

1 **Spatiotemporal patterns of desiccation tolerance in natural populations of *Drosophila***  
2 ***melanogaster***

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12 *Running title:* Drought tolerance in temperate *Drosophila*

13 *Keywords:* Desiccation tolerance, *Drosophila*, Geographical variation, North America.

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18 **Abstract** Water availability is a major environmental challenge to a variety of terrestrial  
19 organisms. In insects, desiccation tolerance varies predictably over various spatial and temporal  
20 scales and is an important physiological basis of fitness variation among natural populations.  
21 Here, we examine the dynamics of desiccation tolerance in North American populations of  
22 *Drosophila melanogaster* using: 1) natural populations sampled across latitudes and seasons in  
23 the eastern USA; 2) experimental evolution in the field in response to changing seasonal  
24 environments; 3) a sequenced panel of inbred lines (DGRP) to perform genome wide  
25 associations and examine whether SNPs/genes associated with variation in desiccation tolerance  
26 exhibit patterns of clinal and/or seasonal enrichment in pooled sequencing of populations. In  
27 natural populations we observed a shallow cline in desiccation tolerance, for which tolerance  
28 exhibited a positive association with latitude; the steepness of this cline increased with  
29 decreasing culture temperature, demonstrating a significant degree of thermal plasticity. No  
30 differences in desiccation tolerance were observed between spring and autumn collections from  
31 three mid-to-northern latitude populations, or as a function of experimental evolution to  
32 seasonality. Similarly, water loss rates did not vary significantly among latitudinal, seasonal or  
33 experimental evolution populations. However, changes in metabolic rates during prolonged  
34 exposure to dry conditions indicate increased tolerance in higher latitude populations. Genome  
35 wide association studies identified thirty-six SNPs in twenty-eight genes associated with sex-  
36 averaged drought tolerance. Among North American populations, genes associated with drought  
37 tolerance do not show increased signatures of spatially varying selection relative to the rest of the  
38 genome, whereas among Australian populations they do.

39

40

## 41 **Introduction**

42           Insects exploit and inhabit a wide range of habitats on planet earth which range from hot  
43 deserts to cold arctic regions. In many of these environments presence of water is scarce and  
44 desiccation is a major threat to many terrestrial organisms living there. Insects are particularly  
45 most vulnerable to water related challenges, because of their small size and thus large surface  
46 area to volume ratio. (Gibbs & Rajpurohit 2010). Environmental stresses such as desiccation are  
47 highly variable among these natural habitats, and often vary predictably with such features as  
48 latitude, altitude, and season. Patterns of phenotypic and genetic variation distributed along these  
49 gradients, both within and among taxa, offer means to address the extent to which divergence in  
50 physiologically selected traits are affected by natural selection (Barton 1999; Whitlock &  
51 McCauley 1999).

52           *Drosophila* species constitute good models for studies relating to population ecology and  
53 physiological adaptations (Parsons 1983; Lemeunier et al. 1986). During their evolutionary  
54 history, different *Drosophila* species have adapted to diverse climatic conditions. This has been  
55 clearly demonstrated on multiple continents where multiple *Drosophila* species exhibit  
56 pronounced clines for multiple fitness traits (e.g., Hoffmann & Parsons 1991; Hoffmann &  
57 Parsons 1993; Blows & Hoffmann 1993; Karan et al. 1998; Gilchrist et al. 2004; Schmidt et al.  
58 2005; Wittkopp et al. 2011; Rajpurohit & Nedved 2013). In various taxa, high levels of  
59 desiccation resistance are associated with adaptation to arid habitats (e.g., David et al. 1983;  
60 Hoffmann and Parsons 1991; Gibbs et al. 2003). Several studies have reported geographical  
61 variation for desiccation tolerance in *Drosophila* species inhabiting arid environments

62 characterized by elevated temperatures (e.g., Hoffmann & Harshman 1999; Matzkin et al. 2007;  
63 Rajpurohit & Nedved 2013; Rajpurohit et al. 2013b).

64 In the cosmopolitan *D. melanogaster*, latitudinal clines have been observed for a wide  
65 range of fitness-associated traits including thermal tolerance, body size, basic life histories,  
66 incidence of reproductive dormancy, cuticular hydrocarbon composition, and aspects of behavior  
67 (e.g., David 1975; Coyne and Beecham 1987; James et al. 1995, 1997; Zwaan et al. 2000;  
68 Schmidt et al. 2005; De Moed et al. 1997; Mitrovski and Hoffmann 2001; Karan et al. 1998;  
69 Eanes 1999; Verrelli and Eanes 2001; Lemeunier et al. 1986; Rajpurohit & Schmidt 2016).  
70 While such patterns may be influenced by demography (Kao et al. 2015; Bergland et al. 2016),  
71 latitudinal clines in *D. melanogaster* are commonly interpreted as resulting from spatially  
72 varying selection and local adaptation to climatic and associated variables.

73 However, the role of desiccation tolerance in the adaptation of *D. melanogaster* to  
74 environmental heterogeneity has not been comprehensively addressed (Hoffmann et al. 2001;  
75 Hoffman et al. 2005; Telonis-Scott et al. 2006; Parkash et al. 2008); there has been little work  
76 done on *D. melanogaster* populations inhabiting temperate latitudinal ranges, in which humidity  
77 and associated desiccation stress are predicted to vary both with latitude and season. Similarly,  
78 the genetic basis of variation in desiccation tolerance remains unresolved (but see Telonis-Scott  
79 et al. 2016). We therefore carried out an integrated study on desiccation tolerance in natural and  
80 experimental populations in a genetic model organism, utilizing a combination of phenotypic,  
81 physiological and plasticity analyses, population level sequencing and genome wide association  
82 studies. Our results demonstrate thermally mediated plasticity, associations among latitude,

83 physiology and stress tolerance, and identify a set of candidate SNPs that may underlie variation  
84 in desiccation tolerance in natural populations.

85

## 86 **Material & Methods**

### 87 *Fly collection and maintenance*

88 Six natural populations of *D. melanogaster* were collected from fruit orchards located  
89 along the east coast of the United States by a combination of aspiration and baiting/sweeping  
90 with aerial nets (see Fig. 1). Gravid females were immediately sorted into isofemale lines in the  
91 field; once the resulting progeny eclosed, lines were typed to species. Approximately 150  
92 isofemale *D. melanogaster* lines were collected from each locale. Populations sampled were:  
93 Bowdoin, Maine (44.03N, 73.20W); Media, Pennsylvania (40.04N, 76.30W); Charlottesville,  
94 Virginia (38.03N, 78.48W); Athens, Georgia (32.84N, 83.66W); Jacksonville, Florida (30.33N,  
95 81.66W); and Homestead, Florida (25.46N, 80.45W). Seasonal collections were done in Media,  
96 Pennsylvania, Lancaster, Massachusetts (42.455N, 71.67W) and Charlottesville, Virginia  
97 (38.03N, 78.48W) orchards in June and October of 2012. Long-term maintenance of all  
98 populations was done at 24 °C, 12:12 light:dark photoperiod, with a generation time of  
99 approximately 21 days. Flies were maintained on regular cornmeal-molasses-agar media.

### 100 *Establishment of clinal cages*

101 To establish population cages for each spatial and temporal collection, we pooled 25  
102 isofemale lines. We created two population cages for each of the spatial and temporal collections

103 (see above). Each cage was created using independent sets of 25 isofemale lines by releasing ten  
104 mated females from each line into 12x12x12 inch insect enclosures (Live Monarch Foundation,  
105 Boca Raton, Florida, USA). These lines were maintained in mass culture and allowed to outcross  
106 for 5 generations; subsequently, samples were collected for the phenotypic assays described  
107 below.

### 108 *Desiccation tolerance assay*

109 Four-to-five day old virgin flies were transferred to empty vials in groups of ten, and  
110 restricted to the lower half of the vials by a foam stopper. Silica gel was then added above the  
111 stopper to maintain low humidity, and the vial was sealed with Parafilm<sup>TM</sup>. Mortality was  
112 recorded at hourly intervals until all flies were dead. Fifteen to twenty-five isofemale lines per  
113 population were assayed. Throughout the desiccation tolerance assay vials were kept in a 25 °C  
114 incubator.

### 115 *Thermal plasticity and diapause treatment*

116 To examine thermal plasticity for desiccation tolerance in the latitudinal collections, we  
117 examined tolerance as a function of 1) three culture temperatures (18, 25 and 28 °C), and 2) after  
118 exposure to conditions that elicit reproductive dormancy (3-week exposure to 11 °C, 10L:14D  
119 photoperiod). Each replicate population cage for each of the geographic regions was allowed to  
120 oviposit for a 2-3h at 25 °C in successive culture vials; egg density was standardized at 40-50  
121 eggs per vial by manual removal. Replicate vials were then randomly assigned to one of the four  
122 experimental treatments and cultured in Percival I36VL incubators. On emergence, adults were  
123 transferred to fresh food vials and aged for 4-5 days before subjecting them to the desiccation

124 tolerance assay; for the reproductive dormancy assay, flies were immediately evaluated for  
125 desiccation tolerance following the 3-week period of exposure.

### 126 *Experimental manipulation of desiccation tolerance evolution*

127 To examine how desiccation tolerance evolves under field conditions, we established 20  
128 experimental mesocosms at a field site in Philadelphia, PA USA. Each mesocosm was an 8m<sup>3</sup>  
129 outdoor insect rearing enclosure (Bioquip Products, Gardena, CA) surrounding a mature (dwarf)  
130 peach tree. Each mesocosm was seeded with 1000 individuals (500 males, 500 females) derived  
131 from a collection made in 2012 from the same PA orchard as described above. This progenitor  
132 population was created by pooling 86 independent inbred lines, allowing them to recombine and  
133 expand for 10 generations in the laboratory, and then maintained at large census size (~10<sup>6</sup>) in  
134 the laboratory for the duration of the experiment (July 1 – November 1, 2014). Each cage was  
135 randomly assigned to a treatment: seasonally evolving (E) or non-seasonally evolving (F). In the  
136 E treatment, populations were supplied with fresh food/oviposition sources (500ml standard  
137 cornmeal-molasses medium) every 2d, and the populations were allowed to evolve and adapt to  
138 seasonal environmental conditions. In the F treatment, fresh food/oviposition sources were also  
139 supplied: however, all eggs laid by the experimental flies were counted, removed, and replaced  
140 with the same number of eggs from the progenitor laboratory population that was maintained  
141 under aseasonal, laboratory conditions. Thus, the F populations were maintained under normal  
142 demographic trajectories and flies were directly exposed to the field environment, but these  
143 populations were not allowed to evolve to the field conditions; the F treatment represents the  
144 progenitor laboratory population while including any potential epigenetic marks that may be  
145 elicited upon exposure to field conditions during direct development as well as any evolution in

146 the aseasonal, laboratory environment. At the end of the experiment, a sample of approximately  
147 2000 eggs was collected from each of the 20 cages and brought back to the laboratory. These  
148 collections were allowed to develop in the laboratory and subsequently passed through two  
149 generations of common garden, density controlled culture. In the F3 generation subsequent to  
150 the field collection, experimental animals were collected and processed for respirometry  
151 measurements.

### 152 *Respirometry measurements*

153         Within 3 hours of adult eclosion, virgin males and females were collected, and sets of  
154 fifteen were placed in fresh food vials. A total of 24 sets were assayed for the latitudinal  
155 extremes of Maine and Florida (2 geographic regions  $\times$  2 sexes  $\times$  2 replicate populations  $\times$  3  
156 experimental replicates), 24 for seasonal comparisons in the Pennsylvania orchard (2 seasons  $\times$  2  
157 sexes  $\times$  2 replicate cages  $\times$  3 repeats) and 40 for experimental evolution (2 treatments  $\times$  2 sexes  
158  $\times$  10 replicate cages). Respirometry was carried out at 4-6 days post-eclosion on sets of 10-15  
159 individuals. Flies were transferred directly from their food vials to a 4 mL glass metabolic  
160 chamber with aluminum stoppers, which was covered with a black cardboard sleeve to reduce  
161 activity in the chamber. Flow through respirometry at 25°C was carried out using two channels  
162 of a flow multiplexer (RM-8, Sable Systems International, Las Vegas, NV, USA), where dry,  
163 CO<sub>2</sub>-free air was supplied to the chambers at 50 mL·min<sup>-1</sup> using factory-calibrated mass flow  
164 controllers (MC-500 sccm; Alicat Scientific, Tuscon, AZ, USA), and excurrent air from the  
165 measured chamber was passed through a LI-7000 CO<sub>2</sub>/water vapor dual analyzer (Li-Cor  
166 Biosciences, Lincoln NE, USA). The flies were acclimated to the experimental chambers and air  
167 flow for 15 minutes during which flies from the alternative chamber were measured. Identical

168 empty chambers were used for baselining. Analyzer voltage output was recorded, stored and  
169 analyzed using UI-2 data acquisition interphase and Expedata software (Sable Systems  
170 International). Recording rate was set to 1 Hz and only data from the last 5 minutes of each 15  
171 min run were averaged for analysis.

172 We used an additional experimental approach to compare the Florida and Maine  
173 populations for which significant differences in desiccation resistance were found (see results). It  
174 was previously reported that water loss rates of *D. melanogaster* under similar experimental  
175 conditions stabilize only after >2h of exposure (Gibbs et al. 1997), and therefore we carried out  
176 respirometry on additional sets of flies (2 geographic regions  $\times$  2 sexes  $\times$  2 replicate populations  
177  $\times$  6 repeats), which were randomly allocated to six multiplexer channels, with a seventh used for  
178 baselining. The measurement sequence of 20 min measurements was as follows: baselining, 3  
179 experimental chambers, baselining, three additional experimental chambers and finally  
180 baselining again for a total file recording of 3 h, which was immediately followed by another  
181 recording at the same sequence. As each set of flies was added to the respirometry setup 20 min  
182 prior to the initial measurement, and only data from the last 10 minutes were analyzed, the flies  
183 were assayed 30 min and following additional 3h during which the flies were exposed to  
184 experimental temperature and dry air flow conditions.

#### 185 ***Desiccation assays for the Drosophila Genetic Reference Panel (DGRP)***

186 DGRP lines (Mackay et al. 2012) were obtained from the Bloomington Drosophila Stock  
187 Center and maintained in the lab on yeast-cornmeal-sucrose medium at 25°C. For assays, 4-6  
188 day old flies were sorted by sex and held on fresh media for two days. On the days experiments  
189 began, flies were lightly anesthetized with CO<sub>2</sub> and ten groups of five flies each were rapidly

190 transferred to empty shell vials. A foam plug was inserted halfway down the vial, silica gel  
191 desiccant was added to the upper half of the vial, and the vial was sealed with Parafilm<sup>TM</sup>. The  
192 vials were transferred to a 25°C incubator with constant illumination, and dead flies were  
193 counted hourly until all flies were dead.

194 Desiccation assays were performed in blocks of ~30 lines per block. Because of the time  
195 required to initiate desiccation stress for all lines and to count dead flies, we recorded the exact  
196 time when desiccant was added to assay vials and the exact times when each line was checked.  
197 Flies that died before the first survival check were assumed to have been injured by handling or  
198 other stress and were not included in the data analysis. Any flies dying between two checks were  
199 assumed to have died midway between them. To assess potential variation among blocks  
200 associated with minor differences in food, incubator temperature, etc., two lines (RAL-315 and  
201 RAL-324) were assayed in each block as internal controls. Variation among blocks in  
202 desiccation resistance of each line was <10%. For these lines, only data from the first block  
203 were included in the overall data analysis.

#### 204 *Genome wide association analysis*

205 We performed genome-wide association analysis on desiccation tolerance using the  
206 Drosophila Genetic Reference Panel (DGRP). Phenotypic line means were uploaded to the  
207 DGRP analysis website (<http://dgrp2.gnets.ncsu.edu>) for genome-wide association analysis  
208 following methods outlined in Mackay et al. (2012) and Huang et al. (2014). Thirty-six (36)  
209 SNPs were associated with sex-averaged desiccation tolerance below nominal  $p$ -value of  $1e-5$   
210 (Fig. S1). These SNPs were annotated as being in or nearby 36 genes (Table S1, Table S2, and  
211 Table S3).

212 We tested whether these 36 genes were more likely to show signals of spatially varying  
213 selection than expected relative to the rest of the genome. We examined patterns of spatially  
214 varying selection using whole genome resequencing of populations sampled along the east coasts  
215 of North America (Bergland et al. 2014) and Australia (Bergland et al. 2016) following a method  
216 outlined in Daub et al. (2013). This method tests whether gene sets identified *a priori* show  
217 stronger signals of spatially varying selection than sets of control genes. To perform this  
218 analysis, we first estimated genetic differentiation at approximately 500,000 common SNPs with  
219 average minor allele frequency greater than 5% (Bergland et al. 2014) among populations of  
220 *Drosophila melanogaster* along latitudinal transects in North America or Australia using the  $T_{FLK}$   
221 statistic (Bonhomme et al. 2010; Bergland et al. 2016). The  $T_{FLK}$  statistic is a modified version of  
222 the classic Lewontin-Krakauer test for  $F_{ST}$  outliers that incorporates certain aspects of population  
223 structure and has been shown to have a low false positive rate when sampled populations result  
224 from secondary contact, as is likely the case for North American and Australian populations of  
225 *D. melanogaster* (Caracristi & Schlotterer 2003; Duchon et al. 2013; Kao et al. 2015; Bergland et  
226 al. 2016).  $T_{FLK}$  values were  $z$ -transformed and, following Daub et al. (2013), we refer to these  
227 transformed  $T_{FLK}$  values as  $z$ . For each gene in the genome, we calculated the maximum  $z$  value,  
228  $z(g)$ , by considering all SNPs within 10Kb of the beginning and end of the gene.  $z(g)$  was  
229 normalized by gene length (hereafter  $z_{st}(g)$ ) by binning all genes with approximately equal length  
230 on a  $\log_2$  scale following equations (1) and (2) of Daub et al. (2013). Next, we generated 1000  
231 sets of control genes matched to the target set associated with desiccation tolerance. These  
232 control sets were matched by chromosome, inversion status at the large cosmopolitan inversions  
233 that segregate on each chromosome (Corbett-Detig & Hartl 2012), recombination rate (Comeron  
234 et al. 2012), and SNP density per basepair. For target and control gene sets we calculated the sum

235 of  $z_{st}(g)$ ,  $SUMSTAT$ , and estimated the probability that  $SUMSTAT_{target}$  is greater than  
236  $SUMSTAT_{control}$ .

### 237 ***Other statistics***

238 Desiccation tolerance is presented as means  $\pm$  s.e., and data were analyzed using ANOVA to  
239 model main effects and interaction terms. As the investigated populations encounter different  
240 temperature and humidity conditions in their natural habitats we performed a multiple regression  
241 analysis of trait values as a simultaneous function of  $T_{min}$ ,  $T_{max}$ ,  $T_{ave}$ ,  $RH_{min}$ ,  $RH_{max}$  and  $RH_{ave}$  of  
242 origin of populations. Respirometry data were analyzed with ANOVA when body mass did not  
243 vary significantly between experimental groups. On the rare occasion (see results) that body  
244 mass did vary, data were analyzed using ANCOVA with body mass as a covariate. JMP v12  
245 (SAS Institute, Cary, NC) and Statistica (Statsoft Inc., Release 7.0, Tulsa, OK, USA) were used  
246 for calculations as well as illustrations.

247

## 248 **Results**

### 249 ***Spatiotemporal variation in desiccation tolerance***

250 Populations from six latitudinal localities of *D. melanogaster* (Fig. 1) exhibited a positive  
251 cline for desiccation tolerance in both sexes, and desiccation tolerance increased positively with  
252 latitude (Fig 2ABC). These patterns were affected by culture temperature, demonstrating an  
253 inverse relationship between temperature and desiccation tolerance (Table 1). The significant  
254 interaction terms of population by temperature and temperature by sex showed that patterns of

255 thermal plasticity varied among populations and were distinct between the sexes (Table 1; Fig.  
256 2ABC).

257 Similarly, we observed a strong, positive correlation between desiccation tolerance and  
258 latitude in adult females that had been exposed for three weeks to environmental conditions that  
259 induce reproductive quiescence (Fig. 2D). This demonstrates that physiological plasticity at the  
260 adult life stage also impacts desiccation tolerance. Furthermore, the differences among  
261 populations were maximized when assayed as adults following this three-week exposure,  
262 although this may also be related to geographic differences in the incidence of reproductive  
263 dormancy (Schmidt et al. 2005). The overall duration of survivorship in the desiccation assay  
264 was lower for older, post-dormancy than in young adults, which could simply reflect biological  
265 age.

266 Regression analysis of desiccation tolerance with geographical and climatic variables  
267 ( $T_{\min}$ ,  $T_{\max}$ ,  $T_{\text{ave}}$ ,  $RH_{\min}$ ,  $RH_{\max}$  and  $RH_{\text{ave}}$ ) for both sexes at all three culture temperatures is given  
268 in Table 2. Temperature parameters exhibited stronger associations with desiccation tolerance  
269 than did humidity parameters. Temperature emerged more significant than humidity parameters  
270 for the variations observed in desiccation tolerance along the east coast of USA. Interestingly,  
271 minimum temperature displayed the most robust association with desiccation tolerance in the  
272 geographic collections. This is consistent with the hypothesis that desiccation tolerance may  
273 have stronger effects on performance and fitness during cooler periods in temperate  
274 environments.

275 Seasonal variation in desiccation tolerance for the three sampled populations (MA, PA,  
276 and VA) is depicted in Figure 3. Again, we observed significant variation in desiccation

277 tolerance among populations, and the significant three-way interaction term for population x  
278 season x sex indicated that the change in desiccation tolerance from spring to fall varied by sex  
279 and population. Contrary to our predictions, however, we did not observe any consistent  
280 difference in desiccation tolerance between spring and fall samples (Table 3;  $F=3.01$ ; d.f.=1;  
281  $p=0.08$ ).

### 282 *Water loss rate and metabolic rate in natural and experimental populations*

283 With the exception of Florida and Maine males, we did not find a significant within-sex  
284 difference in mean body mass among all our comparisons (data not shown). The mean body  
285 mass of male and female flies (ca. 0.7-0.8 and 1.1-1.2 mg, respectively) correlated with the well-  
286 documented higher desiccation resistance of females (e.g. Gibbs et al. 1997). We found no  
287 significant difference for either sex in mean  $\dot{V}CO_2$  or water loss rates (WLRs) between Florida  
288 and Maine, early and late season collections from Pennsylvania, or evolving and fixed  
289 populations (ANOVA, with replication as a random factor,  $p>0.05$ ). When comparing Florida  
290 and Maine males, no significant difference was found in mean  $\dot{V}CO_2$  ( $F_{1,7}=1.61$ ,  $p=0.24$ ) or  
291 water loss rates ( $F_{1,7}=2.76$ ,  $p=0.21$ ), even when accounting for body size (ANCOVA, with body  
292 mass as a covariate).

293 The limitation of our original respirometry approach in detecting variation in WLRs  
294 prompted us to focus on the two most geographically-distant populations for which significant  
295 variation in desiccation resistance were detected (i.e. Florida vs. Maine; see Fig. 2), using an  
296 alternative approach (see Methods). Still, no significant difference was found in the WLRs  
297 among female (ANOVA;  $F_{1,21}=0.005$ ,  $p=0.95$ ) or male ( $F_{1,21}=0.13$ ,  $p=0.71$ ) flies even when  
298 exposed to dry air flow for 3 h. No significant differences were found in  $\dot{V}CO_2$  between Florida

299 and Maine females ( $p=0.68$  and  $0.34$  after 30 min and 3 h, respectively), but similar values for  
300 males after 30 minutes ( $p=0.46$ ) were followed by significantly lower  $\dot{V}CO_2$  values for Maine  
301 compared with Florida males ( $F_{1,21}=4.34$ ,  $p=0.049$ ). Interestingly, when testing for temporal  
302 changes within sets of flies we found a significant increase in  $\dot{V}CO_2$  of Florida males from initial  
303 values ( $3.16\pm 0.15 \mu\text{L}\cdot\text{fly}^{-1}\cdot\text{h}^{-1}$ ) compared to those recorded after 3 h of exposure to desiccation  
304 ( $3.82\pm 0.13 \mu\text{L}\cdot\text{fly}^{-1}\cdot\text{h}^{-1}$ ) (paired t-test;  $t_{11}=5.04$ ,  $p<0.001$ ). In contrast, values after 30 min and 3 h  
305 of desiccation did not vary significantly for Maine male flies ( $t_{11}=1.89$ ,  $p=0.09$ ) (Fig. 4A).  
306 Among female flies, a significant decrease in  $\dot{V}CO_2$  with exposure to desiccation was recorded  
307 for Maine ( $t_{11}=3.69$ ,  $p=0.004$ ), but not Florida populations ( $t_{11}=0.40$ ,  $p=0.70$ ) (Fig. 4B).

### 308 ***GWAS, enrichment and parallelism of SNPs associated with desiccation tolerance***

309 We observed considerable genetic variability among DGRP lines for desiccation  
310 tolerance using a total of 162 lines (Table S1). Line means are depicted in Fig. 5. Females  
311 survived longer than males under desiccating conditions, as expected. Using this among line  
312 variation in the standard mapping pipeline (Mackay et al. 2012), SNPs associated with  
313 desiccation tolerance mapped to or close to 36 genes (Table S2 and Table S3).

314 Among North American populations, genes associated with desiccation tolerance did not  
315 show increased signatures of spatially varying selection relative to the rest of the genome (Fig. 6;  
316  $p=0.8$ ), whereas they do among Australian populations (Fig. 6;  $p=0.02$ ). The probability that  
317  $SUMSTAT_{target}$  is greater than  $SUMSTAT_{control}$  was 0.74, indicating that genes associated with  
318 desiccation are not more differentiated among North American populations than expected  
319 relative to the rest of the genome.

320

## 321 **Discussion**

322 Desiccation resistance in insects could involve one or more adaptive mechanisms,  
323 including (1) increases in total body water content and/or in haemolymph volume (Folk et al.  
324 2001); (2) increased dehydration tolerance (i.e., tolerance of body water loss before death)  
325 (Telonis-Scott et al. 2006); (3) reduction in rate of water loss (Gibbs et al. 1997). In insects,  
326 higher surface area to volume ratio turn them susceptible to desiccation under drier climatic  
327 conditions. (which varied five-fold among 20 *Drosophila* species) and desiccation tolerance but  
328 not with rate of water loss. Based on these interspecific comparisons, it is generally predicted  
329 that body size can play a role in desiccation resistance. Along the coast of eastern Australia, no  
330 cline was observed for desiccation tolerance in *D. melanogaster* (Hoffmann et al. 2001), whereas  
331 along the Indian latitudes a robust cline has been observed (Karan et al. 1998). Interestingly, a  
332 cline for body size (e.g., thorax and wing length) has been observed on both continents (James et  
333 al. 1995; Bhan et al. 2014). In North America, body size also varies clinally and increases with  
334 increasing latitude (e.g., Coyne and Milstead 1987). Here, we observed a significant but shallow  
335 cline in desiccation tolerance at normal assay temperatures of 25°C (Table S4). While body size  
336 may play a role in desiccation tolerance, the relationship between variation in body size and  
337 desiccation tolerance among *D. melanogaster* populations appears complex (Rajpurohit et al.  
338 2016b, unpublished).

### 339 *Spatiotemporal patterns in desiccation tolerance*

340 Our results, demonstrating a positive cline in desiccation tolerance for populations from  
341 the east coast of the U.S, are consistent with previous studies showing higher desiccation  
342 tolerance in temperate vs. tropical locales (Hoffmann & Harshman 1999). On the Indian  
343 subcontinent, this trend was clearly observed in multiple *Drosophila* species (Karan et al. 1998;  
344 Parkash et al. 2008; Rajpurohit & Nedved 2013), where parallel clines for this trait have been  
345 observed in several *Drosophila* species. On the Indian subcontinent higher latitudes in the north  
346 are characterized by lower temperature and lower humidity during winter, whereas low latitude,  
347 southern locations are warm and humid for most of the year. A meta-analysis approach  
348 concluded that  $T_{cv}$  (coefficient of variance in temperature) was a major climatic component to  
349 support the observed parallel clines for desiccation tolerance in several *Drosophila* species on  
350 the Indian subcontinent (Rajpurohit et al. 2013a). Along the east coast of the U.S., more  
351 temperate populations experience greater environmental fluctuations associated with seasonality  
352 and harsher winter conditions. As the temperate winter is generally associated with a highly  
353 desiccating environment, we predicted that populations collected in the spring would be  
354 characterized by relative increases in desiccation tolerance, similar to patterns observed for other  
355 stress-related traits (Behrman et al. 2015). However, we did not observe any differences in  
356 desiccation tolerance between early and late season collections of *D. melanogaster* from three  
357 temperate populations spanning 38 – 42°N latitude. Similarly, we did not observe any  
358 differentiation in water loss rates in response to experimental evolution to seasonality in our field  
359 experiment. Humidity differences between fall vs spring seasons are not significant for any of  
360 these collection sites (e.g., Fig. S2), and this may be associated with the absence of seasonal  
361 variation in desiccation tolerance.

### 362 *Climatic associations*

363           The widespread occurrence of latitudinal clines for many fitness related phenotypes may  
364 be related to the regularity of climatic changes, and associated parameters, with latitude. We  
365 analyzed climatic data along a south-north transect for the six sampled localities ranging from  
366 Florida (25.48 °N Latitude) to Maine (42.26 °N Latitude). Latitude was negatively correlated  
367 with winter temperature. We also observed that the amplitude of thermal seasonal variation,  
368 estimated by the between-month coefficient of variation (CV), increased with increasing with  
369 latitude, from 3.3% in Florida up to 29.7% in Maine (Schmidt et al. 2005).

370           It is generally assumed that desiccation tolerance is selected during hot and dry  
371 conditions, as heat and desiccation stresses generally co-occur (Hoffmann and Parsons 1991). In  
372 natural habitats on the U.S. east coast, however, such is not the case. Variation in desiccation  
373 tolerance is most strongly associated with minimum temperature of the geographic origin of our  
374 collections, suggesting that desiccation tolerance may be favored in environments characterized  
375 by cold and dry conditions (Leather et al. 1993). Cold and desiccation tolerance may also exhibit  
376 correlated responses (e.g., Bublly and Loeschcke 2005; MacMillan et al. 2009). Thus, it remains  
377 unclear whether the latitudinal patterns we observed are driven primarily by selection directly on  
378 desiccation tolerance or may reflect an indirect response due to selection on a correlated trait.

### 379 ***Thermal plasticity***

380           Populations of *D. melanogaster* grown at lower temperatures slightly increased their  
381 desiccation tolerance. The difference in desiccation tolerance hours was in the direction of  
382 18>25>29 °C (see slope comparison in Table S4). A recent study on the cold-adapted *D.*  
383 *nepalensis* from the western Himalayas found that flies grown at 15 °C show twofold higher  
384 body size, greater melanization, higher desiccation resistance, hemolymph and carbohydrate

385 content as compared to flies reared at 25 °C (Parkash et al. 2014). There is a strong possibility  
386 that *D. melanogaster* populations growing at lower temperatures may also exhibit these plastic  
387 responses that could subsequently affect desiccation tolerance. However, we also observed that  
388 variation among populations became exacerbated at lower temperatures, and was most distinct  
389 following exposure to dormancy inducing conditions. Thus, our data also suggest that patterns of  
390 plasticity in *D. melanogaster* may vary predictably among natural populations and habitats.

### 391 ***Geographic variation in metabolic rate***

392 Desiccation resistance in *Drosophila* is associated with reduced water loss rates under  
393 both natural (Kalra et al. 2014) and laboratory conditions (reviewed by Hoffmann and Harshman  
394 1999). In contrast, evidence for other potential adaptive mechanisms is more equivocal. Higher  
395 body water content was reported for resistant populations in some studies (Gibbs et al. 1997;  
396 Chippindale et al. 1998; Folk et al. 2001; Gefen et al. 2006), but not in others (Hoffmann and  
397 Parsons 1993). The ability to tolerate dehydration has also reported to vary between desiccation-  
398 selected populations and their controls in one study (Telonis-Scott et al. 2006), but not another  
399 (Gibbs et al. 1997). However, this discrepancy could simply reflect an inconsistency in the use of  
400 the term dehydration tolerance (Gibbs and Gefen 2009).

401 We found no evidence for variation in water loss rates that could explain the clinal  
402 variation in desiccation tolerance. Water-loss rates of the northernmost (Maine) and  
403 southernmost (Florida) populations did not differ in either males or females. We also recorded  
404 similar metabolic rates, expressed as CO<sub>2</sub> emission rates ( $\dot{V}CO_2$ ) for females from the two  
405 populations, but higher values for severely desiccated Florida male flies suggest significant  
406 differences in the metabolic response of these populations to prolonged exposure to desiccating

407 conditions. This is in agreement with both intraspecific (Hoffmann and Parsons 1993; Gefen and  
408 Gibbs 2009) and interspecific (Gibbs et al. 2003) reports which showed that desiccation  
409 resistance in *Drosophila* is correlated with reduced activity under stressful conditions; these  
410 results are also consistent with pronounced differences in central metabolism between  
411 populations at the geographic extremes of the U.S. east coast (e.g., Verrelli and Eanes 2001;  
412 Flowers et al. 2007; Lavington et al. 2014). It should be noted that increasing  $\dot{V}CO_2$  values could  
413 reflect a switch to carbohydrate catabolism under desiccation stress (Marron et al. 2003),  
414 independent of changing metabolic rates. However, we did not observe a similar response in  
415 females. Instead, the significant decrease in  $\dot{V}CO_2$  in the more resistant Maine females as they  
416 settled to the dry metabolic chamber environment, and absence of this response for Florida  
417 females, suggests a difference in behavioral response. Both males and females in these  
418 populations do exhibit very distinct behavior in response to thermal variation as well (Rajpurohit  
419 and Schmidt 2016).

420       Variation in activity patterns under stressful conditions is likely to result in correlated  
421 differences in respiratory water losses. The similar WLRs reported here for Florida and Maine  
422 flies can be explained by the considerably higher relative importance of cuticular water loss in  
423 insects (Chown 2002), and may suggest that flies across the experimental populations do not  
424 vary in their cuticular resistance to water loss. Nevertheless, while results in this study do not  
425 confirm an effect of activity level on WLRs and thus on desiccation-resistance, they could well  
426 reflect how stressful the exposure to experimental desiccation is to flies from the respective  
427 populations. If the more susceptible Florida flies have lower body water contents when hydrated  
428 compared with the more resistant Maine flies, then at similar WLRs the former would approach  
429 the minimum tolerable hydration state earlier, which could elicit an increase in activity levels as

430 a result of attempts to seek more favorable conditions. In addition, a delayed escape response in  
431 the more resistant flies could indicate higher dehydration tolerance that would trigger an escape  
432 response at lower body water content.

### 433 *Ecological genetics of desiccation tolerance*

434 Relative to other fitness-associated traits (e.g., body size, Coyne and Milstead 1987),  
435 reproductive dormancy (Schmidt et al. 2005), cuticular hydrocarbons (Rajpurohit et al. 2016c  
436 unpublished), thermal preference (Rajpurohit and Schmidt 2016) and body pigmentation  
437 (Rajpurohit et al. 2016a unpublished), we observed a very shallow cline for desiccation tolerance  
438 across the sampled latitudinal gradient in eastern North America. Our results also demonstrated  
439 pronounced patterns of plasticity in response to temperature, both in terms of developmental  
440 plasticity as well as adult acclimation and subsequent response. However, the observed patterns  
441 suggest that spatially varying selection may be less pronounced on this trait, both in comparison  
442 to other traits in North American populations as well as to desiccation tolerance in Australian  
443 (Telonis-Scott et al. 2006) and Indian (Karan et al. 1998) populations. Work done by Telonis-  
444 Scott et al. (2015), in comparison to our GWAS, further suggests the genetic basis of desiccation  
445 tolerance may be more robust and polygenic in Australian populations relative to North America.  
446 Similarly, our analysis of clinal enrichment demonstrated that genes associated with desiccation  
447 tolerance are enriched for clinality in Australia but not in North America. Differences among the  
448 continents may reflect differential local adaptation and the role of desiccation tolerance in  
449 affecting fitness, variation in colonization history and associated demography (Kao et al. 2015;  
450 Bergland et al. 2016), or a combination of the two.

### 451 *Conclusion:*

452           We observed a shallow cline for desiccation tolerance in populations sampled along the  
453 latitudinal gradient in the eastern U.S.; these patterns of variation among populations exhibited  
454 both developmental and adult plasticity, suggesting that further analysis of desiccation tolerance  
455 should be examined under a range of environmental conditions. Climatic analysis of this cline  
456 indicated that observed patterns of desiccation tolerance were most strongly associated with  
457 lower temperature conditions, suggesting that selection on this trait in temperate populations may  
458 be associated with response to desiccating conditions that co-occur with exposure to reduced  
459 temperatures. GWAS analysis using the DGRP panel of inbred lines identified 36 genes  
460 associated with desiccation tolerance in North American populations, providing a wealth of  
461 candidates for subsequent functional analysis and investigation. These genes were not enriched  
462 for signatures of clinality in North American populations but were in Australian populations,  
463 further suggesting differential dynamics of this trait in various habitats of this cosmopolitan  
464 genetic model organism.

465

#### 466 **Data Accessibility**

467 Raw data will be archived at the Dryad digital repository.

468

#### 469 **Acknowledgements**

470           We would like to thank Kelly Dyer for providing isofemale lines collected from Athens,  
471 GA and Emily Behrman for providing seasonal collections from Media, PA. Research in the

472 Schmidt Lab & Gibbs Lab was supported through the National Institute of Health grant  
473 R01GM100366 and National Science Foundation (EnGen-0723930) respectively. This work was  
474 also supported through a grant from the Peachey Environmental Biology Fund to SR. We are  
475 particularly thankful to undergraduate research assistants in the Gibbs Lab for their 24-hour help  
476 in phenotyping DGRP lines.

477

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659

**Table 1:** ANOVA for desiccation tolerance of males and females from the six geographically distinct populations.

<b>Parameters</b>	<b>SS</b>	<b>d. f.</b>	<b>MS</b>	<b>F</b>	<b><i>p</i></b>
Population (P)	532.2	5	106.4	28.74	<b>0.000000</b>
Temperature (T)	1709.3	2	854.7	230.75	<b>0.000000</b>
Sex (S)	14039.0	1	14039.0	3790.42	<b>0.000000</b>
P x T	104.6	10	10.5	2.82	<b>0.001866</b>
P x S	27.5	5	5.5	1.49	0.191275
T x S	33.1	2	16.6	4.47	<b>0.011721</b>
P x T x S	64.3	10	6.4	1.74	0.068731
Error	3229.7	872	3.7		

**Table 2:** Regression analysis of desiccation tolerance with geographical and climatic parameters. The first row gives correlations (*r* values), the second line the goodness of fit from ANOVA. Statistical significance is depicted by bold typeface.

<b>Temperature</b>		<b>Latitude</b>	<b>T<sub>min</sub></b>	<b>T<sub>max</sub></b>	<b>T<sub>ave</sub></b>	<b>RH<sub>min</sub></b>	<b>RH<sub>max</sub></b>	<b>RH<sub>ave</sub></b>
18 °C Male	<i>r</i>	0.8582	-0.8766	-0.6603	-0.7901	-0.9236	-0.1267	-0.6808
	<i>p</i>	<b>0.02874</b>	<b>0.0219</b>	0.15353	0.06148	<b>0.00854</b>	0.81104	0.13654
18 °C Female	<i>r</i>	0.98115	-0.9531	-0.9651	-0.977	-0.6814	-0.5908	-0.7623
	<i>p</i>	<b>0.00053</b>	<b>0.00325</b>	<b>0.0018</b>	<b>0.00079</b>	0.13612	0.2169	0.07804
25 °C Male	<i>r</i>	0.87516	-0.8453	-0.8305	-0.8544	-0.6351	-0.4225	-0.5673
	<i>p</i>	<b>0.02241</b>	<b>0.03405</b>	<b>0.04064</b>	<b>0.03028</b>	0.17541	0.40394	0.24034
25 °C Female	<i>r</i>	0.78074	-0.82	-0.8713	-0.8602	-0.3712	-0.3493	-0.4461
	<i>p</i>	0.06684	<b>0.04567</b>	<b>0.02377</b>	<b>0.02797</b>	0.46884	0.4974	0.37529
29 °C Male	<i>r</i>	0.76591	-0.8095	-0.6357	-0.742	-0.7053	0.06338	-0.3727
	<i>p</i>	0.07579	0.05098	0.17491	0.09126	0.11748	0.90505	0.46687
29 °C Female	<i>r</i>	0.86563	-0.8094	-0.7261	-0.7852	-0.7537	-0.3719	-0.7134
	<i>p</i>	<b>0.02587</b>	0.05102	0.10226	0.06426	0.08353	0.46788	0.11146

**Table 3:** ANOVA for seasonal variations in desiccation tolerance of males and females from the three geographically distinct populations.

<b>Parameters</b>	<b>SS</b>	<b>d.f.</b>	<b>MS</b>	<b>F</b>	<b><i>p</i></b>
Population (Pop)	109.66	2	54.83	14.95	<b>0.000001</b>
Season (Sea)	11.06	1	11.06	3.01	0.083603
Sex	6188.14	1	6188.14	1686.67	<b>0.000000</b>
Pop*Sea	21.49	2	10.74	2.93	0.055111
Pop*Sex	133.73	2	66.87	18.23	<b>0.000000</b>
Sea*Sex	14.78	1	14.78	4.03	<b>0.045685</b>
Pop*Sea*Sex	67.72	2	33.86	9.23	<b>0.000131</b>
Error	1034.62	282	3.67		

## Legends of Figures

**Fig. 1:** Temperature map of the east coast of the U.S.A. showing the populations of *D. melanogaster* that were collected and assayed. Seasonal collections were done in Lancaster, MA, Media, PA and Charlottesville, VA in 2012.

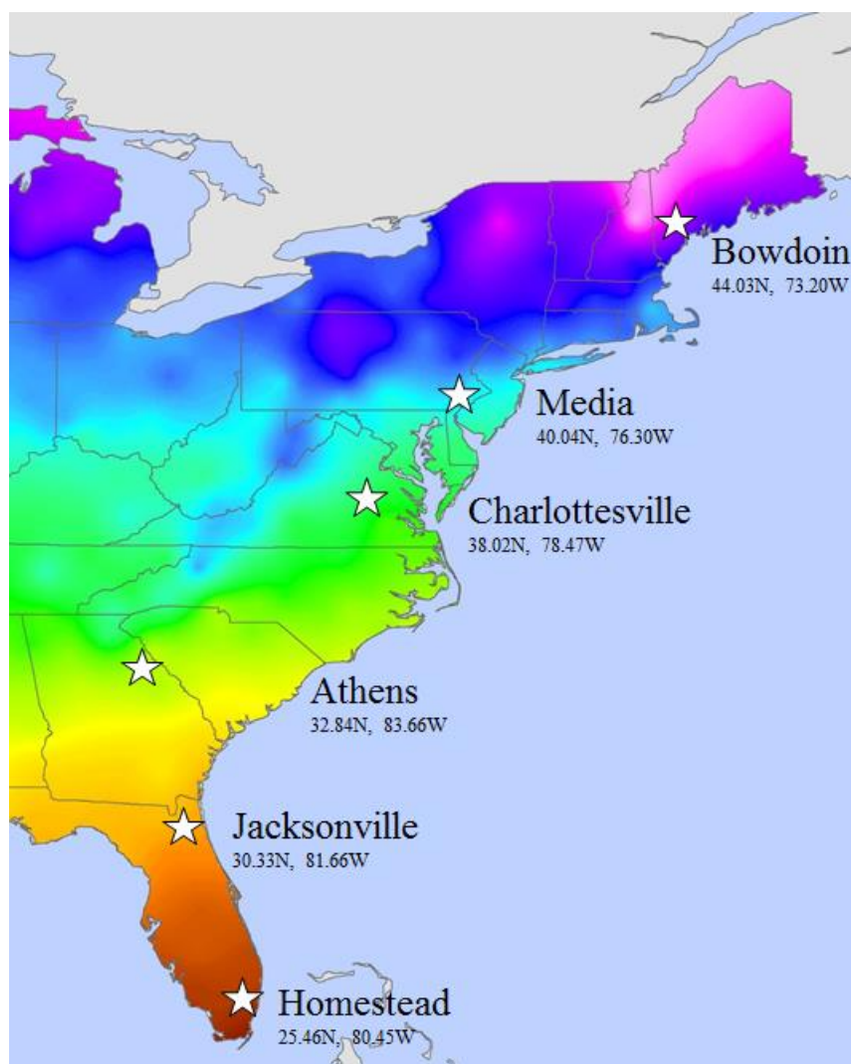
**Fig. 2:** Data (mean $\pm$ s.e.) on desiccation tolerance for males and females of *D. melanogaster* from six geographical locations (see Fig. 1) at four different thermal conditions (18 °C (A), 25 °C (B), 28 °C (C), and Diapause conditions)). To study over-wintering affects (D) over desiccation tolerance flies were kept under dormancy inducing conditions (11 °C and 09:15 photoperiod) for 3 weeks before subjecting them to desiccation tolerance assay (at 25 °C). Males and females are denoted as triangles and rectangles respectively.

**Fig. 3:** Data (mean $\pm$ s.e.) on early and late season desiccation tolerance for males and females of *D. melanogaster* from three geographical locations (Lancaster, MA; Media, PA; Charlottesville, VA). Open, gray and black bars represent MA, PA and VA populations respectively.

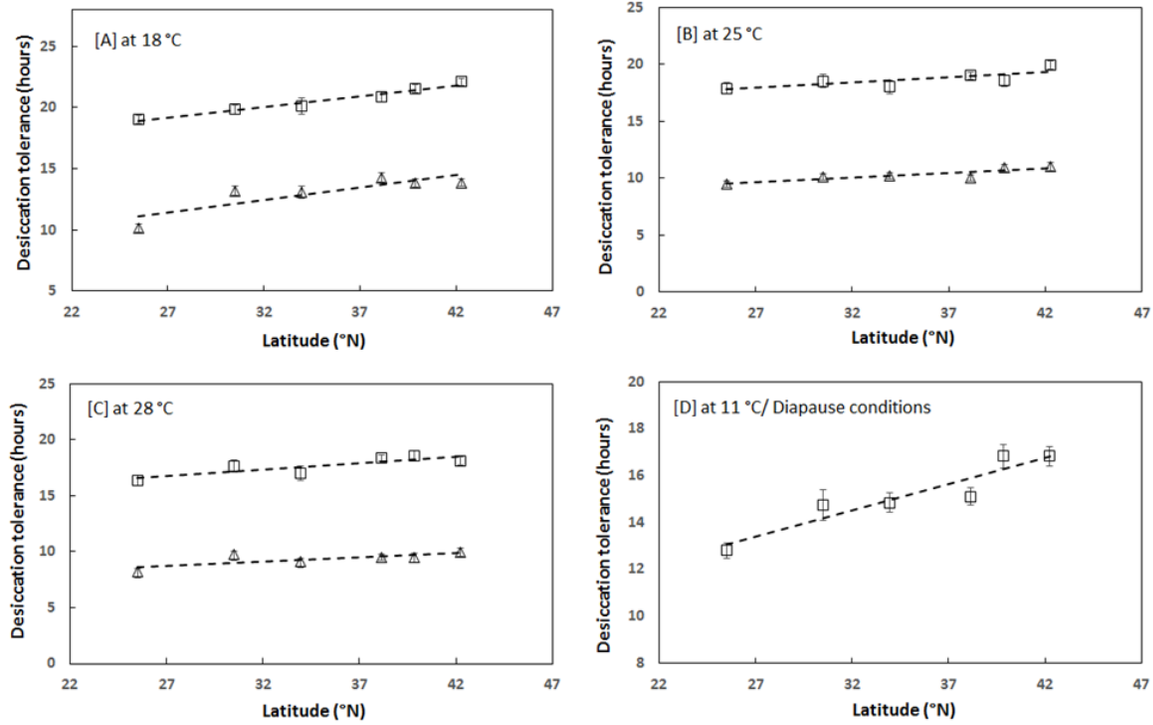
**Fig. 4:** Carbon dioxide emission rates for males (A) and females (B) of *D. melanogaster* populations from Florida (FL) and Maine (MN) locations. Measurements were done at two time points (30 min and 3hr). Asterisks indicate significant difference from 30 min values ( $\alpha=0.05$ ).

**Fig. 5:** Status of desiccation tolerance (line means) in Drosophila Genetic Resource Panel. A total of 162 lines were considered for this association mapping. A considerable genetic variability in desiccation tolerance was observed across the lines. Males (A) survived shorter under desiccating conditions than females (B).

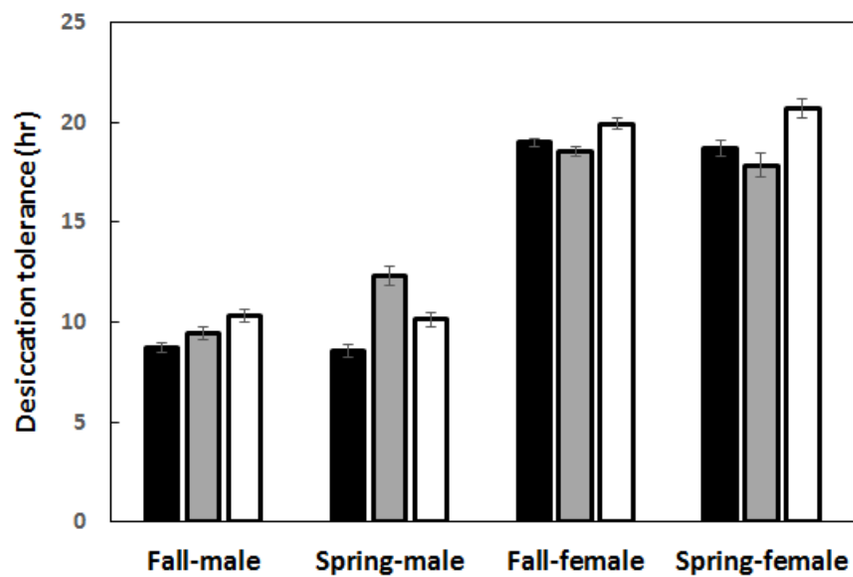
**Fig. 6:** SumStat scores for 1000 sets of control genes (distribution) and observed SumStat score (dashed red line) for genes associated with desiccation tolerance among populations sampled along the east coast of North America (Nam) or Australia (Aus). See Materials and Methods for a description of the SumStat score.



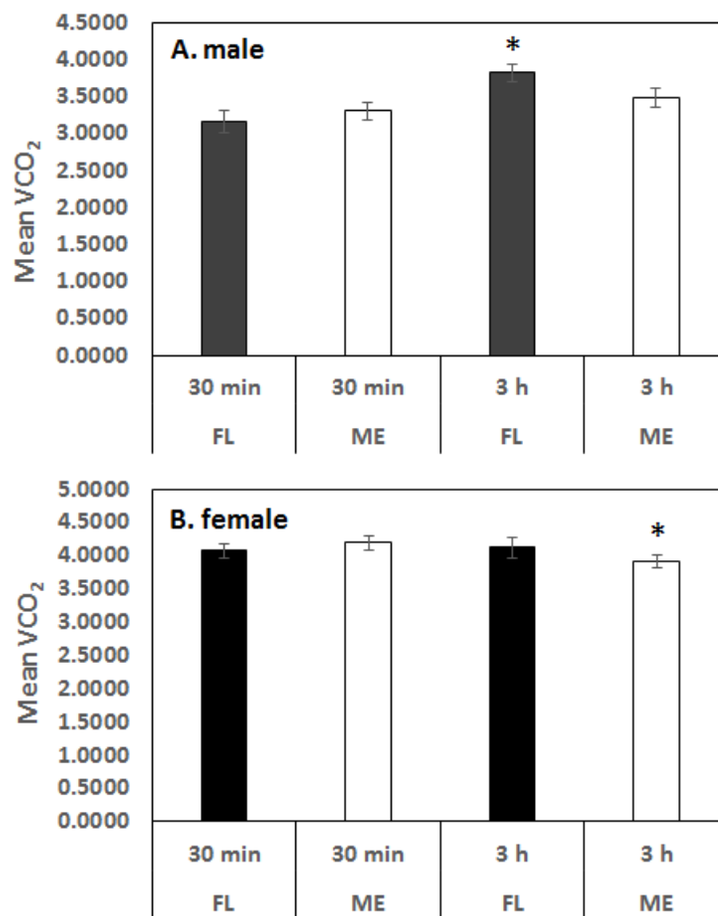
**Figure 1**



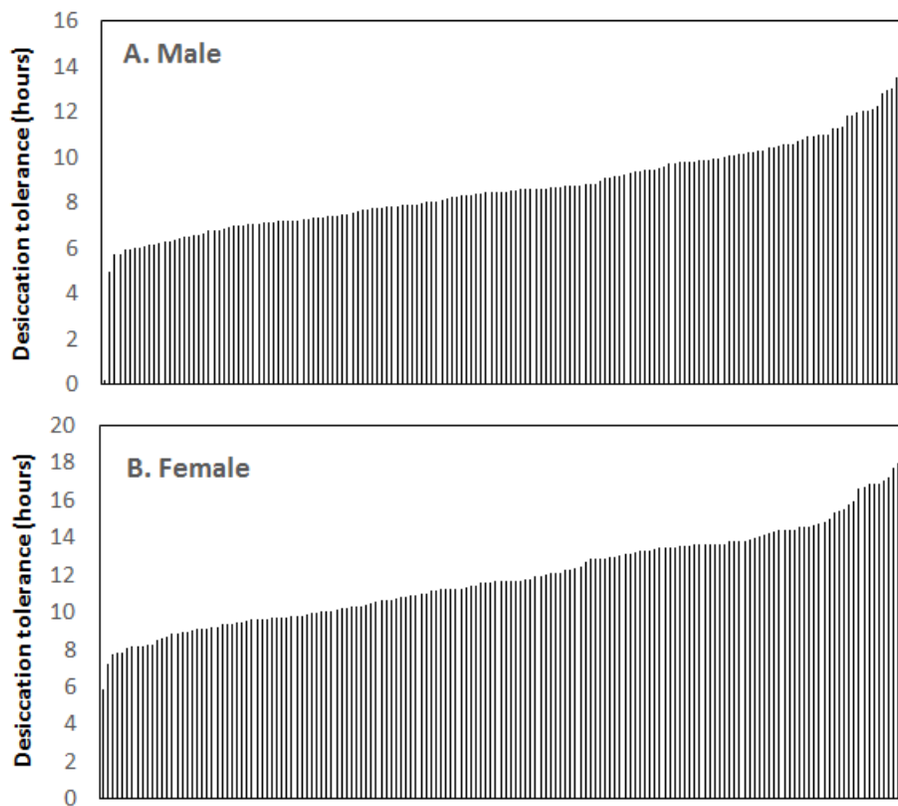
**Figure 2**



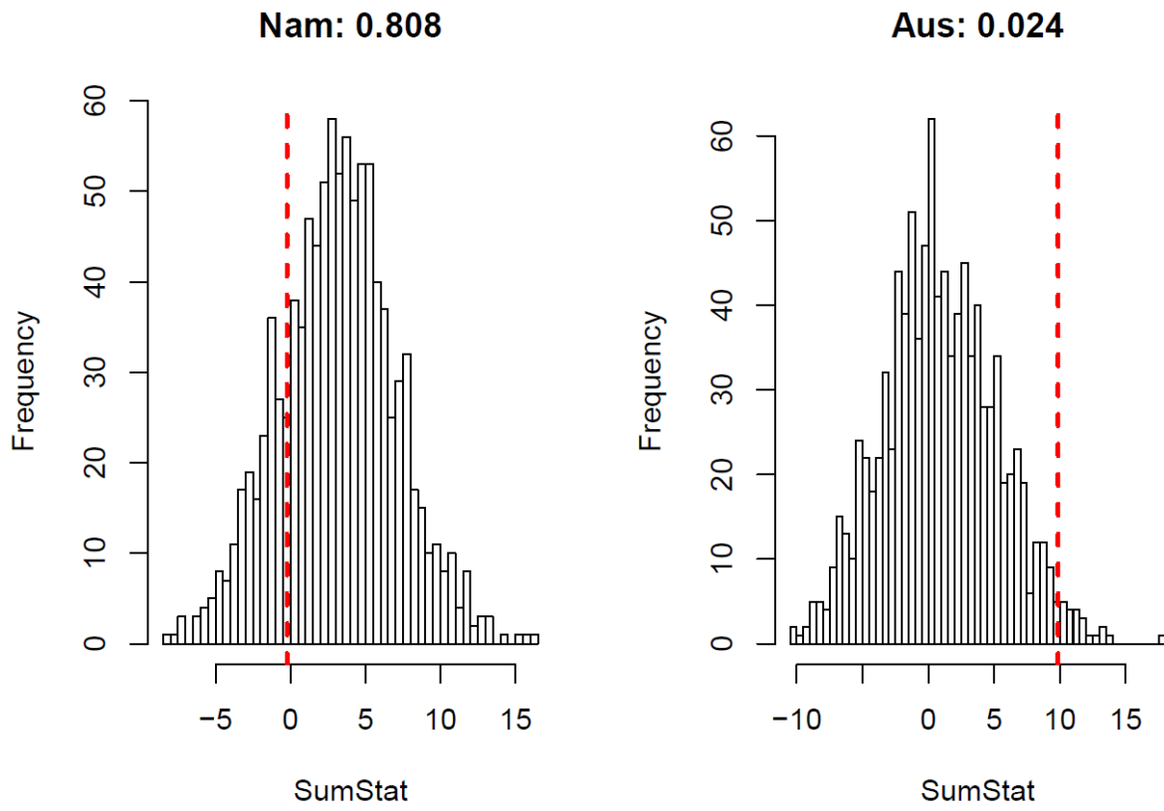
**Figure 3**



**Figure 4**



**Figure 5**



**Figure 6**

## Supporting Figures

**Fig. S1:** qq-plot of genome wide association analysis of intersex averaged desiccation tolerance. Observed p-values are plotted along the y-axis and expected p-values along the x-axis. The solid red line depicts the 1:1 line.

**Fig S2:** 3D Surface plot (A) of  $T_{ave}$  and  $RH_{ave}$  along the latitudes of east coast of U.S.A.  $RH_{ave}$  = Distance weighted least square. Geographical clines in fitness traits and their interactions with climatic variables are well known. To explore these interactions in this study we planned to study the relationship between desiccation tolerance and climatic variables (temperature and relative humidity) of the origin of sites of populations. Temperature and relative humidity data for sample collection sites averaged over 30 years (from 1980 to 2010). Seasonal variations (Spring vs. Fall) in relative humidity of the origin of sites of population collections are also presented (B). All the historical data of temperature and humidity were downloaded from the National Climatic Data Center (NOAA; <http://www.ncdc.noaa.gov/>).