



# Genetic basis of thermal plasticity variation in Drosophila melanogaster body size

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## OPEN ACCESS

Citation: Lafuente E, Duneau D, Beldade P (2018) Genetic basis of thermal plasticity variation in *Drosophila melanogaster* body size. PLoS Genet 14(9): e1007686. https://doi.org/10.1371/journal. pgen.1007686

**Editor:** Gregory P. Copenhaver, The University of North Carolina at Chapel Hill, UNITED STATES

Received: March 29, 2018

Accepted: September 10, 2018

Published: September 26, 2018

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**Data Availability Statement:** All relevant data are within the paper and its Supporting Information files.

Funding: Financial support for this work was provided by the Portuguese science funding agency, Fundação para a Ciência e Tecnologia, FCT: PhD fellowship to EL (SFRH/BD/52171/2013), and research grant to PB (PTDC/BIA-EVF/0017/2014); French research funding agency, Agence Nationale de la Recherche, ANR: Laboratory of Excellence TULIP, ANR-10-LABX-41 and ANR-11-IDEX-0002-02 (support for DD and PB) and French

# **Abstract**

Body size is a quantitative trait that is closely associated to fitness and under the control of both genetic and environmental factors. While developmental plasticity for this and other traits is heritable and under selection, little is known about the genetic basis for variation in plasticity that can provide the raw material for its evolution. We quantified genetic variation for body size plasticity in Drosophila melanogaster by measuring thorax and abdomen length of females reared at two temperatures from a panel representing naturally segregating alleles, the Drosophila Genetic Reference Panel (DGRP). We found variation between genotypes for the levels and direction of thermal plasticity in size of both body parts. We then used a Genome-Wide Association Study (GWAS) approach to unravel the genetic basis of inter-genotype variation in body size plasticity, and used different approaches to validate selected QTLs and to explore potential pleiotropic effects. We found mostly "private QTLs", with little overlap between the candidate loci underlying variation in plasticity for thorax versus abdomen size, for different properties of the plastic response, and for size versus size plasticity. We also found that the putative functions of plasticity QTLs were diverse and that alleles for higher plasticity were found at lower frequencies in the target population. Importantly, a number of our plasticity QTLs have been targets of selection in other populations. Our data sheds light onto the genetic basis of inter-genotype variation in size plasticity that is necessary for its evolution.

## Author summary

Environmental conditions can influence development and lead to the production of phenotypes adjusted to the conditions adults will live in. This developmental plasticity, which can help organisms cope with environmental heterogeneity, is heritable and under selection. Its evolution will depend on available genetic variation. Using a panel of *D. melanogaster* flies representing naturally segregating alleles, we identified DNA sequence variants associated to variation in thermal plasticity for body size. We found that these variants correspond to a diverse set of functions and that their effects differ between body parts and properties of the thermal response. Our results shed new light onto the long discussed genes for plasticity.



research centre, Centre National de la Recherche Scientifique, CNRS: International Associated Laboratory, LIA BEEG-B (support for DD and PB), and the People Programme (Marie Curie Actions) of the European Union's Seventh Framework Programme (FP7/2007-2013) under REA grant agreement (PCOFUND-GA-2013-609102), through the PRESTIGE programme coordinated by Campus France (support for DD). This work was developed with the support of the research infrastructure Congento, project LISBOA-01-0145-FEDER-022170, co-financed by Lisboa Regional Operational Programme (Lisboa 2020), under the Portugal 2020 Partnership Agreement, through the European Regional Development Fund (ERDF), and Foundation for Science and Technology (FCT, Portugal). The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript.

**Competing interests:** The authors have declared that no competing interests exist.

#### Introduction

Body size has a great impact on the performance of individuals [1, 2], as well as that of species [3]. Diversity in this trait is shaped by the reciprocal interactions between the developmental processes that regulate growth, and the evolutionary forces that determine which phenotypes increase in frequency across generations [4]. Body size varies greatly within and between populations [2, 5] and is controlled by both genetic and environmental factors [6–9]. Studies in different animal models have provided insight about the selection agents that shape the evolution of body size (e.g. predators [10, 11], mates [12], thermal regimes [13, 14]), and about the molecular mechanisms that regulate body size and body proportions during development [15–18].

Body size is also a prime example of the environmental regulation of development, or developmental plasticity [19, 20], and it is influenced by different factors, including nutrition and temperature. This plasticity can help organisms cope with environmental heterogeneity and, as such, can have major implications for population persistence and adaptation [20–22]. Thermal plasticity in body size is ubiquitous among insects [23-25], with development under colder temperatures typically resulting in larger bodies, which is presumably advantageous for thermal-regulation [1, 26]. The environmental dependency of body size, and other plastic traits, is often studied using reaction norms, in which phenotypic variation is plotted as a function of gradation in the environment [27]. The properties of these reaction norms, including their shapes and slopes, can differ between genotypes [28-30], and the genes underlying such variation can fuel the evolution of plasticity. Little is known about what these genetic variants are and what types of functions they perform (e.g. perception of environmental cues, conveying external information to developing tissues, or executing actions in developing plastic organs). It is also unclear to what extent the loci contributing to variation in thermal plasticity in size are the same for different body parts, and whether the loci contributing to variation in size plasticity are the same that underlie inter-individual variation in body size at a given temperature.

Studies in *D. melanogaster* have provided much insight about the evolution and development of body size, body proportions, and body size plasticity [7, 31–38]. Size differences among populations, including clinal [31, 39] and seasonal variation [40], and among individuals within a population, are due to the effects of genes, environment, as well as genotype-by-environment interactions [33, 41–43]. While we have increasing detailed knowledge about the genetic basis of adaptation and of natural variation for many adaptive traits in *D. melanogaster* and other species [6, 43–45], little is known about the genetic basis of variation in plasticity. Widely-accessible mapping panels [46, 47] allowed the dissection of the genetic architecture of various quantitative traits in *D. melanogaster* [48–50], including body size [51]. However, with a few recent exceptions [52–55], the genetic basis of phenotypic variation has been investigated under a single environmental condition, precluding assessment of differences between environments and of the genetic basis of plasticity itself. Series of isogenic lines from these mapping panels can be reared under different conditions to characterize reaction norms and ask about the genes that harbor allelic variation for their properties [53].

Here, we use a panel of *D. melanogaster* lines representing naturally segregating alleles from one natural population, the DGRP [46, 48], to characterize genetic variation for thermal plasticity in thorax and abdomen size, and to identify loci contributing to variation in the slopes of their thermal reaction norms. We document correlations between body size and body size plasticity, as well as correlations with other traits, using published data for the same panel. We also ask about the extent of overlap between QTLs for size and for size plasticity, and between QTLs for size plasticity of the different body parts. We then use different approaches



to validate and further characterize the role of selected QTLs, and to ascertain their pleiotropic effects.

#### Results

We measured thorax and abdomen length in adult females from different genotypes reared at two temperatures. We quantified effects of genotype (G), environment (E), and genotype-by-environment (GxE) interactions on body size (Fig 1), and explored correlations between body parts and between temperatures (Fig 2). We then used a GWAS approach to identify DNA sequence polymorphisms associated with variation in body size plasticity (Fig 3). Ensuing functional analyses of candidate QTLs validated and clarified their role in body size variation at different temperatures (Fig 4).

## Partitioning variation in body size

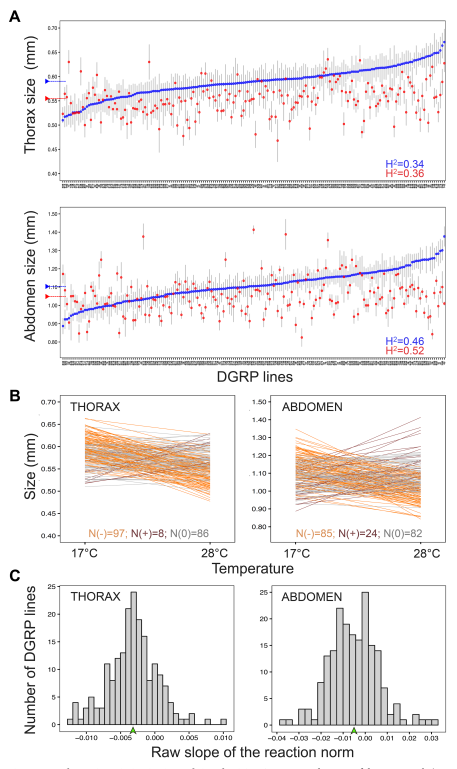
To assess the contribution of genotype and temperature to body size variation, we quantified length of abdomens and thoraxes (Fig 1A, S1A Fig) of adult females from ~196 isogenic lines reared at either 17°C or 28°C (S1 Dataset, S2 Table). We found significant differences between DGRP genotypes and between developmental temperatures, as well as significant GxE interaction effects (thermal plasticity) for the size of both body parts (Fig 1A, S1C Fig). We also found variation between individuals of (presumably) the same genotype and same rearing temperature; the coefficients of variation (CV) varied between 0.6 and 15 for thorax measurements and between 0.3 and 23.8 for abdomen measurements (Table 1). Broad-sense heritabilities for body size (at 17°C and 28°C) and body size plasticity (between-environment variation) varied between 33 and 52% (Table 1).

We computed a correlation matrix to assess relationships between the different components of variation in body size (Fig 2). We found a significant positive correlation between thorax and abdomen size within each rearing temperature (Fig 2A and 2B). We also found significant positive correlations between our measurements of thorax (both temperatures), but not abdomen size, and the size of several body parts measured in other studies [51] for the same genotypes reared at 25°C (S2A Fig). To address the question of association between body size and fitness, we measured correlations between our measurements of body size and a series of fitness related traits quantified in other studies of the DGRPs [46, 49, 50, 56]. We found significant correlations with chill coma recovery (thorax size at both temperatures and abdomen size at 28°C) and with survival upon infection with Metarhizium anisopliae fungi (abdomen size at 17°C), but not with longevity, starvation resistance, and other immune-defense traits (\$2B Fig). To address the question of what might explain the extent of inter-individual variation within genotype and temperature, we calculated correlations between our measurements of CV and of size. We found the CV to be: 1) positively correlated between body parts for flies reared at 28°C, but not for those reared at 17°C, and 2) negatively correlated with mean size for thoraxes, but not for abdomens (Fig 2B).

### Thermal reaction norms for body size

Using reaction norms, we studied the extent and properties of thermal plasticity for body size in the DGRP lines (Fig 1B). We calculated the slope of the regression lines for size across temperatures for each body part and DGRP genotype, and found genetic variation for both the intercept and slope of the reaction norms (Fig 1B, S2 Table). From each reaction norm, we extracted two properties of the thermal plasticity in body size: i) the absolute value of the slope, describing only the magnitude of the response to temperature, and ii) the raw value of the slope, which describes magnitude and direction of the response (Fig 1C, S1D Fig). Using the





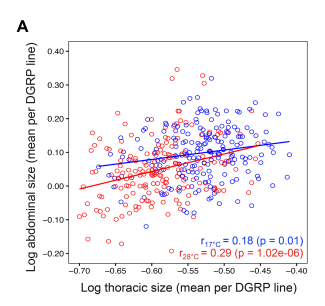
**Fig 1. Natural genetic variation in size and size plasticity. A.** Means and 95% confidence intervals (Y axis) for thorax (upper panel) and abdomen (lower panel) size in the DGRP lines (X axis) reared either at  $17^{\circ}$ C (blue) or at  $28^{\circ}$ C (red). DGRP lines are ranked by their mean size at  $17^{\circ}$ C. Dashed horizontal bar represents the mean value for all DGRP lines at a given temperature. Mean values ( $\mu$ ) and broad sense heritability ( $H^{2}$ ) estimates per body part and temperature can be found in Table 1. **B.** Reaction norms for thorax and abdomen sizes (Y axis) across temperatures (X axis) plotted as the regression fit for the model lm ( $Size \sim Temperature$ ) for each DGRP line. Reaction norms are colored by in relation to slope: slopes significantly different from zero are orange when positive and brown when negative, while slopes that were not significantly different from zero are gray (alpha = 0.05). Counts of each are shown on the bottom of each graph. Broad sense heritability estimates were:  $H^{2} = 0.33$  (thorax plasticity) and  $H^{2} = 0.48$  (abdomen plasticity). **C.** 



Frequency distribution for the raw value of the slope of the reaction norm in the DGRP lines. The mean value for the raw slope of all DGRP reaction norms is indicated with a green arrowhead.

https://doi.org/10.1371/journal.pgen.1007686.g002

reaction norms for 191 DGRP lines, we identified plastic and non-plastic genotypes. Slopes of the thermal reaction norm were significantly different from zero for 55% and 57% of the lines for the size of thoraxes and abdomens, respectively, with most of the plastic genotypes having smaller sizes when reared at higher temperatures (Fig 1B). However, we also found plasticity



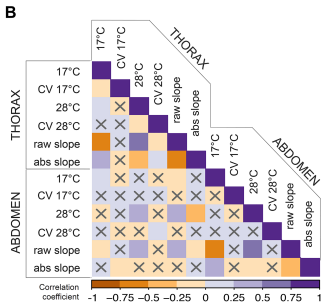
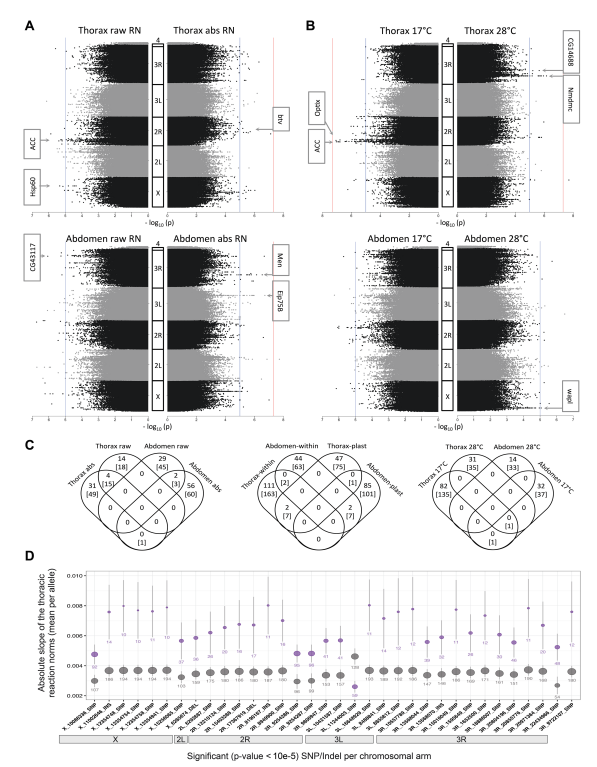


Fig 2. Phenotypic co-variation in size and size plasticity. A. Thoracic (X axis) and abdominal (Y axis) mean size per DGRP line at  $17^{\circ}$ C (blue) and at  $28^{\circ}$ C (red). Pearson correlation, r = 0.18 (p-value = 0.01) and partial Pearson correlation, r = 0.34 (p-value = 1.02e-06) for  $17^{\circ}$ C, Pearson correlation r = 0.29 (p-value < 2e-16) and partial Pearson correlation r = 0.34 (p-value < 2e-16) for  $28^{\circ}$ C. B. Heat map of Pearson's correlation coefficients between our measurements in thoraxes and abdomens: mean sizes and coefficients of variation (CV) at each temperature and raw and absolute slopes of the reactions norms. Non-significant correlations (p-value < 0.01) are indicated with an 'X'.

https://doi.org/10.1371/journal.pgen.1007686.g002





**Fig 3. Genetic variants influencing size and size plasticity. A-B.** Manhattan plots corresponding to the eight GWAS analyses performed. Horizontal lines are p-value < 10e-5 (blue) and p-value < 10e-8 (red). Gene names are shown for a subsample of significant SNPs/Indels, which were selected based on p-value, putative variant effect and associated genes (S3 and S4 Tables). **A.** GWAS for variation in size plasticity in thoraxes (upper panels) and abdomens (lower panels) and for either the raw (left) or the absolute (right) slopes of the reaction norms. **B.** GWAS for variation in size in thoraxes (upper panels) and abdomens (lower panels) at 17°C (left panels) or 28°C (right panels). **C.** Venn diagrams showing the number of candidate SNPs/Indels (number outside the brackets) and candidate genes (number within brackets) harboring those polymorphisms identified in the different GWAS. **D.** Mean



and 95% confidence interval of the absolute slope of the reaction norms for thorax size (Y axis) per allele (major allele in gray, minor allele in magenta) for each candidate plasticity QTL along the chromosomal arms (X axis). The position and identity of the polymorphisms in this figure is given by their annotation with Genome Release v.5.

https://doi.org/10.1371/journal.pgen.1007686.g003

in the opposite direction (i.e. genotypes with smaller flies at lower temperatures), corresponding to a positive significant slope for the thermal reaction norms: 8% of the DGRP reaction norms for the thorax and 22% for the abdomen (Fig 1B).

Even though the levels of thermal plasticity for thorax and abdomen size were significantly positively correlated (Fig 2B), lines having the highest levels of plasticity for one body part were not necessarily the most plastic for the other body part (S1B Fig). Furthermore, for thorax, but not abdomen, we found that genotypes with larger CVs had steeper negative reaction norms. Finally, we also found no correlation between our size plasticity measurements and various fitness-related traits measured in the DGRPs (S2B Fig), including longevity [49], starvation resistance, chill coma recovery [46], and immune-defense traits [50, 56].

# Genetic basis of variation in body size plasticity

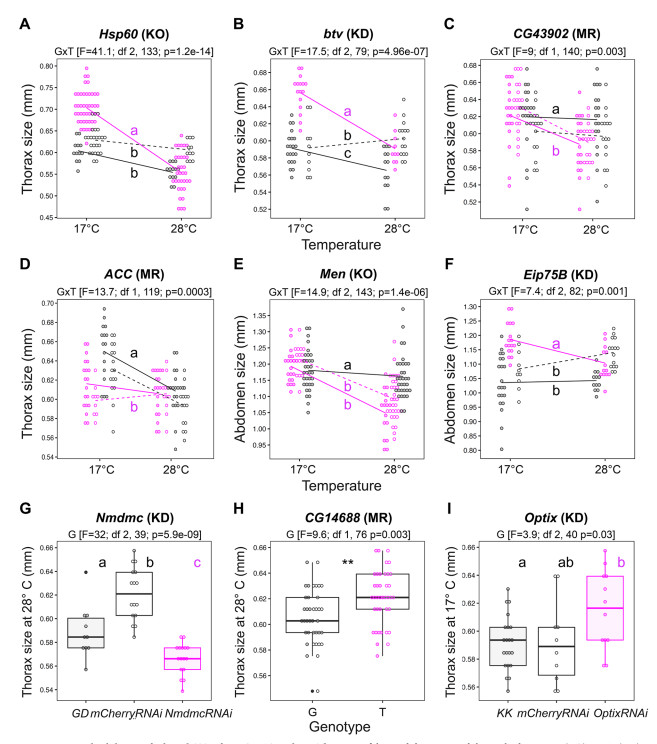
We used a GWAS approach to identify DNA sequence polymorphisms associated with variation in thermal plasticity for thorax and abdomen size in the DGRP. Because the loci carrying allelic variation for the direction and extent of environmental responsiveness are not necessarily the same, we used both the raw and absolute values of the slopes of the DGRP reaction norms as quantitative traits (Fig 3A, S3 Fig). The candidate QTLs significantly associated with variation in plasticity were typically only so in relation to a single property of the reaction norm (raw or absolute slope) or body part (thorax or abdomen; Fig 3C, S3 Table). We also found that these allelic variants fell within different genomic regions (e.g. UTR, intronic, coding) within or nearby 192 different putative genes (Table 2, S3 Table). Gene ontology enrichment analysis of the candidate QTLs showed an over-representation of vesicle-mediated processes (e.g. phagocytosis and endocytosis; S6A Fig), while network enrichment analyses (protein-protein interaction network followed by a KEGG pathways enrichment analysis) revealed an over-representation for SNARE interactions and Notch pathways (S6B Fig), both of which have been implicated in diverse biological functions [57-60]. We also found that in the vast majority of cases, alleles associated to increased environmental responsiveness were at lower frequencies in the DGRP (Fig 3D, S5 Fig).

To explore to what extent loci that contribute to variation in size plasticity also contribute to variation in body size within environments, we also performed GWAS analyses using body size at 17°C and 28°C as quantitative traits (Fig 3B, S4 Fig, S4 Table). This analysis revealed QTLs that were mostly environment- and body part-specific (Fig 3C, S4 Table), including no overlap between our candidate QTLs associated with variation in thorax and abdomen size and those reportedly associated with head size at 25°C [51]. Moreover, we found little overlap between candidate QTLs contributing to variation in size plasticity and those contributing to within-environment size variation (Fig 3C), but some overlap in terms of network enrichment (S6 Fig). None of our size traits or plasticity therein was affected by chromosomal inversions (p-value > 0.01), or by the genetic relatedness among DGRP lines (low and non-significant coefficients of phylogenetic signal Blomberg's K and Pagel's λ; S2C Fig).

#### Validation of selected GWAS hits

We selected a number of significant QTLs for validation via different approaches (Fig 4, Table 2, S2 Dataset). To test selected candidate genes, we used available null mutants and inducible gene knock-downs (with RNA interference using the Gal4/UAS system). If a





**Fig 4. Functional validation of selected QTLs from GWAS analyses.** The name of the candidate gene and the method—mutant (KO), RNAi (KD), or Mendelian Randomization (MR)—to test the different candidate QTLs is shown above each plot. For validations via mutant or RNAi, genotypes with impaired gene function are shown in magenta and control genotypes are shown in black. Similarly, for validations via Mendelian Randomization, the two populations fixed for the minor allele are shown in magenta and the two populations for the major allele are shown in black. **A.** Reaction norms in mutant Hsp60A/+ and controls Canton-S (filled circles, solid line) and Fm7a/Canton-S (empty circles, dashed line). **B.** Reaction norms in btv-RNAi/bab-Gal4 and control lines KK (filled circles, solid line) and mCherry-RNAi/bab-Gal4 (empty circles, dashed line). **C.** Reaction norms in the four MR populations for SNP 2R:7983239 within gene ACC. **E.** Reaction norms in Men mutants MenEx3/+ (filled circles, solid line) and MenEx55/+ (empty circles, dashed line) and control line w1118. **F.** Reaction norms in Eip75B-RNAi/bab-Gal4 and control lines KK (filled circles, solid line) and Mcherry-RNAi/bab-Gal4 (empty circles,



dashed line). **G.** Size at 28°C in *Nmdmc-RNAi/tub-Gal4* and control lines GD (filled circles) and *mCherry-RNAi/tub-Gal4* (empty circles). **H.** Size at 28°C in the four MR populations for SNP 3R:10678848 within gene *CG14688*. **I.** Size at 17°C in *Optix-RNAi/bab-Gal4* and control lines KK (filled circles) and *mCherry-RNAi/bab-Gal4* (filled circles). For the validations of plasticity QTLs, we tested the model lm (*Size* ~ *Genotype*\* *Temperature*) and for the validations of within-environment QTLs we tested the model lm (*Trait* ~ *Genotype*). Results from the statistical models are shown above each plot and, when significant, indicated by asterisks in the plot (where p-values < 0.001 and < 0.01 are denoted by '\*\*\*' and '\*\*', respectively). Differences for more than two groups were estimated by post hoc comparisons (Tukey's honest significant differences) and are indicated by different letters in each plot (p-value < 0.01). For all tested SNPs/genes, the phenotype of the DGRP lines carrying the minor versus the major allele at the target QTL can be found in S7 Fig.

https://doi.org/10.1371/journal.pgen.1007686.g004

candidate gene affects plasticity, we expected to see a difference in the slope of thermal reaction norms between genotypes with abolished (mutant) or reduced (knock-down) gene function in relation to the corresponding control genotypes (with "wildtype" gene function). To test specific significant SNPs/Indels, we used a SNP-based validation approach, hereafter called Mendelian Randomization (see Materials and methods), that allowed comparisons between genotypes fixed for the candidate SNP (minor versus major alleles) but not for any other significant SNP for the same trait. If a candidate SNP affects plasticity, we expected a difference in slope of the reaction norms between newly-established genotypes carrying the minor versus the major allele at the target SNP.

Using these methods, we confirmed a role in thermal plasticity for six out of seven candidate QTLs (Fig 4, S7 Fig, Table 2). For genes *Hsp60* (Fig 4A), *btv* (Fig 4B), *Men* (Fig 4E), and *Eip75B* (Fig 4F), plasticity was different between genotypes with impaired gene function (knock-out or knock-down) versus controls. For SNPs in genes *CG43902* (Fig 4C) and *ACC* (Fig 4D), plasticity was different between new genotypes with minor versus major allele. We did not find a difference in abdomen size plasticity between genotypes for candidate QTL *CG43117* (S7G Fig, Table 2, S2 Dataset). For all the confirmed candidates for plasticity, the genotypes with impaired gene function were more plastic than their respective controls, and the DGRP genotypes harboring the minor allele were more plastic than those harboring the major allele (S7 Fig). We also validated three out of four candidate QTLs for within-temperature variation in body size (Fig 4, S7 Fig, Table 2, S2 Dataset). Genes *Nmdmc* (Fig 4G) and *Optix* (Fig 4I) affected thorax size at 28°C and at 17°C, respectively, while a SNP in gene *CG14688* (Fig 4H) affected thorax size at 28°C. We did not find a difference in abdomen size at 28°C between genotypes for candidate QTL *Wapl* (S7K Fig, Table 2, S2 Dataset).

Table 1. Measurement summary and broad-sense heritability estimates for size and size plasticity.

	•		•				
	Trait 1		Range <sup>2</sup>	CV <sup>3</sup>	H <sup>2 4</sup>	Effects of G or GxT 5	
Within environment	Т	17°C	0.51-0.67	0.6-12.1	0.34	F = 6.6; df = 193,2104; p<2.2e-16	
		28°C	0.47-0.64	0.9-15.0	0.36	F = 7.4; df = 196,2271; p<2.2e-16	
	A	17°C	0.89-1.38	0.4-17.7	0.46	F = 10.9; df = 193,2148; p<2.2e-16	
		28°C	0.82-1.41	0.3-23.8	0.52	F = 14.3; df = 196,2391; p<2.2e-16	
Plasticity	Plasticity T RN		-1.3e-02-9.9e-03	NA	0.33	F = 5.2; df = 190,4375; p<2.2e-16	
	A	RN	-3.8e-02-3.1e-02		0.48	F = 10.3; df = 190,4539; p<2.2e-16	

<sup>&</sup>lt;sup>1</sup> Subsets used to perform the calculations: thorax (T) or abdomen (A) per temperature (17°C or 28°C) for within-environment parameters, and raw slope of the reaction norm (RN) for plasticity parameters.

https://doi.org/10.1371/journal.pgen.1007686.t001

<sup>&</sup>lt;sup>2</sup> Minimum and maximum size mean value (in mm) or minimum and maximum slope of the reaction norm.

<sup>&</sup>lt;sup>3</sup> Minimum and maximum coefficient of variation.

<sup>&</sup>lt;sup>4</sup> Broad-sense heritability estimates (see <u>S1 Table</u> for the variance components used to estimate heritability values).

<sup>&</sup>lt;sup>5</sup> Results of the statistical models used to test for the effect of genotype on size (model lm (*Size* ~ *Genotype*); per body part and temperature) or the interaction between genotype and temperature on size (model lm (*Size* ~ *Genotype*\* *Temperature*); per body part).



Table 2. Details of candidate QTLs selected for functional validation.

Gene ID 1	Polymorphism <sup>2</sup>	Position <sup>3</sup>	Type <sup>4</sup>	GWAS <sup>5</sup>	Validation <sup>6</sup>	CBP effect <sup>7</sup>
CG43902	SNP	X:10192303	mis	T—absolute RN	YES (MR)	NO
CR44894			int			
ACC	SNP	2R:7983239	int	T—raw RN, 17°C	YES (MR)	NO
CG43117	SNP	3R:26454823	mis	A—raw RN	NO (MR)	NA
CG14237			ups			
CG14238						
CG5447			down			
Hex-t1						
Hex-t2						
Pdf						
Men	8 SNPs, 1MNP	3R:12719133-3R:12726343	int, ups	A—1 for raw RN, 8 for abs RN	YES (KO)	NO
Hsp60	1 INS	X:11108613	5'UTR	T—absolute RN	YES (KO)	NO
			ups			
Eip75B	13 SNPs	3L:17997317-3L:18012599	5'UTR int, ups	A—absolute RN	YES (KD)	NO
btv	2 SNPs	2L:17961008-2L:17969392	down	T—raw RN	YES (KD)	YES
				T—absolute RN		
Nmdmc	2 SNPs,	3R:9042750	5' UTR, syn, int, ups	T—28°C	YES (KD)	NO
	1 DEL	- 3R:9052391		T—28°C		
CG14688	SNP	3R:10678848	mis	T—28°C	YES (MR)	NA
CG6465			ups			
CR44230						
Optix	SNP	2R:8032561	int	T—17°C	YES (KD)	NA
Wapl	SNP	X:2152985	int	A—28°C	NO (KD)	NA

<sup>&</sup>lt;sup>1</sup> Putatively affected gene given as ID gene symbol.

https://doi.org/10.1371/journal.pgen.1007686.t002

To explore the pleiotropic effect of validated plasticity QTLs, we investigated whether the plastic response was also seen in the body part for which the SNP/gene had not been significantly associated to in the GWAS analysis (S7 Fig, Table 2). Out of the six validated plasticity QTLs, we only found cross-body part effects for gene *btv*; initially implicated in variation in plasticity of thorax size, this gene was also found to influence variation in plasticity of abdomen size (S7B Fig, Table 2).

#### **Discussion**

Body size and body size proportions are traits closely associated to fitness [1, 2, 5], which vary greatly between species and populations, as well as between sexes and between same-sex individuals [11, 39, 62]. Environmental conditions, such as temperature or food availability, can work both as inter-generational selective agents impacting body size evolution, and as intragenerational instructive agents that affect body size during development [7, 9, 33, 63]. Studies

<sup>&</sup>lt;sup>2</sup> Type of polymorphism (SNP = Single Nucleotide Polymorphism, INS = Insertion, DEL = Deletion, MNP = Multiple Nucleotide Polymorphism).

<sup>&</sup>lt;sup>3</sup> Genomic position (FlyBase release FB2017\_06 [61]).

<sup>&</sup>lt;sup>4</sup> Putatively affected gene regions (mis = coding missense substitution, int = intronic, ups = upstream, down = dowstream, syn = coding synonymous substitution).

<sup>&</sup>lt;sup>5</sup> Body part (T = thorax, A = abdomen) and trait for which the QTL was a hit in the GWAS.

<sup>&</sup>lt;sup>6</sup> Results of the validation ("YES/NO" for whether validation confirmed or not the role of the target QTL on phenotype; results in Fig 4 and S7 Fig) including information of the approach used (MR = Mendelian Randomization, KO = mutant, KD = RNAi).

<sup>&</sup>lt;sup>7</sup> Results of the test for pleiotropic effects ("YES/NO" for whether we found cross body part (CBP) effects; results in S7 Fig).



in various species have explored what shapes inter- and intra-specific differences in size, including the physiological basis of body size regulation [16, 36, 38, 64] and the genetic basis of variation in body size [6, 65, 66].

## Components of variance in body size

Studying a panel of populations representing naturally-segregating alleles, the DGRP, we quantified effects of G, E, and GxE interactions on the size of two body parts (thorax and abdomen). As is well documented for various species of insects [32, 67, 68], most *D. melanogaster* genotypes we analyzed showed larger bodies when flies were reared at our lower temperature. However, we also documented cases of genotypes showing no plasticity (robustness) or showing size plasticity in the opposite direction. We also found a positive correlation in the levels of plasticity for the two body parts. The strong associations between the sizes of different body parts and plasticity therein is likely to reflect the tight regulation of body proportions, which is key for organismal performance [7, 69].

It is unclear to what extent a genotype's responsiveness to environmental conditions (i.e. its plasticity) is associated with inter-individual differences found within a given environment for that same genotype (quantified with the coefficient of variation). While the latter is presumably un-accountable for by the effects of G, E, or GxE, it could reflect small genetic differences between individuals (e.g. derived from somatic mutation), micro-environmental variation (e.g. differences within a vial), or stochasticity in phenotype expression (e.g. developmental noise). We found that genotypes that were more plastic for thorax size (but not abdomen) also had higher levels of intra-genotype, intra-environment variation (i.e. higher coefficient of variation). Whether this component of phenotypic variance is assignable to micro-environmental variation and whether it has its own genetic basis has started to be investigated [53, 70, 71] and will, undoubtedly, be a topic of targeted future research.

# QTL for size and size plasticity of different body parts

By using the raw and absolute values of the slopes of reaction norms as quantitative traits, we identified loci associated with variation in size plasticity. Genetic variation for environmental responsiveness could, in principle, involve different types of molecular players and could affect multiple traits in a similar or different manner. We described QTLs influencing size plasticity corresponding to different functions in terms of putative SNP effects (e.g. missense, regulatory, or synonymous mutations; Table 2, S3 Table), and described molecular function and biological process for corresponding genes. These genes could potentially be mediating environmental effects at different levels, from the perception of the environmental cue (e.g. gene *btv*, which has been implicated in sensory perception [72]), to the transmission of that information to developing tissues (e.g. gene *Eip75B*, coding for an ecdysone receptor [73]), or the execution of the information on those tissues (e.g. genes *Men*, *ACC*, and *Hsp60A*, coding for two metabolic enzymes and a chaperone [73], respectively).

Presumably, genes higher up in the process of responding to the environment (e.g. those involved in the perception of external conditions versus those responding in specific tissues) would be more likely to affect multiple plastic traits in a concerted manner. With the exception of *btv*, none of our validated plasticity QTLs affected plasticity for other than the body part they had been identified as QTL for. Even for the complete set of candidate loci, we found very little overlap between QTLs for plasticity of different body parts (thorax versus abdomen), as well as for different properties of the reaction norms (raw versus absolute value of slopes). Furthermore, we also documented mostly private QTLs influencing variation in size at any given environment (i.e. body part and temperature-specific). Sex-, body part-, and environment-



specific QTL effects had been previously documented for various traits in different models [51, 74–76]. In *D. melanogaster*, for example, different loci have been associated to variation in bristle number in different body parts [76], and to variation in size in different environments [77]. Such private QTLs can potentially facilitate independent evolution of the traits.

Previous works exploring the genetic basis of environmentally sensitive variation have mostly focused on investigating QTLs whose effect vary across environments (QTL-by-environment interactions) for a variety of traits in different species [66, 78-80]. Much less attention has been paid to unraveling the allelic variants contributing to differences in plasticity itself. Exceptions include mapping of the genetic basis of thermal plasticity in life-history traits in Caenorhabditis elegans [81], of photoperiodic plasticity in multiple traits in Arabidopsis thaliana [82] and, more recently, of thermal plasticity in cold tolerance in D. melanogaster [53]. Our results revealed little overlap between the QTLs that contribute to variation in trait within environments and the QTLs that contribute to variation in trait plasticity, assessed from the slope of reaction norms. We documented loci underlying variation in size plasticity (i.e. properties of reaction norms) that are different from those underlying variation in size at any temperature (i.e. at 17°C and at 28°C). These results shed light onto a long-standing discussion about the genetic underpinnings of plasticity, which argue that either the genetic control of phenotypic plasticity happens via specific loci determining plastic responses or via the same loci that control trait values at a given environment [83-86]. Our data show that the genetic basis for trait plasticity, to a large extent, differs from the genetic basis for phenotypic variation in the trait itself.

# Evolution of size and size plasticity

Plasticity can be adaptive in that it helps populations cope with environmental heterogeneity, and it has even been argued that it can promote phenotypic and taxonomic diversification [20–22, 27]. Theoretical models highlight the ecological conditions that should influence the evolution of plasticity, such as the predictability of environmental fluctuations [87] and costs of plasticity [88, 89]. Plasticity is generally presumed to be costly and only selected for in predictably heterogeneous environments, such as seasons [90]. The absence of a correlation between our thermal plasticity measurements and various fitness-related traits measured for the same genotypes could not identify any such cost. These potential costs might involve traits that have not been considered here, or these same traits but under (environmental) conditions that were not those assayed.

The ability to respond or resist environmental perturbation, and the balance between both processes, can be crucial for fitness in variable environments. In the DGRP, even though some degree of environmental responsiveness is maintained, we found that the alleles contributing to increased levels of plasticity occur nearly always at lower frequencies (i.e. the genotypes with the minor allelic variant having steeper reaction norms than those with the major allele). It is unclear to what extent this is the result of natural selection by the thermal regime that the natural population from which the DGRP was derived was exposed to, and/or the result of the process of deriving the mapping panel in the laboratory. It is also unclear to what extent QTLs for size plasticity in the DGRPs are those under selection in other populations. While it is conceivable, if not likely, that different QTLs contribute to variation in plasticity (or other quantitative traits) in different populations, we did find that a number of our plasticity QTLs have been targets of selection in other populations. Specifically, some of our candidate genes for size plasticity appear to have been selected in experimental populations of *D. melanogaster* evolving under different fluctuating thermal regimes [91]. Among our 192 candidate QTLs for thermal plasticity, eight genes (including the validated *btv*) had changes in the populations evolving



under hot and cold temperatures fluctuations [91], nine genes (including *Men*) had changes in populations evolving under hot fluctuations, and 25 genes had changes in the populations evolving under cold temperatures fluctuations (see all overlaps in S3 Table). Gene *Eip75B* has also been previously implicated in the response to artificial selection for body size [92] and in differentiation between clinal populations [93, 94], which typically represent different thermal environments.

Altogether, our results shed light onto the nature of inter-genotype variation in plasticity, necessary for the evolution of plasticity under heterogeneous environments. We showed that QTLs for size plasticity: 1) bear alleles for increased plasticity at low frequencies, 2) correspond to polymorphisms in different genomic regions and within genes of a multitude of functional classes, and 3) are mostly "private QTLs", with little overlap between our various GWAS analysis. The latter underscores the potential for independent evolution of trait and trait plasticity (different QTLs for size plasticity and for within-environment size variation), plasticity of different body parts (different QTLs for size plasticity of thorax and of abdomen), and even properties of the environmental response (different QTLs for raw and absolute slopes of thermal reaction norms).

#### Materials and methods

## Fly stocks and rearing conditions

Data for the GWAS was collected from adult female flies of the Drosophila Genetic Reference Panel (DGRP) obtained from the Bloomington Stock Center. The DGRP is a set of fully sequenced inbred lines collected from a single population in Raleigh, NC, USA [46, 48]. The number and the details of the lines included in the GWAS for each trait can be found in \$2 Table. Mutant stocks for the functional validations were: Hsp60A (stock 4689 from Bloomington), MenEx3 and MenEx55 (obtained from the T. Merritt lab). Control genetic backgrounds were w1118 (stock 5905, from Bloomington) and Canton-S (obtained from C. Mirth lab). UAS-Gal4 and UAS-RNAi lines used for validations were: stocks 6803 for bab-Gal4, 5138 for tub-Gal4, 28737 for btv-RNAi, and 35785 for mCherry-RNAi (all obtained from the Bloomington stock center), and stocks v108399 and v110813 for Eip75B and Optix-RNAi, respectively (obtained from the VDRC stock center).

Fly stocks were maintained in molasses food (45 gr. molasses, 75gr sugar, 70gr cornmeal, 20 gr. Yeast extract, 10 gr. Agar, 1100 ml water and 25 ml of Niapagin 10%) in incubators at 25°C, 12:12 light cycles and 65% humidity until used in this study. For the experiments, we performed over-night egg laying from ~20 females of each stock in vials with *ad libitum* molasses food. Eggs were then placed at either 17°C or 28°C throughout development. We controlled population density by keeping between 20 and 40 eggs per vial. We reared 200 DGRP lines and quantified thorax and abdomen size of 5 to 20 females per line, temperature and replicate. For 135 DGRP lines, we ran two replicates and for 33 lines we ran three replicates. The total number of flies used varied among DGRP lines due to differences in mortality at one or both of the temperatures. For some specimens, we could only quantify size of one body part if, for example, the individual was not properly positioned in the image or was damaged. Full details on the stocks used and the number of flies used per stock and temperature can be found in \$1 Dataset and \$2 Table. Rearing conditions for the validations of candidate QTLs were similar to those used for the DGRP lines.

## Phenotyping body size and plasticity

Adult female flies (8–10 days after eclosion) were placed in 2ml Eppendorf and killed in liquid nitrogen followed by manual shaking to remove wings, legs and bristles. Bodies were mounted



on Petri dishes with 3% Agarose, dorsal side up, and covered with water to avoid light reflections. Images containing 10 to 20 flies were collected with a LeicaDMLB2 stereoscope and a Nikon E400 color camera under controlled imaging conditions of light and white-balance. Images were later processed with a customized Mathematica macro to extract size measurements. For this purpose, we drew two transects per fly, one in the thorax and one in the abdomen, using body landmarks (as shown in S1A Fig). Size of each body part was initially quantified as the number of pixels in the transect and later converted to millimeters. For abdominal transects, when necessary, we performed an additional step that involved the removal of pixels corresponding to the membranous tissue that is sometimes visible between segments.

## Genome-Wide association study

For each body part (thorax and abdomen), we performed four independent Genome-Wide Analyses (GWAS): two for thermal plasticity (raw and absolute values of the slopes of the reaction norms), and two for within-environment variation (length at 17°C and length at 28°C). Slopes of the reaction norm were calculated as the slope of the linear model lm (Size ~ Temperature) for each body part and DGRP line. We used linear mixed-effects models (lmer) in lme4 R package to test which polymorphisms explained variation in size plasticity or in size. The GWAS analyses for variation in thermal plasticity tested for effects of fixed factor Alelle (corresponding to each polymorphic site), and random factors Wolb (corresponding to Wolbachia status (0 or 1) of each DGRP line [7,8]) and DGRP (i.e. genetic background) to variation in the dependent variable Slope (i.e. raw or absolute values of the reaction norms). This corresponds to notation lmer (Slope ~ Allele + (1|Wolb/DGRP)) in lme4 R package [95]. The GWAS analyses for within-environment variation tested the model lmer (Size ~ Allele + (1|Wolb/DGRP), where Size is the dependent variable (thorax or abdomen length from flies reared at 17°C or at 28°C) and all the other terms are the same as described above for the plasticity GWAS. All the GWAS were performed by using polymorphisms where we had information for at least ten DGRP lines per allele. We did not find an effect of Wolbachia in any of our GWAS analyses.

For each of the GWAS, we annotated the SNPs with an arbitrary and commonly used p-value < 10e-5 using the FlyBase annotation (FlyBase release FB2017\_05; [73]). For the same SNPs, we performed first, a gene ontology enrichment analysis using the publicly available GOrilla Software [96, 97] and second, a network enrichment analysis using gene-enrichment and pathway-enrichment analyses were done using the publicly available NetworkAnalyst Software [98, 99]; using all nodes from first order network generated with IrefIndex Interactome settings.

Analyses of the overlap in significant QTLs (p-value < 10e-5) from different GWAS analyses was done for individual polymorphic sites (considered to be the same based on genomic position and type) and for the genes those sites were annotated to (using FlyBase release FB2017\_05; [73]).

We tested for the effect of the chromosomal inversions (In\_3R\_K, In\_3R\_P, In\_2L\_t, In\_2R\_NS and In\_3R\_Mo) on our thorax and abdomen traits by using linear models (Im (*Mean Size* ~ *Inversion*) for within-environment size variation and Im (*Slope* ~ *Inversion*) for size plasticity variation).

Genetic distance matrix for the DGRPs was obtained from http://dgrp2.gnets.ncsu.edu/data.html and was used to perform a cluster hierarchical dendogram using *ape* and *phylobase* R packages. We estimated the phylogenetic signal and statistical significance for each of our traits using Blomberg's K [100] and Pagel's  $\lambda$  [101] metrics with the *phylosig* function in the *phytools* R package [102].



#### **Functional validations**

The subsample of significant QTLs to be validated was taken from a first list of candidates (\$3 and \$4 Figs) selected based on p-value and corresponding peaks in the Manhattan plots (clear peaks prioritized), putative effect (missense and regulatory variants prioritized over intergenic variants), associated genes (annotated and known function prioritized). We used three methods for validation, depending on QTL properties: null mutants and RNAi (Gal4/UAS system) for genes containing several significant SNPs and/or containing SNPs corresponding to missense variants, and Mendelian Randomization (MR) for SNPs in genes with little or no information available. Mutant and RNAi test that no or low levels of peptide affect variation in the quantitative trait for which the gene was identified as a candidate QTL while MR tests for sufficiency and independence from genetic background of the specific allele. Following these criteria we tested a total of 11 candidate SNPs/genes.

Validations by null mutants were done by comparing the phenotype in the heterozygous mutant stock with its respective genetic background. Validations by RNAi were done by comparing, for each Gal4 driver line, the phenotype of the gene of interest knockdown with the corresponding control cross using UAS-*mCherry*RNAi as well as with the corresponding control genetic background for the UAS line. We always used two different driver lines for our validations by RNAi: *tub*-Gal4 and *bab*-Gal4. Gene *tub* is ubiquitously expressed throughout development and gene *bab* is expressed in multiple tissues during different stages of development. Both genes include expression in developing thoraxes and abdomens (see FlyBase reports FBgn0003884 and FBgn0004870, respectively). However, for all candidate genes selected for RNAi validation, except *Nmdmc*, the crosses between RNAi line and *tub*-Gal4 were lethal.

The identity of the SNPs tested by MR is given by their annotation with Genome Release v6. For each candidate SNP, we first selected 10 DGRP lines to make a population with the minor allele fixed and 10 others to make a population with the major allele fixed. The 10 DGRP lines used to create each population were checked for having only one of the significant QTLs (p-value < 10e-5) fixed and not the others. These lines were used to generate four populations, two fixed for the major allele and two for the minor allele (S2 Table). Each population was established by crossing 8 virgin females from each of 5 of the same-allele lines to 8 males of the other 5 lines. Reciprocal crosses were used to set two independent populations per allele. These populations were allowed to cross for eight generations to randomize genetic backgrounds. We confirmed by Sanger sequencing that those populations had our candidate allele fixed. Primer sequences used to confirm the allele in each population were:

Gene *CG43902*—forward primer: ACCACCAACATCAGCGTTTC; reverse primer: TGGTTT CGGCGTAGTTGTTG.

Gene *ACC*–forward primer: TGGGAAAAACCGGCCTAAGA; reverse primer: ATTTGTGGC TGTGGATTGCG.

Gene CG43117 – forward primer: TAAGCAAAATGTGGCGTGCA; reverse primer: TTAACA TGGATCCTGCGCAC

Gene CG14688 – forward primer: CATACTTTGACAGACGGCCG; reverse primer: CGGCTA CATTGTCATCGAGG

#### Statistical analyses

All statistical analyses were performed with R Statistical Package version 3.3.1 [103]. For each body part and temperature, we used linear mixed-effects model (lmer) in *lme4* R package [95]



to test for the effect of replicate on size (model lmer ( $Size \sim Replicate + (1|DGRP/Replicate)$ )), that were found to be non-significant (p-value > 0.05). Individuals from different replicates were pooled for all other analyses. We then used linear models (lm) to test for the within-environment effect of genotype (DGRP) on size (model lm ( $Size \sim DGRP$ )). We also used linear models to test for plasticity in size (i.e. the interaction between genotype (DGRP) and temperature (model lm ( $Size \sim DGRP^*Temperature$ ))). Reaction norms for each DGRP line were calculated by using the linear model lm ( $Size \sim Temperature$ ). We extracted two properties of the reaction norms per DGRP line and body part: the absolute value of the slope as a measurement of thermal sensitivity, describing only the magnitude of the response to temperature, and the raw value of the slope as a measurement which describes also the direction of that response. We defined plastic genotypes as those DGRP lines whose reaction norm slope was significantly different from zero (p-value < 0.05).

We used the total number of phenotyped individuals for each DGRP and temperature to perform the calculations of the summary statistics in S2 Table. We used Pearson correlations (alpha = 0.99) to test for linear correlation in size between body parts, controlling for DGRP lines. We also used Pearson correlations to test for linear correlations among our measured traits and between those and other available datasets for the DGRPs. For this, we used the mean value per DGRP line for each trait and the *corrplot* R package [104]. We report both correlation coefficient and significance levels. Available DGRP phenotypes that were used to correlate with our traits were: size measurements at 25°C [51], longevity [49], starvation resistance, chill coma recovery [46], tolerance to infection with *Providencia rettgeri* bacteria [56] and resistance to infection with *Metarhizium anisopliae* fungi or with *Pseudomonas aeruginosa* bacteria [50].

Broad sense heritability for size at each temperature was estimated as  $H^2 = \sigma_A^2/(\sigma_A^2 + \sigma_W^2)$  where  $\sigma_A^2$  and  $\sigma_W^2$  are the among-line and within-line variance components, respectively. Heritability of plasticity was calculated as  $H^2 = \sigma_{G^*E}^2/\sigma_{TOTAL}^2$  where  $\sigma_{G^*E}^2$  and  $\sigma_{TOTAL}^2$  are the variance associated with the genotype by environment interaction and total variance components, respectively, as proposed in Scheider and Lyman (1989). Variance components were extracted using *varcomp* R package.

For the functional validations of within-environment SNPs and genes we used the linear models  $\operatorname{lm}(Size \sim Allele)$  and  $\operatorname{lm}(Size \sim Genotype)$ , respectively. For the validations of plasticity SNPs and genes we used the models  $\operatorname{lm}(Size \sim Genotype^*Temperature)$  and  $\operatorname{lm}(Size \sim Allele^*Temperature)$ , respectively. In all cases, significant differences among groups were estimated by post hoc comparisons (Tukey's honest significant differences).

### Supporting information

**S1 Fig. Variation in size plasticity in the DGRP lines. A.** Image of an adult female *D. melanogaster* fly showing the thoracic and abdominal transects. **B.** Slope and confidence interval of the reaction norms in the DGRP lines, calculated as the regression model lm (*Size* ~ *Temperature*) in the thoraxes (grey) and abdomens (green) of each DGRP line (Y axis). Slopes are ranked by their value in the thorax. Horizontal bars represent the mean of all DGRP lines for the raw slope of the reaction norm (dashed bar) and the absolute slope of the reaction norm (solid bar) per body part. **C.** Histograms showing the frequency of the size measurements in thoraxes and abdomens of all DGRP lines reared at 17°C (blue) and 28°C (red). Dashed line represents the mean value for all DGRP lines at a given temperature. **D.** Histograms for the absolute slope of the reaction norms (calculated as the absolute value for the slope of the regression lm (*Size* ~ *Temperature*) in thoraxes and abdomens. Mean value for the absolute slope of all DGRP lines is indicated with a green arrowhead. (PDF)



**A.** Heat map of Pearson's correlation coefficients between our within-environment size measurements ( $17^{\circ}$ C and at  $28^{\circ}$ C) and size measurements at  $25^{\circ}$ C. Non-significant correlations (p-value > 0.01) are indicated with an 'X'. **B.** Heat map of Pearson's correlation coefficients between our traits (mean size at each temperature and raw and absolute slopes of the reaction norms) and fitness-related traits. Non-significant correlations (p-value > 0.01) are indicated with an 'X'. **C.** Dendogram of the genetic distance between DGRP lines. Corresponding trait values are shown as a heat map and scaled for each trait independently. Coefficients of Blomberg's K phylogenetic signal were: K = 0.24; p-value = 0.17 (thorax at  $17^{\circ}$ C), K = 0.23; p-

S2 Fig. Phenotypic co-variation in size and size plasticity with fitness and genetic distance.

value = 0.37 (thorax at 28 °C), K = 0.22; p-value = 0.78 (thorax raw slope), K = 0.24; p-value = 0.20 (thorax absolute slope), K = 0.24; p-value = 0.24 (abdomen at 17 °C), K = 0.22; p-value = 0.67 (abdomen at 28 °C), K = 0.23; p-value = 0.55 (abdomen raw slope) and K = 0.21; p-value = 0.89 (abdomen absolute slope). Pagel's  $\lambda$  coefficient of phylogenetic signal was  $\lambda$  = 6.88e-05; p-value = 1, for all the traits in both body parts.

(PDF)

S3 Fig. GWAS for variation in size plasticity per chromosomal arm. Manhattan plots corresponding to the four GWAS performed for variation in size plasticity: raw slopes of the reaction norms (grey dots) and absolute slope of the reaction norms (black dots) for thorax (left side) and abdomen (right side) size. For each trait and body part, the GWAS was done testing the model  $\lim (Slope \sim Allele + (1|Wolb/DGRP))$ . The significance level for each SNP along the chromosomal arms is shown as the  $\log_{10}$  p-value. Some of the genes associated to SNPs/Indels with a p-value < 10e-5 and that we consider as particularly interesting are shown. The position and identity of the polymorphisms in this figure is given by their annotation with Genome Release v.5. (PDF)

S4 Fig. GWAS for variation in size per chromosomal arm. Manhattan plots corresponding to the four GWAS performed for within-environment variation in size: at  $17^{\circ}$ C (blue dots) and at  $28^{\circ}$ C (red dots) for thorax (left side) and abdomen (right side) size. For each trait and body part, the GWAS was done testing the model  $\lim (Size \sim Allele + (1|Wolb/DGRP))$ . The significance level for each SNP along the chromosomal arms is shown as the  $\log_{10}$  p-value. Some of the genes associated to SNPs/Indels with a p-value < 10e-5 and that we consider as particularly interesting are shown. The position and identity of the polymorphisms in this figure is given by their annotation with Genome Release v.5. (PDF)

S5 Fig. Effect and frequency of candidate SNPs/Indels for variation in size plasticity. A. Mean and confidence interval of the absolute slope of the reaction norms for abdomen size (Y axis) per allele (minor allele in grey, minor allele in magenta) at each candidate plasticity SNP/Indel (p-value < 10e-5) along the chromosomal arms (X axis). **B.** Mean and confidence interval of the raw slope of the reaction norms for size (Y axis) per allele (minor in grey, minor in magenta) at each candidate plasticity QTL (p-value < 10e-5) along the chromosomal arms (X axis) per body part. The position and identity of the polymorphisms in this figure is given by their annotation with Genome Release v.5. (PDF)

S6 Fig. Functional relationships between candidate genes influencing size and size plasticity. A-B. Gene ontology and network enrichment analyses for genes harboring allelic variants significantly associated with variation in size plasticity. All SNPs/Indels with a p-value < 10e-5 from the GWAS for the raw and absolute slopes of the reaction norms of thoraxes and abdomen



were pooled to perform these analyses. **A.** Results of the gene-ontology enrichment analysis for genes harboring allelic variants significantly associated with variation in size reaction norms. **B.** Network enrichment analyses (KEGG gene enrichment analyses of the protein-protein interactions) for genes harboring allelic variants significantly associated with variation in size reaction norms. All our candidate QTLs for variation in size plasticity were pooled for this analysis. This corresponds to the QTLs from four independent GWAS: raw and absolute values of the slopes of the reaction norms for thoracic and abdominal size. **C.** Network enrichment analyses (KEGG gene enrichment analyses of the protein-protein interactions) for genes harboring allelic variants significantly associated with variation in size reaction norms. SNPs with p-value < 10e-5 from the GWAS for variation at 17°C and for variation at 28°C were pooled to perform this analysis. Note that for allelic variants associated with variation in size plasticity, the gene-ontology enrichment analyses showed no enriched GO terms. All our candidate QTLs for variation in size were pooled for this analysis. This corresponds to the QTLs from four independent GWAS: thoracic and abdominal size for flies reared at 17°C and at 28°C. (PDF)

S7 Fig. Allele effect in the DGRP and pleiotropy effect for the QTLs selected for validation. For each candidate QTLs, the effect of the minor and major alleles in the DGRPs are shown in the left panels and the pleitropic effect is shown in the right panels. For validations via mutant or RNAi, genotypes with impaired gene function are shown in magenta and control genotypes are shown in black. Similarly, for validations via Mendelian Randomization (MR), the two populations fixed for the minor allele are shown in magenta and the two populations for the major allele are shown in black. The identity of the SNP/Indels is given by their annotation with Genome Release v.6. as Chromosome:Position. Size measurements are given as length in mm. A. Left panel: slope of the reaction norms in the DGRP lines with the major and the minor alleles for insertion in position X:11108613, within gene Hsp60. Right panel: reaction norms in mutant Hsp60A/+ and controls Canton-S (filled circles, solid line) and Fm7a/Canton-S (empty circles, dashed line). B. Left panel: slope of the reaction norms in the DGRP lines with the major and the minor alleles for SNP in position 2L:17961008, within gene btv. Right panel: reaction norms for size in btv-RNAi/bab-Gal4 and control lines KK (filled circles, solid line) and mCherry-RNAi/bab-Gal4 (empty circles, dashed line). C. Left panel: slope of the reaction norms in the DGRP lines with the major and the minor alleles for SNP in position X:10192303, within gene CG43902. Right panel: reaction norms in the four MR populations corresponding to SNP in position X:10192303, within gene CG43902. D. Left panel: slope of the reaction norms in the DGRP lines with the major and the minor alleles for SNP in position 2R:7983239, within gene ACC. Right panel: reaction norms in the four MR populations for SNP in position 2R:7983239 within gene ACC. E. Left panel: slope of the reaction norms in the DGRP lines with the major and the minor alleles for SNP in position 3R:12720159 within gene Men. Right panel: reaction norms in Men mutants MenEx3/+ (filled circles, solid line) and MenEx55/+ (empty circles, dashed line) and control line w1118. F. Left panel: slope of the reaction norms in the DGRP lines with the major and the minor alleles for SNP in position 3L:17999272, within gene Eip75B. Right panel: reaction norms in Eip75B-RNAi/bab-Gal4 and control lines KK (filled circles, solid line) and mCherry-RNAi/bab-Gal4 (empty circles, dashed line). G. Left panel: slope of the reaction norm in the DGRP lines with the major and the minor alleles for SNP in position 3R:26454823, within gene CG43117. Right panel: reaction norms in the four MR populations for SNP 3R:26454823 within gene CG43117. H. Left panel: size at 28°C in the DGRP lines with the major and the minor alleles for SNP in position 3R:9049187, within gene Nmdmc. Right panel: size at 28°C in Nmdmc-RNAi/tub-Gal4 and control lines GD (filled circles) and mCherry-RNAi/tub-Gal4 (filled circles). I. Left panel: size at



28°C in the DGRP lines with the major and the minor alleles for SNP in position 3R:10678848, within gene CG14688. Right panel: size at 28°C in the four MR populations for SNP in position 3R:10678848, within gene CG14688. J. Left panel: size at  $17^{\circ}$ C in the DGRP lines with the major and the minor alleles for SNP in position 2R:8032561, within gene Optix. Right panel: size at 17°C in Optix-RNAi/bab-Gal4 and control lines KK (filled circles) and mCherry-RNAi/ bab-Gal4 (filled circles). K. Left panel: size at 28°C in the DGRP lines with the major and the minor alleles for SNP in position X:2152985, within gene Wapl. Right panel: size at 28°C in Wapl-RNAi/bab-Gal4 and control lines KK (filled circles, solid line) and mCherry-RNAi/bab-Gal4 (empty circles, dashed line). For the validations of plasticity QTLs (genes Hsp60, CG43902, Men, ACC, btv, Eip75, and CG43117), we tested the model lm (Trait ~ Genotype\* Temperature). For the validations of within-environment QTLs (genes Nmdmc, CG14688, Optix and Wapl), we tested the model lm (Trait ~ Genotype). Results from the statistical models are shown above each plot and, when significant, indicated by asterisks in the plot (where p-values < 0.001 and < 0.01 are denoted by '\*\*\*' and '\*\*', respectively). Differences for more than two groups were estimated by post hoc comparisons (Tukey's honest significant differences) and are indicated by different letters in each plot (p-value < 0.01). (PDF)

S1 Table. Variance components.

(DOCX)

S2 Table. Summary of phenotypic measurements for size and size plasticity in the DGRP. (DOCX)

S3 Table. GWAS results for variation in size plasticity. (DOCX)

**S4** Table. GWAS results for variation in size. (DOCX)

**S1 Dataset. Phenotypic variation in size in the DGRP.** Raw individual data for size measurements (length in mm) in thoraxes and abdomens of females adult flies from the DGRP lines reared at 17°C and at 28°C. (XLSX)

**S2 Dataset. Functional validations of selected GWAS candidates.** Raw data for size measurements (length in mm) in thoraxes and abdomens of flies reared at 17 °C and at 28 °C from the different genetic backgrounds used for functional validations. (XLSX)

# **Acknowledgments**

We are grateful to Thomas Merritt for sharing the *Men* mutants, and to Daniel Sobral and IGC's Bioinformatics Unit and Fly Unit for technical support. We also thank Erik van Bergen for comments on the manuscript. Computations were performed on the EDB-Calc Cluster, which uses a software developed by the Rocks(r) Cluster Group (San Diego Supercomputer Center, University of California, San Diego and its contributors), and is hosted by the laboratory "Evolution et Diversité Biologique" (EDB) and supported by Pierre Solbes.

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