 2004), suggesting that epistasis may generally be fundamental to the evolution of environmental sensitivity (plasticity and canalization). Nonetheless, we currently know little about the quantitative genetic architecture of environmental sensitivity/ plasticity.

Although the gene regulation/ epistasis model of plasticity emphasises physiological epistatic gene interactions between regulatory and trait loci, it is unknown whether these interactions generate epistatic variance and influence the evolution of plasticity (Via et al. 1995). Several QTL studies have found evidence for dominance and/ or epistatic genetic effects for environmental sensitivity/ plasticity (Wu 1998; Li et al. 2014; Mendez-Vigo et al. 2016) suggesting that non-additive genetic effects may underpin plastic responses. However, the extent to which these loci generate epistatic variance (rather than additive variance) that influences the evolution of plasticity, especially in natural populations, is not clear. Although line cross analyses (Mather and Jinks 1982) and crosses examining outbreeding depression (or F<sub>2</sub> breakdown) (Fenster et al. 1997) do not measure standing levels of additive, dominant

and epistatic genetic variance, and thus cannot predict the contribution of non-additive genetic effects to any future short term responses to selection, they can detect whether epistatic interactions were important in the evolution of divergence in means across populations/ lines (Fenster et al. 1997). A small number of controlled crossing experiments have found evidence that dominance and/ or epistatic genetic effects underlie divergence in environmental sensitivity/ plasticity (Westerman 1971b, a; Perkins and Jinks 1973; Connolly and Jinks 1975; Pooni et al. 1987). However, most studies were conducted on inbred or artificially selected lines, rather than natural populations/ populations recently collected from the field (but see Westerman 1971a). Furthermore, past studies have also failed to clearly distinguish contributions from dominance compared to different types of digenic epistatic effects; nor were they able to account for maternal effects, which if present, may lead to the erroneous detection of epistasis (Gilchrist and Partridge 1999; Kennington et al. 2001). Importantly, to our knowledge, no studies have used a comprehensive crossing design to investigate the genetic architecture underlying divergence in plasticity in populations collected from different biogeographical locations. This information would provide valuable insight into the extent to which non-additive genetic effects are important for adaptive evolution of plasticity in nature, and whether current plasticity models that assume that additive genetic effects underlie evolution are sufficient.

For *Drosophila*, temperature is a major environmental factor influencing the geographic distribution of species, and adult tolerance to thermal extremes (heat and cold resistance) provides a good predictor of current species distributions (Overgaard et al. 2014). In *Drosophila melanogaster*, opposing genetic clines in heat and cold resistance have been observed along the east coast of Australia; tropical populations have higher heat resistance

than temperate populations, while temperate populations recover faster from a cold stress than tropical populations (Hoffmann et al. 2002; Sgro et al. 2010; Cockerell et al. 2014). Similar to many other species (Chown and Gatsos 2010; Meiri and Dayan 2003), body size also varies with latitude, with size increasing at higher latitudes (James et al. 1995; van Heerwaarden and Sgro 2011).

In *Drosophila* body size, heat and cold resistance also respond plastically to temperature. Warmer rearing temperatures result in smaller flies (Atkinson 1994; James et al. 1997; Azevedo et al. 1998; van Heerwaarden and Sgro 2011) and increased heat resistance, while cooler rearing temperatures commonly increase cold resistance and size (Hoffmann et al. 2005; Cockerell et al. 2014). Although clinal patterns for the mean value of these traits in two species, *D. melanogaster* and *D. simulans*, do not change significantly across different developmental temperatures, there is evidence for genetically based differences in plasticity for body size and heat and cold resistance along the eastern Australia cline in both species (van Heerwaarden and Sgro 2011; Cockerell et al. 2014). Furthermore, as phenotypically plastic responses to temperature mirror clinal patterns (i.e. plastic changes in these traits in response to rearing temperature change in the same direction as genetic patterns in response to temperature changes along the latitudinal gradient), these responses may be adaptive (Hoffmann et al. 2005, Huey and Berrigan 1996, Fallis et al. 2014). Line-cross and clinal analyses have been performed between tropical and temperate populations of *D. melanogaster* and *D. simulans* from the east coast of Australia to investigate whether clinal patterns and the genetic effects contributing to clinal divergence in mean wing centroid size, thorax length, wing-to-thorax ratio, cold and heat resistance differ under different developmental temperatures (18, 25 and 29 °C) (van Heerwaarden and Sgro 2011).

Specifically, we (van Heerwaarden and Sgro 2011), showed that the genetic basis of these traits is environment specific. However, we did not explore the genetic basis of divergence in the thermal plasticity of these traits.

The aim of this study was to use line cross analyses to investigate the quantitative genetic basis of clinal divergence in plasticity for thermal stress resistance (heat and cold resistance) and body size (wing centroid size and thorax size) in *D. melanogaster* and *D. simulans*. Specifically, as current plasticity evolution models ignore non-additive genetic variance, we were interested in determining the extent to which additive and non-additive genetic effects, particularly epistasis, underlie divergence in plastic responses. We took a reaction norm approach to studying plasticity, where different functions (parameters) of the reaction norm can be used to analyse average performance (trait mean, elevation), and the degree of plasticity - (slope) and the shape (curvature) - of the reaction norm (David et al. 1997; Berger et al. 2014; Murren et al. 2014). This approach enabled us to investigate whether the genetic basis of divergence in mean performance of each trait (elevation) was similar to the genetic basis of divergence in plasticity (slope and curvature). Although line cross analyses cannot directly distinguish which genetic model (allelic sensitivity or genetic regulation hypothesis) may underlie clinal divergence in plasticity, as the gene regulation genetic model involves physiological epistasis, we were interested in investigating whether statistical epistasis contributed more frequently to divergence in environmental sensitivity (plasticity/ canalization) than mean performance. Finally, we also explored whether the genetic effects underlying curvature and slope differed to explore whether different aspects of plastic responses evolve via different genetic mechanisms to further understand the evolution of plasticity.



## METHODS

### Experimental populations

Populations of *D. melanogaster* and *D. simulans* were collected from a tropical (Gordenvale, north-eastern Queensland, 17°10'05"S, 145°49'55"E) and temperate (Melbourne, Victoria, 37°47'30"S, 145°26'05"E) location in January 2008. Twenty single field-collected females of each species from each location were used to found 20 isofemale lines for each population. Three generations after field collection, a mass-bred population was initiated with 20 males and 20 females from each of the 20 isofemale lines, per species per location. Each mass-bred population was maintained at 25 °C under a 12:12 light: dark cycle at a census population size of approximately 1000 individuals across 3 x 250 ml bottles containing 20 ml of potato, yeast and sucrose media. Line crosses were performed after six (*D. simulans*) or nine (*D. melanogaster*) generations of mass breeding.

### Experimental design

Line crosses were performed to examine the relative contribution of different composite genetic effects (CGEs, e.g., additive, dominance, and epistatic gene effects) to the divergence in mean performance and plasticity for heat knock down time, chill coma recovery time (*D. melanogaster*), wing centroid size and thorax length, between tropical and temperate populations of *D. melanogaster* and *D. simulans*. Line crosses are a widely used quantitative genetics method for estimating the genetic architecture underlying divergence in a phenotype of interest between two strains or populations. This approach involves crossing two parental

strains to produce an F1 cohort, and performing subsequent crosses (e.g., F2, backcrosses, reciprocal crosses) to generate cohorts that have different combinations of parental genes. The observed mean phenotypic values of the parental and subsequent cross cohorts are then compared to the means expected from different genetic models that include additive, dominance, epistatic, maternal and/ or cytotype effects (maternal effects inherited from the organelles (e.g. mitochondria) or micro-organisms in the cytoplasm (Mather and Jinks 1982; Kearsey and Pooni 1996; Fox et al. 2004).

To generate cohorts that have different combinations of parental genes, we set up 14 crosses, similar to the procedure outlined in Gilchrist and Partridge (1999) (see Table 1), and as described in van Heerwaarden and Sgro (2011). Crosses were initiated between the two parental lines (temperate and tropical populations of each species) and the subsequent F1 and F2 generations. The F1 generation was then backcrossed to the parents. Each cross included a reciprocal cross. Re-establishing the parental and F1 crosses each generation allowed all 14 cohorts and parentals to be tested simultaneously after three generations of crossing. All crosses were performed at 25 °C, and were initiated with 100 virgin females and 100 males. In the third generation, when all crosses had been initiated, larvae from each cohort were picked over two subsequent days (one day for thermal stress traits and one day for body size traits) and placed into 18 replicate vials total (nine per day) at a density of 50 larvae per vial. Once the larvae from each block were picked, six replicate vials per cohort, were placed at 18°, 25° and 29 °C to develop, so that offspring could be measured for thermal stress resistance (day 1, three replicate vials) and for all morphological traits (day 2, three replicate vials) at each temperature.

### Climatic stress traits

Heat resistance was scored as knock down time (Hoffmann et al. 2002), while cold resistance was scored as chill coma recovery time (David et al. 1998) and both were measured on females only. Females were separated from males under CO<sub>2</sub> anaesthesia 48 hours prior to stressing (thus females were assumed to have mated), and heat and cold resistance was measured on seven and eight day old flies respectively. For heat resistance, individual flies were placed in 10 ml dry vials and submerged in a water bath heated to 38.5 °C and heat resistance was scored as the time taken (to the nearest second) for flies to be knocked down. For cold resistance, individual flies were placed in 10 ml dry vials and submerged in a water bath filled with 10% glycol solution and cooled to 0 °C for three hours. Chill coma recovery was assayed by scoring the time (to the nearest second) to recover (the ability to stand upright) at 25 °C following a chill coma induced by a cold shock. Ten to fifteen females per replicate vial were scored (total 30 - 45 females per cross). One-way ANOVAs showed no evidence for significant vial effects on cold or heat resistance at any temperature (data not shown).

### Morphological traits

Wing centroid size and thorax length were measured on the same 10 males and 10 females from each of three replicate vials, which had developed at 18°, 25° and 29 °C. The right wing (or the left wing if the right was damaged) was removed from individual flies with fine forceps and mounted on glass slides with double sided tape and protected with a cover slip. Wing images were captured with a Wild M3 dissector microscope attached to a digital

camera and land-marked for the eight junctions of longitudinal veins with the wing margins or cross-veins (Liefting et al. 2009). Their x and y co-ordinates were recorded using the program TPSDIG Version 1.31 written by F.J. Rohlf and wing size was calculated as centroid size, the square root of the sum of the squared interlandmark distances (Hoffmann and Shirriffs 2002). Thorax length was measured as outlined in Hoffmann et al. (2007). To check for measurement error, we measured repeatability for both thorax and wing centroid size estimates. Repeat measures were found to be highly correlated for thorax ( $r > 0.97$ ,  $N = 50$ ) and wing centroid size ( $r > 0.99$ ,  $N = 100$ ). Furthermore, one-way ANOVAs found no evidence for significant vial effects on wing centroid or thorax size at any temperature (data not shown).

## Analysis

We used the following equations to explore the divergence in reaction norm average performance (elevation, equation 1), sensitivity (slope, equation 2) and shape (curvature, equation 3) across the parental populations:

$$\text{Elevation} = \frac{\sum_1^n Z_i}{n} \quad (1)$$

$$\text{Slope} = \frac{\sum_1^{n-1} S_i}{n-1} ; S_i = \frac{Z_{i+1} - Z_i}{T_{i+1} - T_i} \quad (2)$$

$$\text{Curvature} = \frac{\sum_1^{n-2} C_i}{n-2} ; C_i = S_{i+1} - S_i \quad (3)$$

For trait  $Z$ ,  $n$  equals the number of test temperatures (3), and  $i$  represents the focal temperature (see Berger et al. 2014; Murren et al. 2014). Note that contrary to Murren et al. (2014), we have divided phenotypic differences by differences in temperatures ( $T$ ) to compute reaction norm slopes and curvatures, because the widths of the temperature intervals were not identical (i.e. the difference between 29 °C and 25 °C was smaller than between 18 °C and 25 °C). These were calculated separately for each cohort, trait, sex and species. As line cross analyses examine the composite genetic effects (CGEs) underlying population divergence in a phenotypic trait, we first examined divergence in reaction norm elevation, slope and curvature between the parental populations using t-tests. Plastic responses can include phenotypic changes with temperature, or underlying genetic/ physiological changes that maintain the same trait value across temperature (canalization) (Debat and David 2001). As such, we also used a one-way ANOVA, with elevation, slope and curvature as the dependent variable and cohort as a fixed effect, to look for any evidence of hybrid breakdown/ heterosis of canalization across the cohorts. We then further examined the CGEs underlying divergence in the reaction norm parameters for traits where we observed either evidence for divergence across the parental lines, and/ or across the cohorts (see below).

We used the software program SAGA (Software for Analysis of Genetic Architecture) to estimate the CGEs contributing to variation among cohorts for the reaction norm parameters for each trait, in each sex of each species (Blackmon and Demuth 2016). SAGA uses a full information-theoretic approach, that utilises the finite sample size corrected version of the Akaike information criterion (AICc) to explore all possible models and make unbiased and, when appropriate, model-averaged estimates of the contribution of CGEs to cohort means. This approach has the advantage of assessing the potential model space,

quantifying model selection uncertainty, and using model weighted averaging to accurately estimate CGEs. Traditional line cross analysis (e.g. joint scaling tests, Lynch and Walsh 1998) depend on identifying the best model and interpreting the CGEs that are included in that model. SAGA is able to obtain accurate estimates of the CGEs that are not dependent on the ability to specify one overall model as best. If the Akaike weight ( $w_i$ ) of the best model is 0.95 or greater, SAGA will perform parameter estimation under a single model. If no model reaches this threshold, then SAGA constructs a 95% confidence set of models that contain the minimum number of models whose  $w_i$  sum to 0.95. SAGA then computes model-averaged results for the 95% confidence set and provides estimates of variable importance ( $vi$ ) calculated by summing  $w_i$  of all models in which a CGE occurs. The  $vi$  score provides evidence that a CGE is important even if its contribution is small or poorly defined. Although one of the strengths of SAGA is that it alleviates the use of strict arbitrarily defined p-values, Blackmon and Demuth (2016) generally found CGEs with a  $vi$  score greater than 0.5 were most likely to be included in the model containing the 95% confidence model set.

To directly compare models with different CGEs for elevation, slope and curvature, we calculated AICc using the following equation (Burnham and Anderson 2002), where  $n$  is the number of cohorts and  $K$  is the number of parameters being estimated:

$$(4) \quad AICc = AIC + \frac{2K(K+1)}{n-K-1}$$

We first calculated AICc for the model that contained all the CGE's with a  $vi > 0.5$  (or 0.3 if there was high model uncertainty and no CGE's had a  $vi > 0.5$ ) for each reaction norm parameter (e.g. elevation). We then compared that AICc score to the model containing the CGEs with a  $vi > 0.5$  for the other reaction norm parameters for that trait. For example, for elevation in heat knock down in *D. melanogaster*, we compared the AICc for the model containing the CGE's with a  $vi > 0.5$  for elevation to the AICc scores for the models containing the CGE's with a  $vi > 0.5$  for slope and curvature, to ask whether the CGE's identified as important for elevation are a better fit than those identified as important for the other reaction norm parameters.

## RESULTS

The mean elevation, slope and curvature of all 14 cohorts for each interpopulation cross for each trait in each species are shown in Figures S1-3. We found significant divergence between the parental populations and/ or across the cohorts for trait mean - elevation, and plasticity - slope and curvature, for all traits (Table S1, Figures 1 and 2). Thus, we further examined the composite genetic effects (CGEs) underlying clinal divergence in elevation, slope and curvature for all traits in both species. Similar to Blackmon and Demuth (2016), no single model had a  $wi$  sufficient to ignore model selection uncertainty (Akaike weight ( $wi$ ) of the best model  $< 0.95$ ) for elevation, slope or curvature for any trait (Table S2), so for all traits, we used the model-averaged results for the 95% confidence set to get estimates of the CGEs underlying clinal divergence. In general, the lower the  $wi$  of the best model and or the more models contained in the 95% confidence set, the greater the degree of model selection

uncertainty (Blackmon and Demuth 2016). For a small number of traits, there was high model selection uncertainty for some of the reaction norm parameters, evidenced by low  $w_i$  and the high number of models required (Table S2). However, in these situations it is still possible to infer which CGEs are likely to be important, albeit with less confidence in estimating their true magnitude or sign because they depend on the other components in the model and standard errors will therefore overlap zero (Blackmon and Demuth 2016).

### **Patterns in the genetic architecture underlying clinal divergence in elevation, slope and curvature of stress and morphological traits**

Across all traits, autosomal additive effects contributed to clinal divergence in less than a third of crosses, and were detected more frequently for divergence in mean performance - elevation (observed in 55% of crosses), than for plasticity - slope (observed in 9% of crosses) or curvature (observed in 18%) (Table 2). Autosomal additive effects were important for divergence in elevation for heat knock down time in *D. simulans*, wing size in *D. melanogaster* females and males, wing size in *D. simulans* females, thorax size in *D. melanogaster* males and thorax size in *D. simulans* females (Figures 3, 4 and 5). For slope, autosomal additive effects were only important for divergence in thorax size in *D. melanogaster* females, and for curvature, autosomal additive effects contributed to divergence in cold in *D. melanogaster* females, and wing size in female *D. melanogaster* (Figures 3, 4 and 5).

Autosomal dominance effects were less common than autosomal additive effects, contributing to clinal divergence in elevation, slope and/ or curvature in only 21% of crosses



overall (Table 2). Similar to autosomal additive effects, autosomal dominance effects were observed more often for divergence in elevation (27% of crosses), than in slope (9% of crosses), but were observed in a similar frequency for curvature (13%) (Table 2). Autosomal dominance effects were important for divergence in curvature for heat knock down time in *D. melanogaster*, divergence in elevation and curvature for cold in *D. melanogaster*, divergence in curvature and elevation for wing size in *D. simulans* females and males respectively, and divergence in thorax size in *D. melanogaster* and *D. simulans* males (Figures 3, 4 and 5).

Autosomal epistatic and cytotype by autosomal epistatic effects were found more frequently than autosomal additive or dominance effects, detected in 58% and 42% of crosses respectively (Table 2). Autosomal epistatic effects were more common for curvature (found in 82% of crosses) than for elevation (45%) or slope (45%), while cytotype epistatic effects were more common for both measures of plasticity (slope: 64%, curvature: 45%), than for elevation (18%) (Table 2). Autosomal additive by additive epistatic effects were only important for divergence in plasticity, detected for slope and curvature in female heat knock down time in *D. melanogaster* and *D. simulans* females, slope for wing size in *D. simulans* males, and curvature in wing size and thorax size in *D. melanogaster* males (Figures 3, 4 and 5). Autosomal additive by dominance and/ or autosomal dominance by dominance epistasis were important for elevation for heat and cold resistance in *D. melanogaster*, wing size in *D. simulans* females and thorax size in *D. simulans* females and males; slope for heat resistance and wing and thorax size in *D. melanogaster* females; and curvature in heat resistance and thorax size in *D. simulans* females, wing size in *D. simulans* males and cold resistance, wing and thorax size in *D. melanogaster* females (Figures 3, 4 and 5).

Maternal effects were also common for divergence in elevation, slope and curvature and were observed in just under half of all crosses, while cytotypic effects were only observed in 24% of crosses overall. Maternal and cytotypic effects contributed more frequently to divergence in slope (73% and 36% respectively) than for elevation (45% and 18% respectively) or curvature (45% and 18% respectively), (Table 2).

Across all reaction norm parameters, the frequency of additive effects was similar for stress and morphological traits (Table 2). Dominance, autosomal and cytotypic epistasis, and maternal effects were more common for divergence in the stress traits, while cytotypic effects were slightly more common for divergence in the morphological traits (Table 2).

### **Differences in genetic architecture between elevation, slope and curvature**

Generally, different composite genetic effects (CGE's) were identified as important for explaining clinal divergence in the different reaction norm parameters (i.e. elevation, slope and curvature) for each trait (Figures 3, 4 and 5). Indeed, when we compared the AICc scores for the models including CGEs with a variable importance ( $v_i$ ) of more than 0.5 (or  $v_i > 0.3$  if there was high model selection uncertainty) for each reaction norm parameter (e.g. elevation) against AICc scores which included the same CGE's identified for the other reaction norm parameters (e.g. slope and curvature) for each trait we generally found strong support that the models that best described divergence in elevation, slope and curvature were different (AICc best model  $> 2$  AICc than the alternate model) (Tables S3-5), suggesting that the genetic basis underlying divergence in these reaction norm parameter values differ. The only traits where this was not the case were slope (versus elevation and curvature) and

curvature (versus elevation and slope) for wing size in *D. melanogaster* males, slope (versus curvature) for wing size in *D. simulans* males (Table S4), and elevation (versus curvature) for thorax size in *D. simulans* females, and slope (versus curvature) and curvature and elevation for thorax size in *D. simulans* males (Table S5).

We also found little evidence that the genetic basis underlying divergence in elevation, slope and curvature is similar across the sexes. With the exception of curvature for thorax size in *D. simulans* and *D. melanogaster*, and elevation and slope for wing size in *D. melanogaster*, the models that best described divergence in elevation, slope and curvature differed across sexes in both species (AICc best model for females  $> 2$  AICc than the alternate model for males and vice versa) (Tables S6-7). Additionally, we found little evidence that the genetic basis underlying divergence in elevation, slope and curvature is similar across species. With the exception of slope for wing and thorax size in males, and curvature in thorax size in females, all of the models that best described divergence in elevation, slope and curvature differed across *D. melanogaster* and *D. simulans* (AICc best model for *D. melanogaster*  $> 2$  AICc than the alternate model for *D. simulans* and vice versa) (Tables S8-10).

## DISCUSSION

The genetic basis of phenotypic plasticity is not well understood, despite being important for modelling and understanding the evolution of plastic responses in nature. The

extent to which additive or non-additive effects contribute to the evolution of plastic responses is still not clear even though physiological epistasis (that is, epistasis between regulatory and structural loci) is central to evolutionary models of plasticity (including the gene regulation model) and canalization (Scheiner 1993; Wagner et al. 1997; Rice 1998; Flatt 2005). We provide the first detailed examination of the quantitative genetic basis of divergence in mean thermal response and thermal plasticity in locally adapted outbred populations of *D. melanogaster* and *D. simulans*. Our analysis suggests that the genetic basis of population divergence in reaction norm elevation, slope and curvature differs, and that epistasis frequently contributes to divergence in plasticity. We discuss the implication of these findings below.

There has been considerable interest in the extent to which epistasis may contribute to the adaptive divergence of key traits (Wright 1931; Whitlock et al. 1995; Fenster et al. 1997; Wade and Goodnight 1998), as well as its role in the maintenance of genetic variance and the rate of evolution (Hansen 2013). Using crosses between diverged populations or lines/ stocks, several studies have shown that epistasis is indeed important for adaptive divergence in morphological, stress and fitness traits (Hard et al. 1992; Armbruster et al. 1997; Gilchrist and Partridge 1999; Schiffer et al. 2006; van Heerwaarden and Sgro 2011). However, only a small number of studies have used this approach to examine the quantitative genetic basis of plasticity (e.g. Perkins and Jinks 1973; Connolly and Jinks 1975; Pooni et al. 1987). These studies found evidence that dominance and epistasis contribute to both mean performance and environmental sensitivity (plasticity), suggesting that non-additive effects may commonly underlie the evolution of plasticity. Nonetheless, these studies did not consider maternal effects, which may contribute significantly to  $F_2$  breakdown, nor did they use a

comprehensive crossing design that allowed the partitioning of different epistatic effects. Furthermore, no such studies have been performed to understand the genetic basis of divergence in plasticity in outbred populations originating from, and adapted to, different biogeographical habitats. While we used flies that had been in the laboratory for 6 - 9 generations before starting the experiment, and may thus have undergone some level of laboratory adaptation (Santos et al. 2012), this is the first study to our knowledge that has used a comprehensive crossing design to examine the genetic architecture underlying plasticity using locally adapted outbred populations recently collected from different biogeographical locations. Consistent with these earlier studies (e.g. Perkins and Jinks 1973; Connolly and Jinks 1975; Pooni et al. 1987), overall we observed that additive, dominance, epistatic and maternal effects all contributed to divergence in the mean (elevation) and plasticity (slope and curvature) of body size and/or thermal tolerance in *D. melanogaster* and *D. simulans*. Thus, both additive and non-additive genetic effects underlie the evolution of the mean of these traits (van Heerwaarden and Sgrò 2011) and their plastic responses to temperature (this study).

In both species, we also observed that the relative contribution of non-additive genetic effects underlying the divergence in trait mean (elevation) and thermal plasticity (slope and curvature) differed. Autosomal additive effects were more common for divergence in mean values (elevation), than for plasticity (slope or curvature), while autosomal dominance effects contributed more frequently to divergence in elevation and curvature than for slope. Cytotype and maternal effects were detected more frequently to divergence in slope than elevation or curvature. Importantly, epistatic effects were more common for plasticity than trait mean (elevation), consistent with Connolly and Jinks (1975). Across all traits, autosomal and/ or

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cytotype epistasis was observed in 91% of crosses for both slope and curvature (c.f. to 45% of crosses for elevation) (Table 2), suggesting that epistasis may be crucial to the evolution of plastic responses. This finding has important implications for genetic models of plasticity, which generally ignore non-additive genetic variation (Via et al. 1995; Berrigan and Scheiner 2004). It is also important for understanding and predicting how traits and their plasticity evolve (Carter et al. 2005; Hallander and Waldmann 2007). This is because studies have shown that epistasis can alter additive genetic variance of traits under selection, and some models suggest that epistasis may accelerate or constrain evolutionary responses to selection (Carter et al. 2005; Carlborg et al. 2006; Hallander and Waldmann 2007). Specifically, positive epistasis, where genes tend to reinforce each other's effects in the direction of selection, will increase additive genetic variance and accelerate the response to selection, while negative epistasis, where genes tend to diminish each other's effects in the direction of selection and reduce additive genetic variance, will reduce the response. (Hansen and Wagner 2001). Thus, if epistasis is pervasive, predicted evolutionary responses in traits or their plasticity based on estimates of additive genetic variance alone may be inaccurate.

In addition to differences between the genetic basis of mean (elevation) and plasticity (slope), we also found that the quantitative genetic basis of the two reaction norm parameters that describe plasticity (slope and curvature) differs. Although quantitative genetic models have not explored the effect of additive and non-additive genetic effects on the evolution of plasticity separately, de Jong and Gavrillets (2000) predicted that the additive genetic variance for the elevation and slope of a linear reaction norm, as well as their covariance, should depend on the level of variation in their developing environment, decreasing with increasing variation in the environment of development. Furthermore, Lande (2009) found that after

large and sudden changes in environment exceeding typical background environmental fluctuations, the proportion of additive genetic variance of a trait in the new environment may increase due to additive genetic variance in plasticity. Given that plasticity is predicted to evolve when populations experience spatial and/ or temporal environmental heterogeneity (Via and Lande 1985; Gabriel and Lynch 1992; Gabriel et al. 2005), environmental variability may directly influence the evolution of plasticity. Consistent with this prediction, de Jong and Gavrillets (2000) showed that the additive genetic variance for elevation and slope (and their genetic covariance) of morphological traits in *Drosophila* both decrease with increasing variation in the environment of development. Furthermore, as long as some loci that influence only the slope are present (i.e. pleiotropy isn't complete, as predicted under the gene regulation hypothesis), the genetic variance in slope (plasticity) is predicted to decrease faster with increased environmental variance than the genetic variance in elevation (mean) (de Jong and Gavrillets 2000). We have only measured the relative contributions of different genetic effects to population divergence, rather than assessing standing additive and non-additive genetic variance, and are thus unable to directly compare our results to those of de Jong and Gavrillets (2000). However if the additive genetic variance in elevation and slope (and their genetic covariance) does indeed change across environment as predicted (de Jong and Gavrillets 2000), then the differences in the overall quantitative genetic architecture (additive, dominance, epistasis, maternal etc) for mean trait values (elevation) and plasticity (slope and curvature) between populations originating from, and locally adapted to, different environments as observed in our study may be expected.

When exploring the level of evolutionary divergence in different reaction norm parameters across diverged populations and species, Murren et al. (2014) found that

divergence in plasticity (slope and curvature) among closely related species were greater than divergence in trait means (elevation), indicating that microevolutionary changes in plasticity may be more common than evolved shifts in trait means. They also found that differences in curvature among closely related species were greater than differences in slope. Higher levels of divergence in plasticity (slope and curvature) than elevation contrast with quantitative genetic experiments that have shown that heritability for plasticity is lower than for elevation (Scheider 2003), but are consistent with theoretical models, which suggest that plasticity may increase with large and sudden changes in environment, with relatively little change to the additive genetic variance of the trait in the original environment (Lande 2009). These results suggest that there is significant potential for the additive genetic variance and selection responses to differ for trait mean (elevation) and plasticity (slope and curvature). Our study, which revealed differences in genetic architecture for divergence in trait mean and plasticity, are consistent with such predictions. In addition, our results are consistent with models that suggest that epistasis may play an important role in evolutionary shifts in traits and plasticity (Carter et al. 2005).

We also found that the genetic effects underlying divergence in plasticity for heat/cold resistance and body size generally differed, suggesting that the genetic basis of plasticity may be trait-specific. Given the close association between body and environmental temperature in ectotherms (Cossins and Bowler 1987), resistance to temperature extremes is likely to be important for surviving and adapting to different thermal environments, as well as dictating species distributions (Sunday et al. 2011; Overgaard et al. 2014). Overall, we found that dominance, autosomal and cytotypic epistasis, and maternal effects were slightly more common for divergence in the plasticity of heat/cold resistance compared to size. Roff and



Emerson (2006) hypothesised that fitness traits would show higher levels of epistasis and dominance because these traits should be subjected to intense natural selection (Kingsolver et al. 2012), which is predicted to deplete additive genetic variation, and leave segregating loci with primarily dominance and epistatic effects. Indeed, in a meta-analysis of line-cross studies, they observed more epistasis for fitness traits than morphological traits (Roff and Emerson 2006). However, in comparisons between tropical and temperate populations of *Drosophila*, the frequency of non-additive genetic effects was higher for morphological traits than stress traits (van Heerwaarden and Sgro 2011, Gilchrist and Partridge 1999; Kennington 2001). Our data suggests that different genetic architectures underpin the mean and plasticity of morphological and climatic stress resistance traits.

Finally, the genetic effects underlying population divergence in the different reaction norm parameters differed between *D. melanogaster* and *D. simulans*, and across the sexes. Despite sharing similar distributions and thus similar environmental conditions, these results suggest that similar selective forces can cause divergence in traits and their plasticity via quite different types of gene action and interaction. These results are also consistent with evolutionary patterns observed in a recent meta-analysis (Murren et al. 2014), which showed that divergence in reaction norm shape varies between organisms, traits, and environments. Taken together, these findings suggest that the evolution of plasticity is complex, and that this complexity must be considered in future studies of plasticity. While we only examined one population from each latitude for each species, which may limit our ability to make general statements about which particular genetic effects underlie clinal divergence for each trait specifically, our replication at the species level supports our general findings that different genetic effects underlie mean performance and plasticity, and that epistasis contributes

frequently to evolutionary divergence in plasticity more broadly. In addition, the fact that independent studies on *D. melanogaster* have repeatedly shown clinal patterns in the traits we examined (e.g. heat: Hoffmann et al. 2002; Sgro et al. 2010; Cockerell et al. 2014; size: James et al. 1995; 1997; van Heerwaarden and Sgro 2011; cold: Hoffmann et al. 2002, 2005), that also reflect the level of population divergence we see in this study, suggests that the divergence in these traits and their plasticity are likely due to local adaptation.

In conclusion, the genetic basis of geographic divergence in mean performance and both measures of plasticity differs in both *D. melanogaster* and *D. simulans*. In addition, the prevalence of non-additive genetic effects for plasticity, as well as trait means (van Heerwaarden and Sgro 2011) suggests that non-additive genetic effects (particularly epistasis) are important for the evolution of both trait means and trait plasticity. Current evolutionary models of adaptive responses to environmental change largely assume that additive genetic effects will be important for the evolution of both trait mean and plasticity (Fisher 1930; de Jong 1995; Coyne et al. 1997, 2000; Hill et al. 2008). The extent to which predictions of evolutionary shifts in plasticity and trait means will differ if non-additive effects are explicitly considered remains to be assessed.

#### **AUTHOR CONTRIBUTIONS**

C.M.S. and B.V.H. designed the experiments. B.V.H. performed the experiments and analysed the data. Both authors contributed to the final manuscript.

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## DATA ARCHIVING

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Figure 1. Thermal reaction norm of mean heat knockdown time at 38.5 °C in (a) *D. melanogaster* females and (b) *D. simulans* females, and mean chill coma recovery time at 25 °C after 3 hours at 0° C in (c) *D. melanogaster* females under different developmental/ adult acclimation temperatures. Error bars are one standard error.

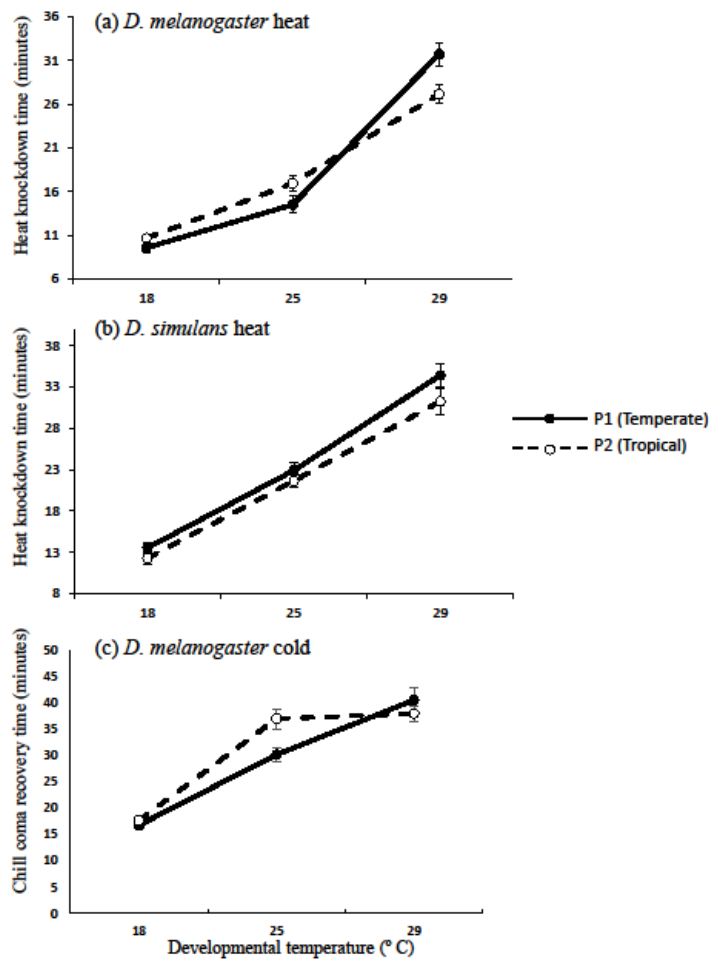


Figure 1

Figure 2. Thermal reaction norm for mean wing centroid size in *D. melanogaster* (a) females and (b) males, mean wing centroid size in *D. simulans* (c) females and (d) males; mean thorax size in *D. melanogaster* (e) females and (f) males; and mean thorax size in *D. simulans* (g) females and (h) males. Error bars are one standard error.

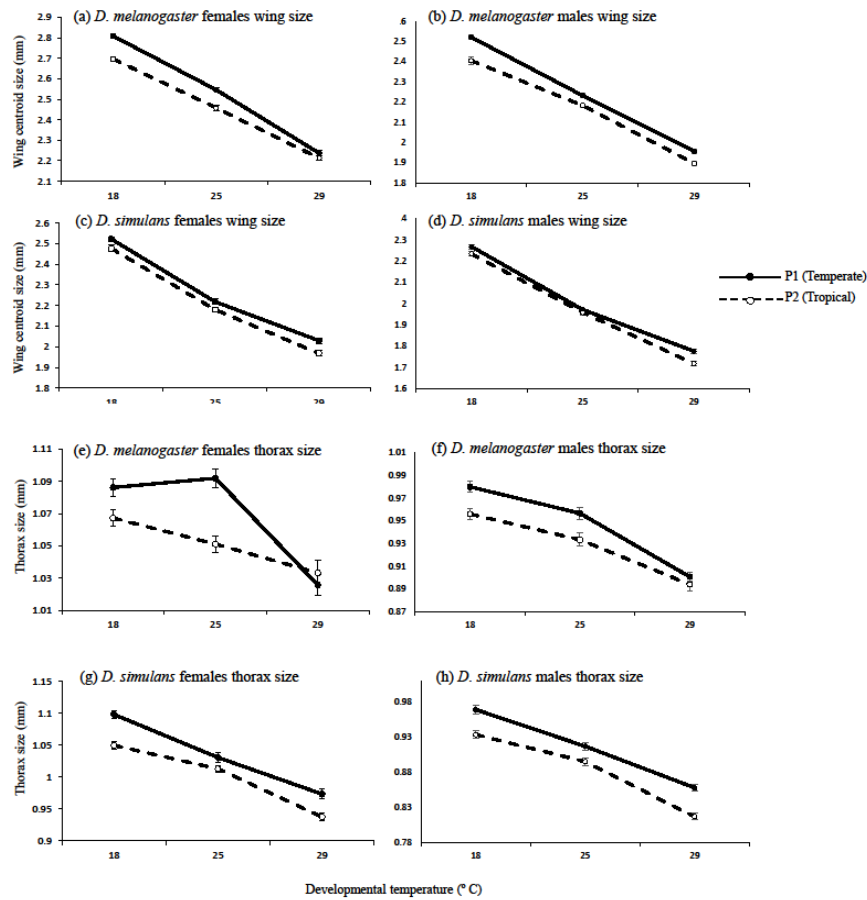


Figure 2

Figure 3. Model weighted average values for the different composite genetic effects (CGEs) contributing to the divergence between tropical and temperate populations in reaction norm elevation, slope and curvature for heat knock down time in *D. melanogaster* (a, b, c) and *D. simulans* (e, f, g), and chill coma recovery time in *D. melanogaster* (h, i, j). Bars are coloured based on  $vi$  (variable importance) score, which provides evidence that a CGE is important even if its contribution is small or poorly defined. Error bars indicate the unconditional standard errors and stars indicate which CGE's are compared across the different reaction norm parameters. The direction of each CGE indicates whether the average effect across all

loci is positive or negative in relation to the reference population, P1. m = mean, Aa = autosomal additive, Ad = autosomal dominance, AaAa = autosomal additive by additive epistasis, AaAd = autosomal additive by dominance epistasis, AdAd = autosomal dominance by dominance epistasis, Mea = additive maternal, Med = dominance maternal, Ca = additive cytotypic, CaAa = additive cytotypic by autosomal additive epistasis, and CaAd = additive cytotypic by autosomal dominance epistasis.

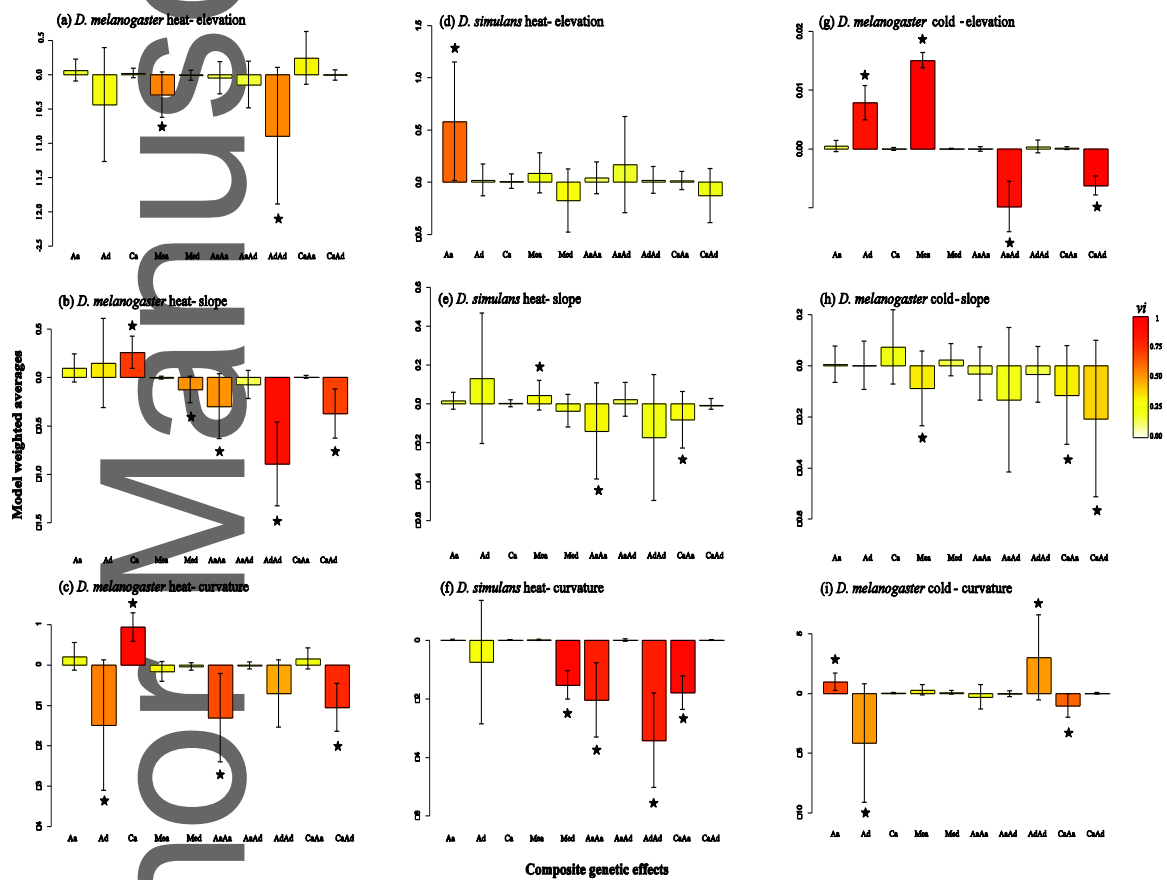


Figure 4. Model weighted average values for the different composite genetic effects (CGEs) contributing to the divergence between tropical and temperate populations in reaction norm elevation, slope and curvature for wing centroid size in *D. melanogaster* females (a, b, c) and males (d, e, f), and *D. simulans* females (g, h) and males (i, j, k). Bars are coloured based on  $v_i$  (variable importance) score, which provides evidence that a CGE is important even if its contribution is small or poorly defined. Error bars indicate the unconditional standard errors



and stars indicate which CGE's are compared across the different reaction norm parameters/ traits/ sex. The direction of each CGE indicates whether the average effect across all loci is positive or negative in relation to the reference population, P1. m = mean, Aa = autosomal additive, Ad = autosomal dominance, AaAa = autosomal additive by additive epistasis, AaAd = autosomal additive by dominance epistasis, AdAd = autosomal dominance by dominance epistasis, Mea = additive maternal, Med = dominance maternal, Ca = additive cytotype, CaAa = additive cytotype by autosomal additive epistasis, and CaAd = additive cytotype by autosomal dominance epistasis.

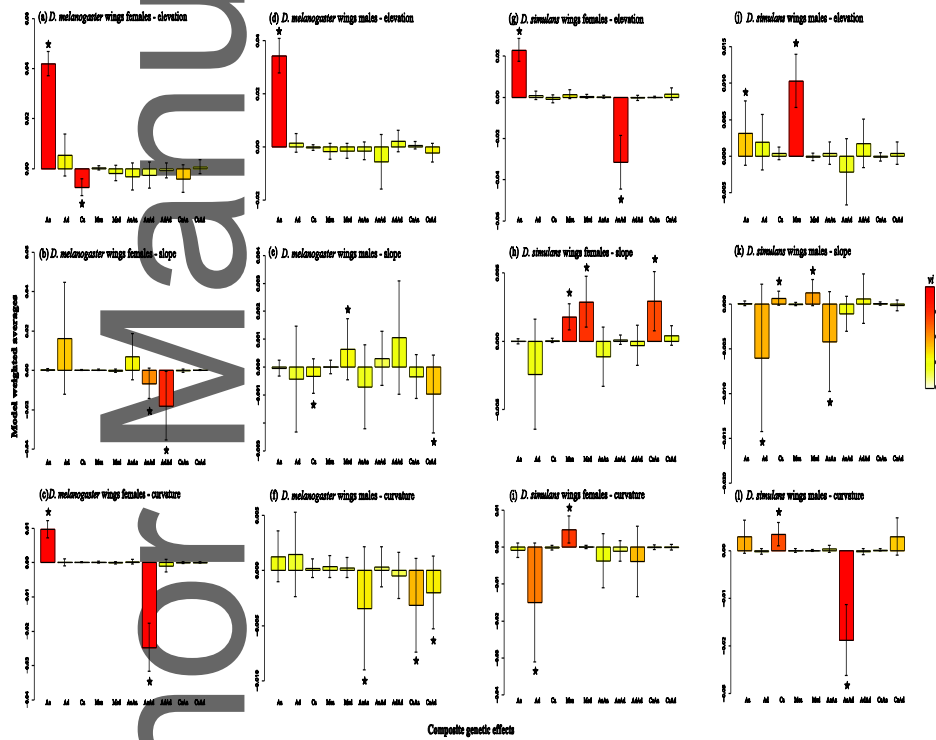


Figure 5. Model weighted average values for the different composite genetic effects (CGEs) contributing to the divergence between tropical and temperate populations in reaction norm elevation, slope and curvature for thorax size of *D. melanogaster* female (a, b, c) and males (d, e), and *D. simulans* females (f, g, h) and males (i, j). Bars are coloured based on  $v_i$  (variable importance) score, which provides evidence that a CGE is important even if its contribution is small or poorly defined. Error bars indicate the unconditional standard errors

and stars indicate which CGE's are compared across the different reaction norm parameters.

The direction of each CGE indicates whether the average effect across all loci is positive or negative in relation to the reference population, P1. m = mean, Aa = autosomal additive, Ad = autosomal dominance, AaAa = autosomal additive by additive epistasis, AaAd = autosomal additive by dominance epistasis, AdAd = autosomal dominance by dominance epistasis, Mea = additive maternal, Med = dominance maternal, Ca = additive cytotype, CaAa = additive cytotype by autosomal additive epistasis, and CaAd = additive cytotype by autosomal dominance epistasis

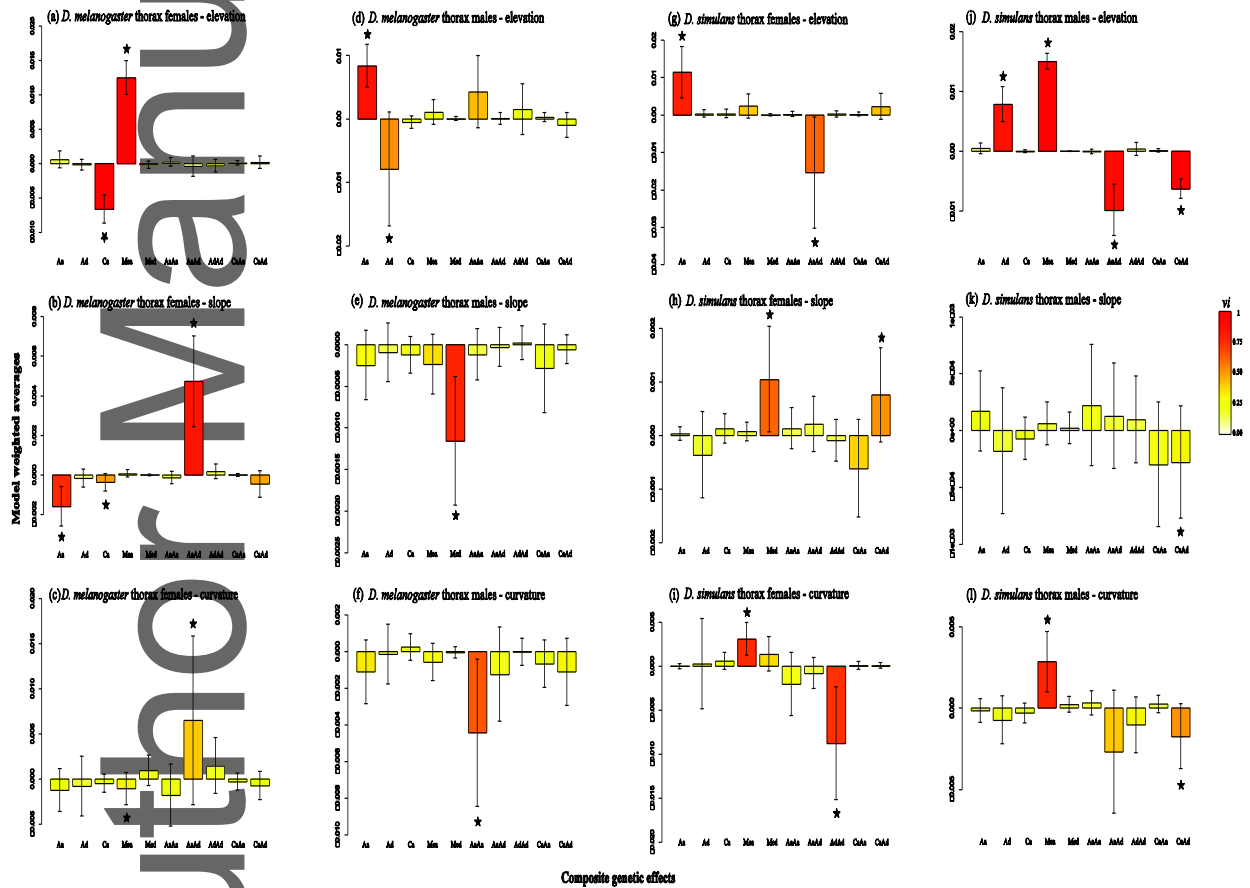


Table 1. Outline of crossing scheme (performed separately for each species), the relative contribution of the composite genetic effects (CGEs) to each cohort and the proportion of P1

Cohort	Cross (dam x sire)	<i>m</i>	<i>Aa</i>	<i>Ad</i>	<i>AaA</i> <i>a</i>	<i>AaA</i> <i>d</i>	<i>AdA</i> <i>d</i>	<i>Me</i> <i>a</i>	<i>Me</i> <i>d</i>	<i>C</i> <i>a</i>	<i>CaA</i> <i>a</i>	<i>CaA</i> <i>d</i>	Proportion of P1 genes
	Temperature population											0	1
P1	Tropical population	1	1	0	1	0	0	1	0	1	1	0	0
P2	P1 x P2	1	-1	0	1	0	0	-1	0	-1	1	1	0.5
F1	P2 x P1	1	0	1	0	0	1	1	0	1	0	-1	0.5
F1R	(P1 x P2) x (P1 x P2)	1	0	5	0	0	0.25	0	1	1	0	0.5	0.5
F2	(P2 x P1) x (P2 x P1)	1	0	5	0	0	0.25	0	1	-1	0	-0.5	0.5
F2R	P1 x (P1 x P2)	1	5	5	0.25	0.25	0.25	1	0	1	0.5	0.5	0.75
B1a	P1 x (P2 x P1)	1	5	5	0.25	0.25	0.25	1	0	1	0.5	0.5	0.75
B1b	(P1 x P2) x P1	1	5	5	0.25	0.25	0.25	0	1	1	0.5	0.5	0.75
B1Ra	(P2 x P1) x P1	1	5	5	0.25	0.25	0.25	0	1	-1	-0.5	-0.5	0.75
B1Rb												-0.5	0.25
B2a	P2 x (P1 x P2)	1	5	5	0.25	0.25	0.25	-1	0	-1	0.5	-0.5	0.25
B2b	P2 x (P2 x P1)	1	5	5	0.25	0.25	0.25	-1	0	-1	0.5	-0.5	0.25
B2Ra	(P1 x P2) x P2	1	5	5	0.25	0.25	0.25	0	1	1	-0.5	0.5	0.25
B2Rb	(P2 x P1) x P2	1	5	5	0.25	0.25	0.25	0	1	-1	-0.5	-0.5	0.25

genes.

m = mean, Aa = autosomal additive, Ad = autosomal dominance, AaAa = autosomal additive by additive epistasis, AaAd = autosomal additive by dominance epistasis, AdAd = autosomal dominance by dominance epistasis, Mea = additive maternal, Med = dominance maternal, Ca = additive cytotype, CaAa = additive cytotype by autosomal additive epistasis, and CaAd = additive cytotype by autosomal dominance epistasis.

**Table 2** Summary of the composite genetic effects underlying divergence in elevation, slope and curvature (averaged across all traits/ sex/ species), in the stress and morphological traits (averaged across sex/ species) and overall. The discrete values are the number of crosses where these effects were found to be important ( $v_i > 0.5$ , or  $v_i > 0.3$  where there was high model selection uncertainty) and the percentage is the fraction of all crosses in a particular category where these effects were observed.

	N	Additive	Dominance	Autosomal Epistasis	Cytotype by Autosomal Epistasis	Epistasis Overall	Cytotype	Maternal
<b>Elevation</b>	16	5 (31%)	3 (19%)	5 (31%)	2 (13%)	5 (31%)	2 (13%)	5 (31%)
<b>Slope</b>	11	1 (9%)	1 (9%)	5 (45%)	7 (64%)	1 (9%)	4 (36%)	8 (73%)
	1			1 (100%)	1 (100%)	0 (0%)	1 (100%)	1 (100%)

<b>Curvatu</b>	1	2	(18	3	(27%)	9	(82	5	(45	1	(91	2	(18	5	(45
<b>re</b>	1		%)				%)		%)	0	%)		%)		%)
<b>Stress</b>	9	2	(22	3	(33%)	7	(78	7	(78	8	(89	2	(22	6	(67
			%)				%)		%)		%)		%)		%)
<b>Morph</b>	2	7	(29	4	(17%)	1	(50	7	(29	1	(71	6	(25	1	(50
	4		%)			2	%)		%)	7	%)		%)	2	%)
<b>Overall</b>	3	9	(27	7	(21%)	1	(58	1	(42	2	(76	8	(24	1	(55
	3		%)			9	%)	4	%)	5	%)		%)	8	%)

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