Spatiotemporal dynamics and genome-wide association genome-wide association analysis of desiccation tolerance in Drosophila melanogaster

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Abstract
Water availability is a major environmental challenge to a variety of terrestrial organisms. In insects, desiccation tolerance varies predictably over spatial and temporal scales and is an important physiological determinant of fitness in natural populations. Here, we examine the dynamics of desiccation tolerance in North American populations of Drosophila melanogaster using: (a) natural populations sampled across latitudes and seasons; (b) experimental evolution in field mesocosms over seasonal time; (c) genome-wide associations to identify SNPs/genes associated with variation for desiccation tolerance; and (d) subsequent analysis of patterns of clinal/seasonal enrichment in existing pooled sequencing data of populations sampled in both North America and Australia. A cline in desiccation tolerance was observed, for which tolerance exhibited a positive association with latitude; tolerance also varied predictably with culture temperature, demonstrating a significant degree of thermal plasticity. Desiccation tolerance evolved rapidly in field mesocosms, although only males showed differences in desiccation tolerance between spring and autumn collections from natural populations. Water loss rates did not vary significantly among latitudinal or seasonal populations; however, changes in metabolic rates during prolonged exposure to dry conditions are consistent with increased tolerance in higher latitude populations. Genome-wide associations in a panel of inbred lines identified twenty-five SNPs in twenty-one loci associated with sex-averaged desiccation tolerance, but there is no robust signal of spatially varying selection on genes associated with desiccation tolerance. Together, our results suggest that desiccation tolerance is a complex and important fitness component that evolves rapidly and predictably in natural populations.

KEYWORDS
cline, desiccation tolerance, Drosophila, plasticity, season

INTRODUCTION
Insects inhabit a wide range of habitats in which water is often scarce and desiccation represents a major physiological challenge. Insects are particularly vulnerable to water-related challenges, due to their small size and thus large surface area to volume ratio (Edney, 1977; Gibbs & Rajpurohit, 2010; Hadley, 1994). Environmental stressors such as desiccation are highly variable among natural habitats and often vary predictably with such features as latitude, altitude and season (Endler, 1977; Hoffmann & Parsons, 1991; Parkash,
Patterns of variation across these gradients (Telonis-Scott, Hoffmann, & Sgro, 2011), particularly when genetic/genomic and phenotypic variation are integrated, offer a means to address the extent to which divergence in physiological traits is affected by natural selection and may represent an adaptive response to environmental heterogeneity (Barton, 1999; Whitlock & McCauley, 1999).

In Drosophila, various species have adapted to diverse climatic conditions during their evolutionary history and thus constitute good models for the integrated study of physiological adaptation and population ecology (Coyne, Bundgaard, & Prout, 1983; Gibbs and Matzkin, 2001; Lemeunier, David, Tsacas, & Ashburner, 1986; Parsons, 1983). High levels of desiccation resistance are associated with adaptation to arid habitats (e.g., David, Allemand, Van Herewege, & Cohet, 1983; Gibbs, Fukuzato, & Matzkin, 2003; Hoffmann & Parsons, 1991). Likewise, geographical variation for desiccation tolerance among populations has been widely observed (Griffiths, Schiffer, & Hoffmann, 2005; Matzkin, Watts, & Markow, 2007, 2009; Rajpurohit & Nedved, 2013; Rajpurohit, Nedved, & Gibbs, 2013; Rajpurohit, Oliveira, Etge, & Gibbs, 2013; Rajpurohit, Zhao, & Schmidt, 2017b). These patterns suggest that desiccation resistance is an important fitness component that varies predictably within and among taxa.

However, the genetic architecture of desiccation tolerance remains unresolved. In Drosophila melanogaster, tolerance evolves rapidly in response to artificial selection in the laboratory (Bradley, Williams, & Rose, 1999; Gefen, Marlon, & Gibbs, 2006; Gibbs, Chippindale, & Rose, 1997; Telonis-Scott, Sgro, Hoffmann, & Griffin, 2016) and appears associated with reduced water loss rates (Rajpurohit, Peterson, Orr, Marlon, & Gibbs, 2016; Telonis-Scott, Guthridge, & Hoffmann, 2006). Selection for increased tolerance results in differential gene expression (Sorensen, Nielsen, & Loeschcke, 2007) and significant allele frequency changes across hundreds of genes (Telonis-Scott, Gane, DeGaris, Sgro, & Hoffmann, 2012; Telonis-Scott et al., 2006), indicating a complex genetic architecture (Foley & Telonis-Scott, 2011; Matzkin et al., 2009; Rajpurohit, Oliveira, et al., 2013). A genome-wide association study in a natural population recently derived from southern Australia identified a number of genetic variants associated with differences in desiccation tolerance, further demonstrating that desiccation tolerance is a highly polygenic trait (Telonis-Scott et al., 2016). However, it remains unclear whether the architecture of this trait is parallel between natural populations exposed to pronounced environmental heterogeneity at various spatial and temporal scales vs. laboratory-based populations subjected to strong artificial selection over shorter timescales.

The observed clines in desiccation tolerance suggest a robust association with climate and imply spatially variable selection on tolerance in natural populations. However, whether the observed phenotypic clines are mirrored by allele frequency clines at identified candidates (e.g., Telonis-Scott et al., 2016) is unknown. Furthermore, clines may be generated by processes other than selection (e.g., Bergland, Tobler, Gonzalez, Schmidt, & Petrov, 2016; Caracristi & Schlötterer, 2003; Kao, Zubair, Salomon, Nuzhdin, & Campo, 2015), and the timescale over which clines are generated and maintained is often unclear. If variation in desiccation tolerance is driven by spatially varying selection mediated by climate, and may respond rapidly to climate change (Telonis-Scott et al., 2016), then tolerance might also vary predictably with seasonal variation in climatic parameters in temperate habitats. The results from artificial selection studies suggest that desiccation tolerance can evolve rapidly in the laboratory (<10 generations; Hoffmann & Parsons, 1989); however, whether this occurs in natural populations is unknown.

Here, we examine some basic dynamics of desiccation tolerance in natural populations sampled from a range of temperate habitats in North America. We predicted that, as has been shown in Indian populations, desiccation tolerance would vary with latitude (Karan et al., 1998; Parkash, Rajpurohit, & Ramniwas, 2008; Rajpurohit, Nedved, et al., 2013) and that such patterns among populations would reflect selection for increased tolerance at higher latitudes that are characterized by increased winter severity and length. We tested this by sampling populations across the latitudinal gradient in the eastern United States and examining genetically based variation in tolerance across a range of developmental temperatures. Likewise, we hypothesized that desiccation tolerance would be higher in populations sampled in the spring (following the winter season in which maintenance of water balance in overwintering adults might be an important fitness component) as compared to populations sampled in the fall. This hypothesis stems from recent observations that a variety of fitness-associated traits evolve rapidly in D. melanogaster over seasonal time (Behrman, Watson, O’Brien, Heschel, & Schmidt, 2015; Behrman et al., 2018; Rajpurohit et al., 2017a), and was tested by examining seasonal change in desiccation tolerance for three mid-latitude populations sampled from the eastern United States. To examine whether desiccation tolerance evolves rapidly outside the laboratory environment, we employed a field-based experimental evolution design to examine whether tolerance changes over the course of the agricultural growing season from spring to fall.

Prior work on the genetic basis of variance in desiccation tolerance (Foley & Telonis-Scott, 2011; Telonis-Scott et al., 2012, 2016) has provided a wealth of information regarding the architecture of desiccation tolerance, as well as an identification of candidates for subsequent functional analysis (Griffin, Hangartner, Fournier-Level, & Hoffmann, 2017; Kang, Aggarwal, Rashkovetsky, Korol, & Michalak, 2016). Here, we extend this to include a genome-wide association study (GWAS) using the panel of inbred lines that constitute the Drosophila Genetic Reference Panel (DGRP; Mackay et al., 2012). We compare the variants associated with desiccation tolerance in the DGRP to those identified in Australian populations (Telonis-Scott et al., 2012, 2016) to examine parallelism across continents and data sets. Given the documented latitudinal clines for desiccation tolerance and the inference that such patterns are generated by local adaptation to climatic variables, the simple prediction is that the genes underlying variation in desiccation tolerance will exhibit latitudinal clines in allele frequency and signatures of spatially varying
selection. We test this prediction by examining whether the SNPs associated with variation in desiccation tolerance in the DGRP exhibit evidence of spatially varying selection, using existing population-level PoolSeq data (Bergland, Berhman, O’Brien, Schmidt, & Petrov, 2014; Kolaczkowski, Kern, Holloway, & Begun, 2011; Telonis-Scott et al., 2016).

2 | MATERIALS AND METHODS

2.1 | Patterns of desiccation tolerance in natural, experimental and inbred populations

2.1.1 | Spatiotemporal variation

Six natural populations of Drosophila melanogaster were collected from fruit orchards located along the east coast of the United States by a combination of aspiration and baiting/sweeping with aerial nets (see Figure 1). Gravid females were immediately sorted into isofemale lines in the field; once the resulting progeny eclosed, lines were typed to species. Approximately 150 isofemale D. melanogaster lines were collected from each locale. Populations sampled were as follows: Bowdoin, Maine (ME; 44.03N, 76.30W); Media, Pennsylvania (PA; 40.04N, 76.30W); Charlottesville, Virginia (VA; 38.03N, 78.48W); Athens, Georgia (GA; 32.84N, 83.66W); Jacksonville, Florida (JFL) 30.33N, 81.66W); and Homestead, Florida (HFL) 25.46N, 80.45W). Independent seasonal collections were performed in Media, Pennsylvania (PA), Lancaster, Massachusetts (MA; 42.455N, 71.67W)) and Charlottesville, Virginia (VA) orchards in June and November of 2012 (N = 6 populations). Long-term maintenance of all populations was carried out using standard, discrete generational culture of isofemale lines at 24°C, 12-hr:12-hr light:dark photoperiod, with a generation time of 21 days. Flies were maintained on cornmeal–molasses–agar media; prior to use in experiments, density was standardized at 40–50 eggs per vial. All desiccation tolerance assays were conducted on individual isofemale lines from the latitudinal and seasonal collections (see below).

Geographic clines in fitness traits and their interactions with climatic variables are well known (e.g., Rajpurohit & Nedved, 2013). To explore these interactions, we examined the relationship between desiccation tolerance and various climatic parameters (temperature and relative humidity) associated with the origin of the sampled populations. As the investigated populations encounter different temperature and humidity conditions in their natural habitats, we performed a multiple regression analysis of trait values as a simultaneous function of $T_{min}$, $T_{max}$, $T_{ave}$, $R_{Hmin}$, $R_{Hmax}$ and $R_{Have}$ of the origin of populations. Climatic data were obtained from the National Oceanic and Atmospheric Administration (NOAA; www.ncdc.noaa.gov). We used 30-year averages (1980–2010) for all the climatic parameters (Supporting Information Table S1). Statistical analyses were conducted in JMP v12 (SAS Institute, Cary, NC) and Statistica (StatSoft Inc., Release 7.0, Tulsa, OK, USA). We did not perform a sequential regression analysis of climatic factors on desiccation tolerance phenotype, as we did not observe any clear relationship between environmental variables and desiccation resistance or plasticity in tolerance.

2.1.2 | Thermal plasticity

To examine thermal plasticity for desiccation tolerance in the latitudinal collections, we primarily examined the effects of developmental temperature, where populations were cultured from egg to adult at three different temperatures that reflect a standardized culture range from cool to control to warm (18, 25 and 29°C; all at 12L:12D). We also examined desiccation tolerance in adult females that were exposed to environmental conditions that can elicit reproductive dormancy (11°C, 9L:15D; Saunders, Henrich, & Gilbert, 1989). This treatment was included for two reasons. First, reproductive dormancy is associated with overwintering physiology, including an increase in desiccation tolerance (e.g., Denlinger, 2002); second, we wished to examine one aspect of short-term, physiological plasticity for desiccation tolerance in the adult stage only.

FIGURE 1  Temperature map of the east coast of the United States showing the populations of Drosophila melanogaster that were collected and assayed. Collection sites for the latitudinal and seasonal populations are shown in white stars and black dots, respectively. Bowdoin, Maine (ME); Lancaster, Massachusetts (MA); Media, Pennsylvania (PA); Charlottesville, Virginia (VA); Athens, Georgia (GA); Jacksonville, Florida (JFL), Homestead, Florida (HFL) [Colour figure can be viewed at wileyonlinelibrary.com]
For developmental thermal plasticity, flies from each line \((N = 15–25)\) from each of the six geographic population were allowed to oviposit for 2–3 hr at 25°C in successive culture vials; egg density was standardized at 40–50 eggs per vial by manual removal. Replicate vials \((N = 30)\) of each line were then randomly assigned to one of the three temperature treatments (10 replicate vials per line per population per temperature) and cultured in Percival I36VL incubators. For the dormancy treatment, freshly eclosed adult females from the density-controlled cultures at 25°C were collected in groups of ten from each isofemale line and exposed to low-temperature and short-day photoperiods (11°C, 10L:14D) for 3 weeks. To further examine spatiotemporal variations, thermal plasticity and their interactions with climatic variables, we performed a principal component analysis (PCA) in Xlstat using an unconstrained ordination technique.

For the latitudinal, seasonal and plasticity analyses, four-to-five day-old virgin flies from each isofemale line were transferred to empty vials (narrow plastic Drosophila culture vials, Genesee Scientific) in groups of ten and restricted to the lower half of the vials by a foam stopper. Silica gel (5 g) was then added above the stopper to maintain low humidity (RH ~3%), and the vial was sealed with Parafilm™ (see Gibbs et al., 1997). Mortality was recorded at hourly intervals until all flies were dead. Desiccation tolerance assays were conducted using individual isofemale lines under common garden laboratory conditions of 25°C, 12L:12D in Percival I36VL incubators. Data on latitudinal, seasonal and thermal plasticity variation in desiccation tolerance were analysed using a mixed-model nested ANOVA in JMPv.12 (SAS Institute, Cary, NC), where the response variable was time until death. Isofemale lines were nested within populations and considered a random factor; population, temperature and sex were treated as fixed factors. LT50 means were calculated using sigmoidal logit regression in Origin 9 (Zar, 2010) and were used for visual presentation.

### 2.1.3 Experimental evolution in the field

To examine whether desiccation evolution evolves rapidly in the field in response to changing environmental conditions over seasonal time, we established 10 experimental mesocosms at a field site in Philadelphia, PA, USA. Each mesocosm was an 8-m² outdoor insect rearing enclosure (Bioquip Products, Gardena, CA) surrounding a mature (dwarf) peach tree (Supporting Information Figure S1). Each mesocosm was seeded with 1,000 individuals (500 males, 500 females) derived from a collection made in 2012 from the same PA orchard as described above. 500 ml of standard cornmeal–molasses medium in loaf pans was placed in each enclosure every other day for the duration of the experiment (July 13, 2014–November 1, 2014). Flies were allowed to oviposit on the fresh food for a period of 2 day, after which point the food was sealed with cotton rope/mesh and larvae allowed to develop; upon eclosion, adults were released into the mesocosms. Thus, the experimental populations were cultured under a natural regime of overlapping generations. Temperature and relative humidity in all experimental cages were recorded using HOBO U23 Pro v2 data loggers (Onset Computer Corp., Bourne, MA, USA). Further experimental details are provided in Rajpurohit et al. (2017a). Over the course of the experiment, samples were taken at four time points for phenotypic characterizations: July 25, August 19, September 11 and October 10, 2014. Approximately 2,000 embryos were collected from each of the 10 cages at each time point, brought back to the laboratory, and subsequently passed through two additional generations of common garden, density-controlled culture (25°C, 12L:12D, 40–50 eggs/vial). In the F3 generation subsequent to each field collection (10 cages x four time points), three replicate vials of 10 individuals per sex were assayed and analysed for desiccation tolerance at 5 days of age according to the methods described above. Data were analysed with a mixed-model ANOVA with cage as a random factor, replicate nested within cage as a random factor and time point treated as an ordinal predictor.

### 2.1.4 Inbred panel (DGRP)

The panel of DGRP inbred lines (Mackay et al., 2012), derived from a single population in Raleigh, NC, USA, was obtained from the Bloomington Drosophila Stock Center and maintained in the laboratory on yeast–cornmeal–sucrose medium at 25°C. For assays, four to six day-old flies were sorted by sex and held on fresh media for 2 days. Assays were conducted on ten replicate groups of five flies per sex for each of the independent lines. We analysed 175 DGRP lines in all.

Desiccation assays were performed in blocks of ~30 lines. Because of the time required to initiate desiccation stress for all lines and to count dead flies, we recorded the exact time when desiccant was added to assay vials and the exact times when each line was checked. Flies that died before the first survival check were assumed to have been injured by handling or other stress and were not included in the data analysis. Any flies dying between two checks were assumed to have died midway between them. To assess potential variation among blocks associated with minor differences in food, incubator temperature, etc., two lines (RAL-315 and RAL-324) were assayed in each block as internal controls. Variation among blocks in desiccation resistance of each line was <10%. For lines assayed multiple times, only data from the first block were included in the overall data analysis. Data are presented as line means ± SE. The presence of Wolbachia and cosmopolitan chromosomal inversions, both of which can affect a variety of physiological and life history traits (e.g., Brummel, Ching, Seroude, Simon, & Benzer, 2004; Umina, Weeks, Kearney, McKenchnie, & Hoffmann, 2005), was included in the examination of among-line variation in desiccation tolerance in the DGRP panel. Line-specific data on infection and infection status were obtained from Huang et al. (2014).

### 2.2 Respirometry

Due to logistical limitations, for respirometry measurements (water loss rate and metabolic rate) we utilized population cages as opposed to individual isofemale lines. We created two population cages for each of the spatial (six populations x two replicates) and
temporal (two seasons × three populations × two replicates) collections (see above). Each cage was created using independent sets of 25 isofemale lines by releasing ten mated females from each line into 12 × 12 × 12 inch insect enclosures (Live Monarch Foundation, Boca Raton, Florida, USA). These lines were maintained in mass culture and allowed to outcross for five generations; subsequently, samples were collected for the phenotypic assays described below.

Virgins were collected within 3 hr of eclosion, separated by sex into sets of 15 individuals and placed in fresh food vials. A total of 24 sets were assayed for the latitudinal extremes of ME and HFL (two geographic regions × two sexes × two replicate populations × three experimental replicates), and 24 for seasonal comparisons in Pennsylvania (two seasons × two sexes × two biological replicate cages × three experimental replicates). Respiriometry was carried out at 4–6 days posteclosion. Flies were transferred directly from their food vials to a 4-ml glass metabolic chamber with aluminium stoppers, which was covered with a black cardboard sleeve to reduce activity in the chamber. Flow-through respiriometry at 25°C was carried out using two channels of a flow multiplexer (RM-8, Sable Systems International, Las Vegas, NV, USA), where dry, CO2-free air was supplied to the chambers at 50 ml/min using factory-calibrated mass flow controllers (MC-500 sccm; Alicat Scientific, Tucson, AZ, USA), and recurrent air from the measured chamber was passed through a LI-7000 CO2/water vapour dual analyser (Li-Cor Biosciences, Lincoln NE, USA). Prior to measurement, the flies were acclimated to the experimental chambers and air flow for 15 min during which flies from the alternative chamber were measured. Identical empty chambers were used for baselining. Analyser voltage output was recorded, stored and analysed using UI-2 data acquisition interphase and EXPEDATA software (Sable Systems International). Recording rate was set to 1 Hz, and only data from the last 5 min of each 15-min run were averaged for analysis.

We used an additional experimental approach to compare the HFL and ME populations for which significant differences in desiccation resistance were found (see Section 3). It was previously reported that water loss rates of D. melanogaster under similar experimental conditions stabilize only after >2 hr of exposure (Gibbs et al., 1997), and therefore, we carried out respirometry on additional sets of flies (two geographic regions × two sexes × two replicate populations × six experimental replicates) that were randomly allocated to six multiplexer channels, with a seventh used for baselining. The measurement sequence of 20-min measurements was as follows: baselining, three experimental chambers, baselining, three additional experimental chambers and finally baselining again for a total file recording of 3 hr, which was immediately followed by another recording at the same sequence. As each set of files was added to the respirometry set-up 20 min prior to the initial measurement, and only data from the last 10 min were analysed, the flies were assayed 30 min and following additional 3 hr during which the flies were exposed to experimental temperature and dry air flow conditions. Respiriometry data were analysed with ANOVA when body mass did not vary significantly between experimental groups. On the rare occasion (see results) that body mass did vary, data were analysed using ANCOVA with body mass as a covariate.

### 2.3 GWAS and signatures of clinality

We performed genome-wide association analysis on desiccation tolerance using 175 assayed inbred lines from the Drosophila Genetic Reference Panel (DGRP). Phenotypic line means were uploaded to the DGRP analysis website (http://dgrp2.gnets.ncsu.edu) for genome-wide association analysis following established methods (Huang et al., 2014; Mackay et al., 2012). From this analysis, twenty-five SNPs were associated with sex-averaged desiccation tolerance below the nominal p-value of 1e−5 (Supporting Information Figure S2; see Supporting Information Table S4). These SNPs were annotated as being in or nearby 20 genes and one noncoding region (Supporting Information Tables S4 and S5); the identified SNP located in the noncoding locus (CR43681) was omitted from further analysis. Visual analysis of the qq-plot (Supporting Information Figure S2) suggests that the observed p-value distribution is not overinflated.

We tested whether the set of 20 identified genes were more likely to show signals of spatially varying selection than expected relative to the rest of the genome. We examined patterns of spatially varying selection using whole genome resequencing of populations sampled along the east coasts of North America (Bergland et al., 2014, 2016) and Australia (Kolaczkowski et al., 2011) following a method outlined in Daub et al. (2013). This method tests whether gene sets identified a priori show stronger signals of spatially varying selection than sets of control genes. To perform this analysis, we first estimated genetic differentiation at approximately 500,000 common SNPs with average minor allele frequency greater than 5% (Bergland et al., 2014) among populations of D. melanogaster along latitudinal transects in North America or Australia using the $T_{FLK}$ statistic (Bergland et al., 2016; Bonhomme et al., 2010). The $T_{FLK}$ statistic is a modified version of the classic Lewontin-Krakauer test for $F_{ST}$ outliers (Lewontin & Krakauer, 1973) that incorporates certain aspects of population structure and has been shown to have a low false positive rate when sampled populations result from secondary contact, as is likely the case for North American and Australian populations of D. melanogaster (Bergland et al., 2016; Caracristi & Schlötterer, 2003; Duchen, Zivkovic, Hutter, Stephan, & Laurent, 2013; Kao et al., 2015). $T_{FLK}$ values were z-transformed and, following Daub et al. (2013), we refer to these transformed $T_{FLK}$ values as $z$. For each gene in the genome, we calculated the maximum $z$ value, $z(g)$, by considering all SNPs within 10Kb of the beginning and end of the gene. $z(g)$ was normalized by gene length (hereafter $z_{ST}(g)$) by binning all genes with approximately equal length on a log2 scale following equations (1) and (2) of Daub et al. (2013). Next, we generated 5,000 sets of control genes matched to the target set associated with desiccation tolerance. These control sets were matched by chromosome, inversion status at the large cosmopolitan inversions that segregate on each chromosome (Corbett-Detig & Hartl, 2012), recombination rate (Cameron, Ratnappan, & Bailin, 2012) and SNP density surrounding each gene, and the number of genes that are wholly or partially within
the ±10-kb window surrounding each gene. For each target gene, we calculated the probability that its \( z_d(q) \) was greater than its specific control genes and report one minus this probability as the bootstrap p-value for that gene. In addition, following Daub et al. (2013), for target and control gene sets we calculated the sum of \( z_d(q) \), SUMSTAT, and estimated the probability that SUMSTAT target is greater than SUMSTAT control.

Gene ontology enrichment analysis was performed using GOWINDA (version 1.12). Because the number of sex-averaged desiccation resistance SNPs was so small, we also included SNPs with p-values below \( 10^{-3} \) for either males or females. This total of 103 candidate SNPs (Supporting Information Table S6) was tested against a background of 4,438,427 SNPs found within the DGRP populations (dgrp2.vcf). Gene annotations were derived by converting a general feature format (gff) file from FLYBASE (version r5.49.gff) into a gene transfer format (gtf) file with the Gff2Gtf.py Python script from GOWINDA. Gene aliases used for this step were extracted from the most recent gff release on FLYBASE. Gene ontology associations were downloaded from FUNCAS-associate 3.0. Gene identifiers between the gene annotation and gene transfer format (gtf) file with the Gff2Gtf.py Python script from GOWINDA. Gene ontology associations were downloaded from FUNCAS-associate 3.0. Gene identifiers between the gene annotation and gene ontology association files were synchronized with the SynchronizeGtfGeneIDs.py script from GOWINDA. GOWINDA was set to perform 1e6 simulations, with the –mode parameters set to “gene.”

3 | RESULTS

3.1 | Spatiotemporal variation in desiccation tolerance

Across the six sampled geographic locales (Figure 1), desiccation tolerance for both sexes increased positively with latitude (Figure 2).

These patterns were affected by culture temperature, demonstrating an inverse relationship between temperature and desiccation tolerance (Table 1; raw data are archived at Dryad doi: http://datadryad.org/review?doi=doi:10.5061/dryad.9719r). The significant interaction terms of population by temperature and temperature by sex showed that patterns of thermal plasticity varied among populations and were distinct between the sexes (Table 1; Figure 2). As culture temperature decreased, in general the magnitude of differentiation among the geographic populations increased. The pattern of population-specific developmental plasticity was qualitatively similar across sexes, but males exhibited a slightly more pronounced response to developmental temperature. Likewise, we observed a strong, positive correlation between desiccation tolerance and latitude in adult females that were exposed to environmental conditions that elicit reproductive dormancy (Figure 2). This may be driven by geographic variation in the incidence of dormancy (Schmidt, Matzkin, Ippolito, & Eanes, 2005), but does suggest that desiccation tolerance may also exhibit short-term physiological plasticity.

Regression analysis of desiccation tolerance with latitude and associated climatic variables (\( T_{\text{min}}, T_{\text{max}}, T_{\text{ave}}, R_{\text{Hmin}}, R_{\text{Hmax}} \) and \( R_{\text{ave}} \)) for both sexes at all three culture temperatures is given in Table 2. In an interesting manner, the temperature parameters exhibited stronger associations with desiccation tolerance than did those for relative humidity. In particular, minimum temperature (\( T_{\text{min}} \)) was most strongly associated with the observed variation in desiccation tolerance. To further explore the collinearity of thermal plasticity in desiccation tolerance and various climatic variables, we performed a principal component analysis (Supporting Information Figure S3). The first ordination axis explained 65.94% of variability, and the second axis explained 27.14%. Temperature correlated best with the first axis.

Seasonal variation in desiccation tolerance for the three sampled seasonal populations (MA, PA and VA) is depicted in Figure 3. Again,
Note. Populations were cultured at three different growth temperatures (18, 25 & 29°C). Bold indicates statistical significant (p < 0.05).

...we observed significant variation across the geographically distinct locales and predictable differences in tolerance between the sexes (Table 3a). While the effect of season was not significant, the interaction terms incorporating season were significant or approached significance (Table 3a). To further investigate this, we also analysed the sexes separately (Table 3b). Tolerance for females did not vary between time points one and two as well as between three and four (Table 4; Figure 4). Thus, in response to field culture under heterogeneous environmental conditions, desiccation tolerance in outbred populations of Drosophila melanogaster evolved rapidly over a period of approximately 90 days (~9 generations).

## 3.3 Water loss rate and metabolic rate in natural populations

With the exception of HFL and ME males, we did not find a significant within-sex difference in mean body mass among all our comparisons (data not shown). The mean body mass of male and female flies (ca. 0.7–0.8 and 1.1–1.2 mg, respectively) correlated with the well-documented higher desiccation resistance of females (e.g., Gibbs et al., 1997). We found no significant difference for either sex in mean VCO₂ or water loss rates (WLRs) between HFL and ME, or the seasonal collections from Pennsylvania (ANOVA, with replication as a random factor, p > 0.05; see Supporting Information Table S2). When comparing HFL and ME males, no significant difference was observed for VCO₂ or WLRs.

### 3.2 Experimental evolution in the field

For both males and females, desiccation tolerance increased over time in the field mesocosms (Table 4; Figure 4). As the populations were sampled and then passed through three generations of common garden culture in the laboratory, the observed patterns of increased tolerance over time are inferred to reflect change in the genetic composition of the experimental populations in response to environmental conditions experienced in the field. In females, desiccation tolerance increased rapidly between the first and second sampling time points (a period of 25 days), remained constant over the next interval (23 days), then exhibited a slight decrease at the last sampling time point (29 days). In contrast, desiccation tolerance of males increased at a fairly constant rate; tolerance increased between time points one and two as well as between three and four (Table 4; Figure 4). Thus, in response to field culture under heterogeneous environmental conditions, desiccation tolerance in outbred populations of Drosophila melanogaster evolved rapidly over a period of approximately 90 days (~9 generations).

### Table 2 Regression analysis of desiccation tolerance with geographical and climatic parameters

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<td>−0.9651</td>
<td>−0.977</td>
<td>−0.6814</td>
<td>−0.5908</td>
<td>−0.7623</td>
</tr>
<tr>
<td>p</td>
<td>0.00053</td>
<td>0.00325</td>
<td>0.0018</td>
<td>0.0079</td>
<td>0.13612</td>
<td>0.2169</td>
<td>0.07804</td>
</tr>
<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>r</td>
<td>0.87516</td>
<td>−0.8453</td>
<td>−0.8305</td>
<td>−0.8544</td>
<td>−0.6351</td>
<td>−0.4225</td>
<td>−0.5673</td>
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<tr>
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<td>0.03405</td>
<td>0.04064</td>
<td>0.03028</td>
<td>0.17541</td>
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<td>0.24034</td>
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<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>r</td>
<td>0.78074</td>
<td>−0.82</td>
<td>−0.8713</td>
<td>−0.8602</td>
<td>−0.3712</td>
<td>−0.3493</td>
<td>−0.4461</td>
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<tr>
<td>p</td>
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<td>0.04567</td>
<td>0.02377</td>
<td>0.02797</td>
<td>0.46884</td>
<td>0.4974</td>
<td>0.37529</td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>r</td>
<td>0.76591</td>
<td>−0.8095</td>
<td>−0.6357</td>
<td>−0.742</td>
<td>−0.7053</td>
<td>0.06338</td>
<td>−0.3727</td>
</tr>
<tr>
<td>p</td>
<td>0.07579</td>
<td>0.05098</td>
<td>0.17491</td>
<td>0.09126</td>
<td>0.11748</td>
<td>0.90505</td>
<td>0.46687</td>
</tr>
<tr>
<td>29°C female</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>r</td>
<td>0.86563</td>
<td>−0.8094</td>
<td>−0.7261</td>
<td>−0.7852</td>
<td>−0.7537</td>
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<td>0.10226</td>
<td>0.06426</td>
<td>0.08353</td>
<td>0.46788</td>
<td>0.11146</td>
</tr>
</tbody>
</table>

Note. The first row gives correlations (r values); the second line the goodness of fit from ANOVA. Statistical significance is depicted by bold typeface.
found in mean \( VCO_2 \) (\( F_{1,7} = 1.61, p = 0.24 \)) or water loss rates (\( F_{1,7} = 2.76, p = 0.21 \)), even when accounting for differences in body size (ANCOVA, with body mass as a covariate).

The limitation of our original respirometry approach in detecting variation in WLRs prompted us to focus on the two most geographically distant populations for which significant variation in desiccation resistance was detected (i.e., HFL vs. ME; see Figure 2), using an alternative approach measuring WLR over a longer time frame (see Section 2). Still, no significant difference was found in the WLRs among female (ANOVA; \( F_{1,21} = 0.005, p = 0.95 \)) or male (\( F_{1,21} = 0.13, p = 0.71 \)) flies even when exposed to dry air flow for 3 hr. No significant differences were found in \( VCO_2 \) between HFL and ME females (\( p = 0.68 \) and 0.34 after 30 min and 3 hr, respectively), but similar values for males after 30 min (\( p = 0.46 \)) were followed by significantly lower \( VCO_2 \) values for ME compared with HFL males (\( F_{1,21} = 4.34, p = 0.049 \)). In an interesting manner, when testing for temporal changes within sets of flies we found a significant increase in \( VCO_2 \) of HFL males from initial values (3.16 ± 0.15 μl fly\(^{-1}\) hr\(^{-1}\)) compared to those recorded after 3 hr of exposure to desiccation (3.82 ± 0.13 μl fly\(^{-1}\) hr\(^{-1}\)) (paired \( t \) test; \( t_{11} = 5.04, p < 0.001 \)). In contrast, values after 30 min and 3 hr of desiccation did not vary significantly for ME male flies (\( t_{11} = 1.89, p = 0.09 \)) (Figure 5a), which could reflect their enhanced desiccation tolerance (Figure 2). Among female flies, a significant decrease in \( VCO_2 \) during exposure to desiccation was recorded for ME (\( t_{11} = 3.69, p = 0.004 \)), but not HFL populations (\( t_{11} = 0.40, p = 0.70 \)) (Figure 5b).

### 3.4 GWAS & GO enrichment

We observed considerable genetic variability in desiccation tolerance among the 175 tested DGRP lines (Supporting Information Table S3). Line means are depicted in Figure 6. Females survived longer than males under desiccating conditions, as expected. There was significant heterogeneity among lines with respect to sex-specific differences and relative ranking (Figure 6). Neither Wolbachia infection status nor the presence of chromosomal inversions (Huang et al., 2014) had any effect on desiccation tolerance (Table 5; Supporting Information Tables S3 and S4).

<table>
<thead>
<tr>
<th>TABLE 3</th>
<th>ANOVA for seasonal variations in desiccation tolerance of Drosophila melanogaster adults from the three geographically distinct populations</th>
</tr>
</thead>
<tbody>
<tr>
<td>(a) Parameters</td>
<td>SS</td>
</tr>
<tr>
<td>Population (Pop)</td>
<td>109.66</td>
</tr>
<tr>
<td>Season (Sea)</td>
<td>11.06</td>
</tr>
<tr>
<td>Sex</td>
<td>6,188.14</td>
</tr>
<tr>
<td>Pop*Sea</td>
<td>21.49</td>
</tr>
<tr>
<td>Pop*Sex</td>
<td>133.73</td>
</tr>
<tr>
<td>Sea*Sex</td>
<td>14.78</td>
</tr>
<tr>
<td>Pop<em>Sea</em>Sex</td>
<td>67.72</td>
</tr>
<tr>
<td>Error</td>
<td>1,034.62</td>
</tr>
</tbody>
</table>

| (b) Parameters | Male | | Female | |
| | SS | df | MS | \( F \) | \( p \) | SS | MS | \( F \) | \( p \) |
| Population (Pop) | 130.11 | 2 | 65.05 | 34.53 | <0.0001 | 113.29 | 56.64 | 10.38 | <0.0001 |
| Season (Sea) | 25.71 | 1 | 25.71 | 13.64 | 0.000315 | 0.13 | 0.13 | 0.025 | 0.875397 |
| Pop*Sea | 75.65 | 2 | 37.83 | 19.87 | <0.0001 | 13.55 | 6.78 | 1.243 | 0.291793 |
| Error | 265.58 | 141 | 1.88 | | | 769.03 | 5.45 | | |

Note. Sex was included as a predictor in (a); and in (b), males and females were analysed separately.
Using among-line variation in the standard mapping pipeline (Mackay et al., 2012), we identified 25 SNPs that were significantly associated with desiccation tolerance; these SNPs were associated with 20 genes and one noncoding locus (Supporting Information Tables S5 and S6). Gene ontology (GO) enrichment analysis placed these 20 genes in various GO biological categories that broadly included metabolism, signalling, cuticle development and transport. However, no enrichment for any biological category was observed (Eden, Navon, Steinfeld, Lipson, & Yakhini, 2009); this was expected, given the small number of genes identified. We did not observe any overlap at the gene level between our association study using the DGRP panel and two studies conducted on D. melanogaster populations from Australia (see table 1 in Telonis-Scott et al., 2016; see also Telonis-Scott et al., 2012).

### 3.5 Clinal differentiation of SNPs associated with desiccation tolerance

Among North American populations, the genes we identified as being associated with desiccation tolerance in the DGRP did not show increased signatures of spatially varying selection relative to with 20 genes and one noncoding locus (Supporting Information Tables S5 and S6). Gene ontology (GO) enrichment analysis placed these 20 genes in various GO biological categories that broadly included metabolism, signalling, cuticle development and transport. However, no enrichment for any biological category was observed (Eden, Navon, Steinfeld, Lipson, & Yakhini, 2009); this was expected, given the small number of genes identified. We did not observe any overlap at the gene level between our association study using the DGRP panel and two studies conducted on D. melanogaster populations from Australia (see table 1 in Telonis-Scott et al., 2016; see also Telonis-Scott et al., 2012).

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the rest of the genome (Figure 7a; \( p = 0.769 \)). Although there has not been any latitudinal cline reported for desiccation tolerance in Australian \( D.\ melanogaster \) (Hoffmann, Hallas, Sinclair, & Mitrovski, 2001; Hoffmann, Shirriffs, & Scott, 2005), we observed an indication of increased spatial genetic differentiation at the loci identified in the DGRP (Figure 7c; \( p = 0.0926 \)) among Australian populations. This prompted us to examine per-gene \( z_{st}(g) \) values relative to sets of genomic matched control genes. Several of the candidate genes (\( Whamy, \ CG18302, \ CG5953 \)) for desiccation tolerance had relatively high \( z_{st}(g) \) values, indicated by low bootstrap \( p \)-values, in the Australian populations (Figure 7d). However, none of bootstrap \( p \)-values for these genes fell below the Bonferroni correction threshold at \( \alpha < 0.05 \).

4 | DISCUSSION

Desiccation resistance in insects can involve one or more physiological mechanisms, including (a) increases in total body water content and/or in haemolymph volume (Folk, Han, & Bradley, 2001); (b) increased dehydration tolerance (i.e., tolerance of body water loss before death; Telonis-Scott et al., 2006); (c) reduction in the rate of water loss (Gibbs et al., 1997); or (d) any modification of behaviour, including changes in activity level and associated metabolism. Body size plays a significant role in desiccation resistance: In 20 examined \( Drosophila \) species, desiccation tolerance varies fivefold among taxa and is associated with interspecific variation in size (Matzkin et al., 2009). Along the coast of eastern Australia, no cline was observed for desiccation tolerance in \( Drosophila\ melanogaster \) (Hoffmann et al., 2001), whereas along the Indian latitudes a robust cline has been observed (Karan et al., 1998). In an interesting manner, a cline for body size (e.g., thorax and wing length) has been observed on both continents (Bhan, Parkash, & Aggarwal, 2014; James, Azevedo, & Partridge, 1995). In North America, body size also varies clinally and increases with increasing latitude (e.g., Coyne & Milstead, 1987). Here, we observed a significant but shallow cline in desiccation tolerance at normal assay temperatures of 25°C (Figure 2). While body size may play a role in desiccation tolerance, the relationship between variation in body size and desiccation tolerance among \( D.\ melanogaster \) populations appears complex (Rajpurohit et al., 2016).

4.1 | Spatiotemporal patterns in desiccation tolerance

Our results, demonstrating a positive cline in desiccation tolerance for populations from the east coast of the United States, are

TABLE 5 ANOVA on desiccation tolerance in the DGRP lines

<table>
<thead>
<tr>
<th>Parameters</th>
<th>SS</th>
<th>df</th>
<th>MS</th>
<th>F</th>
<th>( p )</th>
</tr>
</thead>
<tbody>
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<td>548.29</td>
<td>114.702</td>
<td>10^{-6}</td>
</tr>
<tr>
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<td>0.08</td>
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<tr>
<td>Inversions</td>
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<td>1</td>
<td>7.27</td>
<td>1.522</td>
<td>0.218271</td>
</tr>
<tr>
<td>Sex*Wolbachia</td>
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<td>1</td>
<td>0.05</td>
<td>0.011</td>
<td>0.917563</td>
</tr>
<tr>
<td>Sex*Inversions</td>
<td>0.05</td>
<td>1</td>
<td>0.05</td>
<td>0.011</td>
<td>0.915927</td>
</tr>
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<td>1.14</td>
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<td>0.625584</td>
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<tr>
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<td>1.22</td>
<td>1</td>
<td>1.22</td>
<td>0.254</td>
<td>0.614309</td>
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<tr>
<td>Error</td>
<td>1,510.53</td>
<td>316</td>
<td>4.78</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Note. The Wolbachia infection and inversion status of DGRP lines have also been incorporated in the full model. Sex-specific analyses are presented in the supplementary material (Supporting Information Table S4).
consistent with previous studies showing higher desiccation tolerance in temperate vs. tropical locales (Hoffmann & Harshmann, 1999). On the Indian subcontinent, this trend was clearly observed in multiple Drosophila species (Karan et al., 1998; Parkash et al., 2008; see also Rajpurohit et al., 2017b), where parallel clines for this trait have been observed (Rajpurohit, Nedved, et al., 2013). In India, higher latitudes are characterized by lower temperature and lower humidity during winter, whereas low latitude, southern locations are warm and humid for most of the year. A meta-analysis approach concluded that the coefficient of variance in temperature was a major climatic component to support the observed parallel clines for desiccation tolerance in several Drosophila species on the Indian subcontinent (Rajpurohit, Nedved, et al., 2013). Along the east coast of the United States, more temperate populations experience greater environmental fluctuations associated with seasonality and harsher winter conditions. As adult overwintering is associated with long-term exposure to a desiccating environment, we predicted that populations collected in the spring would be characterized by relatively higher desiccation tolerance, similar to patterns observed for other stress-related traits (Behrman et al., 2015). However, seasonal variation in relative humidity is not pronounced at the sites we sampled (e.g., Supporting Information Figure S4). We did not observe robust differences in desiccation tolerance between early and late season collections of D. melanogaster from three temperate populations spanning 38–42°N latitude; only males exhibited seasonal variation in tolerance, and this pattern was pronounced in only one of the three sampled geographic locales.

Although humidity may not vary significantly among seasons (Supporting Information Figure S4), the regulation of water balance in natural populations may be driven primarily by temperature, or the interaction between temperature and humidity (Kingsolver, 1979; Rajpurohit, Nedved, et al., 2013). In the field mesocosms, relative humidity was highly variable over diurnal timescales, but average humidity was not distinct among sampling time points. Temperature, in contrast, varied predictably over time and was distinct across the sampled intervals from July to October. The patterns of desiccation tolerance observed in the field mesocosms mirror temperature profiles, with some deviation between males and females in the environmental tracking. The evolution of increased desiccation tolerance in the field experiment cannot be explained by response to humidity, as this was consistently higher in the field than under typical conditions of laboratory culture (Heerwaarden & Sgro, 2014).

4.2 Climatic associations

It is generally assumed that desiccation tolerance is selected during hot and dry conditions, as heat and desiccation stresses often co-occur (Hoffmann & Parsons, 1991). In natural habitats on the US east coast, however, such is not the case. Variation in desiccation tolerance is most strongly associated with minimum temperature (and length of the winter season) of the geographic origin of our collections, suggesting that desiccation tolerance in temperate populations may be favoured in environments characterized by cold and dry conditions (Leather, Walters, & Bale, 1993). Cold and desiccation tolerance may also exhibit correlated responses (e.g., Bubliy & Loeschcke, 2005; MacMillan, Walsh, & Sinclair, 2009). Thus, it remains unclear whether the latitudinal patterns we observed are driven primarily by selection directly on desiccation tolerance or may reflect an indirect response due to selection on a correlated trait. In contrast to the observed patterns for the geographic collections, our field mesocosm experimental data indicate a role of higher temperature, where desiccation tolerance increases as temperature increases. These contrasting patterns may reflect complex dynamics associated with various spatial and temporal scales, as well as the aforementioned correlations between desiccation tolerance and other traits. It would be of particular interest to generate long-term longitudinal data on desiccation tolerance in natural and experimental populations to further explore the associations between tolerance and various climatic parameters.
4.3 | Thermal plasticity

We observed that decreasing culture temperature in the laboratory resulted in elevated desiccation tolerance. A recent study on the cold-adapted *D. nepalensis* from the western Himalayas found that flies grown at 15°C demonstrate a twofold increase in body size, a greater degree of melanization, higher desiccation resistance, and increased haemolymph and carbohydrate content as compared to flies reared at 25°C (Parkash, Lambhod, & Singh, 2014). There is a strong possibility that *D. melanogaster* populations growing at lower temperatures may also exhibit these plastic responses that could subsequently affect desiccation tolerance. We did observe that variation among populations became exacerbated at lower temperatures and was most distinct following exposure to dormancy-inducing conditions. Thus, our data also suggest that patterns of plasticity in *D. melanogaster* may vary predictably among natural populations and habitats.

4.4 | Geographic variation in metabolic rate

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

We found no evidence for variation in water loss rates that could explain the observed clinal variation in desiccation tolerance. Water loss rates of the northernmost (ME) and southernmost (HFL) populations did not differ in either males or females. We also recorded similar metabolic rates, expressed as CO2 emission rates (VCO2), for females from the two populations; however, higher values for severely desiccated HFL male flies suggest significant differences in the metabolic response of these populations to prolonged exposure to desiccating conditions. This is in agreement with both intraspecific (Gibbs & Gefen, 2009; Hoffmann & Parsons, 1993) and interspecific (Gibbs et al., 2003) reports that demonstrated desiccation resistance in *Drosophila* is correlated with reduced activity under stressful conditions; these results are also consistent with pronounced differences in central metabolism between populations at the geographic extremes of the US east coast (e.g., Flowers et al., 2007; Lavington et al., 2014; Verrelli & Eanes, 2001). It should be noted that increasing VCO2 values could reflect a switch to carbohydrate catabolism under desiccation stress (Marron, Markow, Kain, & Gibbs, 2003), independent of changing metabolic rates. However, we did not observe a similar response in females. Instead, the significant decrease in VCO2 in the more resistant ME females as they settled to the dry metabolic chamber environment, and absence of this response for HFL females, suggests a difference in behavioural response. Both males and females in these populations do exhibit very distinct behaviour in response to thermal variation as well (Rajpurohit & Schmidt, 2016).

Variation in activity patterns under stressful conditions is likely to result in correlated differences in respiratory water losses. The similar WLRs reported here for HFL and ME flies can be explained by the considerably higher relative importance of cuticular water loss in insects (Chown, 2002) and may suggest that flies across the experimental populations do not vary in their cuticular resistance to water loss. Nevertheless, while results in this study do not confirm an effect of activity level on WLRs and thus on desiccation resistance, they could well reflect how stressful the exposure to experimental desiccation is to flies from the respective populations. If the more susceptible HFL flies have lower body water contents when hydrated compared with the more resistant ME flies (Chippendale et al., 1998; Folk et al., 2001; Gefen et al., 2006), then at similar WLRs the former would approach the minimum tolerable hydration state earlier, which could elicit an increase in activity levels as a result of attempts to seek more favourable conditions. In addition, a delayed escape response in the more resistant flies could indicate higher dehydration tolerance that would trigger an escape response at lower body water content. We speculate that geographically distinct populations exhibit distinct behaviours in response to desiccation exposure. This hypothesis needs to be tested in future work.

4.5 | Candidate genes

The GWAS of the DGRP panel identified 20 genes associated with the sex-averaged among-line variance in desiccation tolerance. A GOWINDA analysis using 103 SNPs with sex-averaged or sex-specific associations found no significant GO term enrichment (lowest FDR >0.4). While there was no associated ontology enrichment, these identified genes do represent candidates for subsequent functional analysis. For example, 5-HT2 is a 5-hydroxytryptamine (5-HT, serotonin) receptor, and serotonin is an important diuretic hormone in insects (Martini et al., 2004). Two additional candidates (mg1, CG4835) are functionally linked to chitin metabolism. Serotonin also mediates cuticular plasticization (Reynolds, 1975); together, these findings suggest that cuticular permeability may be an important factor affecting desiccation tolerance. Metabolic differences also appear to be important, as SNPs mapped to or close to three genes implicated in lipid metabolism (klar, CG18302, CG31089). However, we emphasize that the identification of these genes is based on GWA of a single panel of inbred lines and that the generation of a comprehensive list of genes associated with variance in desiccation tolerance would require further investigation as well as integration across data sets and experimental approaches.

4.6 | Ecological genetics of desiccation tolerance

Relative to other fitness-associated traits in North American populations (e.g., body size (Coyne & Milstead, 1987); reproductive
dormancy (Schmidt et al., 2005); cuticular hydrocarbons (Rajpurohit et al., 2017a); and thermal preference (Rajpurohit & Schmidt, 2016), we observed a shallow cline for desiccation tolerance across the sampled latitudinal gradient in eastern North America. Our results also demonstrated pronounced patterns of plasticity in response to temperature, both in terms of developmental plasticity as well as adult acclimation and subsequent response. However, the observed patterns suggest that spatially varying selection may be less pronounced on this trait, both in comparison with other traits in North American populations as well as to desiccation tolerance in Australian (Telonis-Scott et al., 2006) and Indian (Karan et al., 1998; Parkash et al., 2008; Rajpurohit, Oliveira, et al., 2013) populations. Our analysis of clinal enrichment demonstrated that genes associated with desiccation tolerance are more strongly enriched for clinality in Australia than in North America. However, it is clear that there is no robust signal, above expectation, that genes associated with desiccation tolerance evidence patterns consistent with spatially variable selection. This may reflect the weak associations between latitude and desiccation tolerance, limitations in the identification of genes underlying variance in the trait, or a combination of the two.

5 | CONCLUSION

We observed a shallow cline for desiccation tolerance in populations sampled along the latitudinal gradient in the eastern United States; these patterns of variation among populations exhibited both developmental and adult plasticity, suggesting that further analysis of desiccation tolerance should be examined under a range of environmental conditions. Climatic analysis of this cline indicated that observed patterns of desiccation tolerance were most strongly associated with lower temperature conditions, suggesting that selection on this trait in temperate populations may be associated with response to desiccating conditions that co-occur with long-term exposure to reduced temperatures. However, our field experiment also demonstrated that desiccation tolerance increases over short timescales under field conditions of increasing temperature, suggesting that the evolutionary dynamics of desiccation tolerance may vary over distinct temporal and spatial scales. GWAS of the DGRP panel of inbred lines identified 20 genes and one non-protein-coding locus associated with desiccation tolerance in North American populations; although underpowered and of perhaps limited scope in associations with patterns in natural populations or in the field, this does provide candidates for subsequent functional analysis and investigation. Overall, patterns of variation for desiccation tolerance appear complex and more fine scaled, mechanistic dissections should shed further light on the evolutionary and ecological dynamics of this trait in natural populations.

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AUTHOR CONTRIBUTIONS


DATA ACCESSIBILITY

All raw data, including data not presented in the paper, are archived at Dryad. https://doi.org/10.5061/dryad.9719r.


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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article.