# A multi-dimensional selective landscape drives adaptive divergence between and within closely related *Phlox* species

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#### 30 Abstract

Natural selection shapes diversity across micro and macro-evolutionary scales. Selection 31 causes local adaptation across populations within species and is simultaneously responsible for 32 33 much of the divergence between species. However, it is unclear if either the force of or the response to selection is the same or different across these scales. Here we show that natural 34 35 selection drives adaptive divergence between closely related species in a pattern that is distinct from local adaptation within a species. We use reciprocal common-garden transplant 36 experiments across three species of *Phlox* wildflowers to show widespread adaptive divergence 37 38 causing species to have highest fitness in their native habitats. Using provenance trials, we also find that one of these *Phlox* species has strong local adaptation between populations. We 39 40 compare the axes of divergence and selection between these two scales of diversity and discover that one suite of traits predicts fitness differences between species and that an independent suite 41 of traits predicts fitness variation within species across individuals. Our work reveals how forces 42 of selection can both drive key divergence between species, allowing for and causing speciation, 43 while simultaneously causing extensive diversity that is maintained across populations within a 44 species. The selection landscape is complex and multidimensional 45

#### 46 Introduction

Ecological adaptation contributes to the origin and maintenance of biodiversity (1-4).
Evolution by natural selection drives local adaptation among populations within a species that
occupy different biotic and abiotic environmental conditions (5-8). Similarly, strong ecological
selection can drive divergence between closely related species and cause reproductive isolation,
leading to ecological speciation (1, 9, 10). Although extensive research has investigated local
adaptation within species and investigated ecological divergence between species, little is known

about if and how these evolutionary processes are related. Are the axes of selection favoring
adaptive divergence between species the same or different than the axes of selection favoring
local adaptation within a species? Characterizing local adaptation both within and between
closely related species can offer insight into how ecological adaptation generates diversity from
the micro- to macroevolutionary scale.

58 Disparate populations within wide-ranging species often evolve to become adapted to the 59 local ecological conditions (7). Across the tree of life there are striking examples of variation in morphology, physiology, and phenology within species across populations that span significant 60 61 gradients of temperature, water availability, seasonality, and types of biotic interactions (11-16). Local adaptation among populations within a species is not inevitable, and depends on the 62 63 relative strengths of selection and migration (17), the presence or absence of fitness trade-offs in different habitats (6), and the genetic correlation between traits (18-20). Nevertheless, meta-64 65 analyses of the local adaptation studies conclude that adaptive divergence between populations 66 of a species is widespread in nature (6, 21).

As taxa become reproductively isolated, they tend to evolve suites of diverged traits that 67 lead to higher fitness (survival or reproduction) in local or native habitats compared to habitats of 68 69 closely related taxa. Adaptive divergence often leads to a similar pattern of reciprocal local vs. foreign advantage between closely related species as the pattern that we see between locally 70 71 adapted populations within a single species. Even when closely related species are in broad 72 sympatry with extensive geographic overlap, we may expect a pattern of adaptive divergence. Interspecific competition for resources can select for ecological divergence and niche 73 74 partitioning driving either species-wide patterns of differentiation or leading to patterns of 75 character displacement in sympatry (2, 22). Therefore, we expect many of the traits that

differentiate species from each other to be the suites of traits that lead to differential fitness and thus ecological reproductive isolation between the species. In this way, adaptive divergence not only causes phenotypic differentiation between taxa but also contributes to the cessation of gene flow between taxa, leading to speciation. For this reason, ecological adaptation is considered important during the speciation process. In fact, environmental divergence and ecological divergence are often added to, and portrayed as parallel to, the speciation continuum from no reproductive isolation to complete reproductive isolation (23, 24).

83 The idea of a continuum of adaptive ecological divergence -- from producing and 84 maintaining diversity within species to causing reproductive isolation between species -- invites us to consider how patterns of adaptation within and between species may or may not be related 85 86 (2, 25). Under one notion of this continuum, the adaptive divergence that we see between species is an extreme case of the local adaptation we see within species across populations and therefore 87 88 could be due to similar axis of selection and involve similar types of trait divergence (Fig. 1, top 89 & bottom right). Alternatively, the types of selection driving divergence between species could 90 be distinct from the selection pressures favoring local adaptation within a species (Fig. 1, top & bottom left). Under this latter scenario, the trait divergence that differentiates species is different 91 92 from the variation that we see within a species. Importantly, ecological divergence and 93 speciation unfolds over evolutionary time, and the snapshot of divergences we see now between 94 populations and species does not directly tell us about how the process of speciation did or will 95 proceed in this system (24). Nonetheless, comparing patterns of phenotypic divergence and axes 96 of selection across phylogenetic scales can help us understand how phenotypic diversity is 97 generated and maintained under different scales of geographic range and genetic exchange.

98 Phlox pilosa subsp. pilosa (hereafter "pilosa"), P. amoena subsp. amoena (amoena), and P. pilosa subsp. deamii (deamii) are three closely related perennial wildflower taxa inhabiting 99 100 the eastern U.S. that provide a promising system in which to evaluate patterns of ecological 101 differentiation, both within and between species (26). The ranges of these three Phlox taxa 102 overlap in western Kentucky, Tennessee, and Indiana, but they rarely co-occur in the same 103 locality, suggesting differences in habitat preference (26-28). Here, we use a combination of reciprocal transplant and provenance trial approaches to evaluate the presence and strength of 104 local adaptation between and within species. Since the foundational experiments of Turesson 105 106 (29), and the team of Clausen, Keck, and Heisey (30), reciprocal transplants have been the 107 standard for studying ecological differentiation within and between species. Provenance trials 108 can also be used to study local adaptation, in which the performance of a diverse panel 109 representing population variation throughout a species' range is evaluated in a common ecological setting (31-33). Here we: 1.) model and compare the ecological niches of the Phlox 110 111 species; 2.) determine whether there is adaptive divergence between the three species; 3.) infer if 112 there is local adaptation within *Phlox* species; and 4.) evaluate patterns of phenotypic diversity 113 across all three species and compare axes of selection driving divergence between and within 114 species. Collectively, this study provides unique insights into how selection operates to drive 115 diversity across scales of micro- and macro-evolution.

116

117 **Results** 

118 *Ecological niche modeling* 

We built robust ecological niche models for the two widespread *Phlox* species, amoenaand pilosa, using available occurrence data and biologically relevant environmental variables

(Fig. 2A, Fig S1, TableS1 & S2). The predicted extents of suitable habitat conform well to the
described geographic ranges of these species with a broad range of sympatry from Georgia to
Kentucky (27, 28). We cannot build an ecological niche model for the third species, deamii, due
to the low number of known occurrence points. It is hypothesized that deamii populations
experience a narrow range of environmental conditions and are broadly sympatric with both
amoena and pilosa (27).

From a principal component analysis (PCA) of the environmental variables used to build 127 128 our niche models, we find that pilosa inhabits a greater breadth of ecological variation than does 129 amoena (Fig. 2B). While both species occupy a similar amount of variation on PC2, amoena 130 occupies a subset of the variation covered by pilosa on PC1. We find that the median conditions 131 occupied by amoena and pilosa are significantly different on PC1 but not on PC2 (Fig S1C.). Of 132 note, the common garden sites chosen to represent amoena and pilosa habitats in our reciprocal 133 transplant experiment described below differ on PC1 as well (colored diamonds in Fig. 2B). The 134 reciprocal transplant experiment includes individuals sampled from populations that reasonably 135 encompass the environmental variation experienced by these species (black edged circles Fig. 2B, Table S5). 136

137

#### 138 Adaptive divergence between taxa

We find strong evidence of adaptive divergence between *Phlox* species from our
reciprocal transplant experiments. Multiple individuals sampled from source populations
throughout the ranges of these three perennial *Phlox* species (black diamonds and circles Fig.
2A) were clonally replicated into common gardens in the native sympatric range of these species.
Our experiment included three garden sites each adjacent to a wild population of one of the focal

144 taxa (Fig. 2C, D & E, Table S6). We quantified five fitness-related traits: herbivory, fruit number, flower number, biomass, and survival and find that a significant taxon-by-garden 145 146 interaction predicts trait values, indicating that the relative success of the species depends on the 147 garden in which they are grown (Fig. 3; Table 1). Adaptive divergence is evidenced by either the 148 local species having higher fitness than the foreign species in the local species' garden, or by a 149 focal species having highest fitness in its home garden compared to all other away gardens. 150 All significant local vs. foreign comparisons match the prediction of adaptive divergence 151 between taxa with the local taxon outperforming the foreign taxa (Table 1). In the amoena 152 habitat, amoena had higher survival and experienced significantly less herbivory than both pilosa 153 and deamii, and produced more fruits than pilosa plants. In the deamii habitat, deamii had 154 significantly higher survival than pilosa. In the pilosa habitat, pilosa plants produced 155 significantly more fruits, and survived at a higher rate than both amoena and deamii plants. Home vs. away comparisons (comparing across habitats for each taxon) showed some 156 157 significant differences in the direction predicted by local adaptation (Table 1). Pilosa had the 158 highest fitness in the home garden compared to in the other gardens on all five fitness traits. 159 Deamii had less herbivory and set more fruits in the home garden compared to the amoena 160 garden. We also found some patterns of success that did not indicate highest success at home-161 sites. For instance, deamii and amoena had fewer flowers and fruits in their home gardens 162 compared to either of the other gardens. 163 Due to limited availability, tests for local adaptation with deamii suffered from low power. We found some qualitative patterns consistent with local adaptation (Fig 3); notably, the 164 165 proportion of plants that survived to the end of the experiment matches the prediction of adaptive 166 differentiation for all local vs. foreign and home vs. away comparisons. If we consider post-hoc

167 comparisons just between amoena and pilosa (excluding deamii due to its lower sample size) a
168 clearer pattern of adaptive differentiation between these species emerges, with significant local
169 vs. foreign comparisons matching the prediction of local adaptation in herbivory, fruit number,
170 and survival (Table 1).

171

#### 172 Local adaptation within species

173 We find strong evidence of local adaptation across populations within pilosa. We used 174 statistical models to estimate the contribution of the source population to variation in the five 175 fitness-related traits for amoena and pilosa. We could not include deamii in this analysis because 176 there was insufficient variation across the three populations. Local adaptation was evidenced by 177 a negative relationship between the estimated population effect on fitness and distance of the 178 population from the common garden. This relationship was tested for geographic distance, genetic distance (as measured by F<sub>ST</sub> using data from Goulet-Scott et al. 2021 (26)), and 179 180 environmental distance (as measured in climate PC space) between populations (Table S3 & S4). 181 For the pilosa species within the pilosa garden, the population effect on flower and fruit 182 fitness traits is strongly negatively correlated with geographic distance, environmental distance, 183 and genetic distance, as predicted by local adaptation. Final biomass in pilosa also shows a strong negative correlation with geographic distance (Fig. 4; Table S4). In other words, 184 185 individuals from populations further away from the common garden did worse than individuals 186 from populations closer to the garden. In the amoena habitat, pilosa populations also show strong negative correlations between biomass and geographic distance, and between flower number and 187 188 all three distances (Table S4). In the amoena garden, pilosa populations show a strong positive 189 correlation between the proportion of plants without herbivore damage and both genetic and

190	geographic distance; this represents the only signal in our data that does not support local
191	adaptation. Within the deamii garden, pilosa populations show a negative correlation between the
192	number of flowers produced and geographic distance while amoena populations show negative
193	correlations between survival and both genetic and environmental distance. There was
194	insufficient variation in herbivory among amoena populations and survival among pilosa
195	populations to model population effects suggesting no evidence for local adaptation in these two
196	traits.
197	

### 198 Selection between and within species

199 Adaptive differentiation between *Phlox* species and local adaptation within species occur along different axes of variation in leaf morphology and physiology, likely driven by different 200 201 axes of selection. For each individual genotype used in the experiment, we measured or 202 calculated six phenotypic traits including: leaf length, leaf width, leaf length/width ratio, leaf 203 area, leaf chlorophyll content, and specific leaf area (SLA), and summarized phenotypic 204 variation in a principal components analysis (Fig. 5). For the subset of fitness traits that showed 205 evidence of adaptive divergence between species and local adaptation within pilosa species we 206 asked if phenotypic PC1 or PC2 predicted fitness variation between and within species in the 207 Pilosa-habitat common garden (Table 2).

The principal components analysis summarizing the phenotypic variation across species sharply divides pilosa from amoena and deamii individuals along PC1 (Fig. 5a.). In fact, taxon identity explains 72% of the variation along this first principal component (F(2, 318)=417.34, p<0.001). PC1 explains 45% of trait variation and describes leaf shape. Long narrow leaves and low chlorophyl content are at one end of the PC axis (pilosa-like), and short wide leaves with

high chlorophyl content are at the other (amoena-like). All species show extensive and
overlapping variation along PC2, which explains 27.8% of the phenotypic variation and
corresponds to variation in size of the leaf (area and SLA). Taxon identity explains none of the
variation along PC2 (F(2,318)=0.8, p=0.451).

217 We investigated how leaf trait variation (PC1 and PC2) explained variation in fitness 218 traits (fruit set, flower set, and biomass) in the pilosa garden both between and within species 219 using two sets of models. First, we modeled variation in fitness traits as explained by each trait 220 PC while controlling for taxon and the interaction between taxon and trait PC. For PC1, we 221 found that taxon identity predicted fitness related traits consistent with our tests of adaptive 222 divergence previously discussed. Due to the collinearity between taxon identity and value at 223 PC1, this trait PC is not significant in our model when controlling for taxon (Table S7). Pilosa 224 individuals have both higher values long PC1 and high fitness in the pilosa garden. For PC2 the interaction term in our model is significant indicating that the relationship between PC2 and 225 226 fitness traits varied across species (Table S7). With our second set of models, we evaluated how 227 each leaf trait PC predicts fitness traits across all the species and within each of the species 228 (Table S8). PC1 does not predict fitness variation within any of the three species; it is only when 229 individuals from all three species are included in the model that we see a significant relationship between PC1 and fitness-related traits (Fig. 5b.,d.,f., Table S8). In contrast, we find that within 230 231 pilosa and amoena PC2 strongly predicts fitness trait and that this variation explains the 232 significant relationship between PC2 and fitness in the combined dataset. (Fig. 5c.,e.,g., Table 233 S8) Together our models indicate that leaf shape (PC1) differs significantly between Phlox 234 species and it is therefore these differences that correspond to fitness differences between species

in the common garden. In contrast, leaf size (PC2) varies within species and significantly
predicts *within* species variation in fitness in the common garden (Table S8).

#### 237 Discussion

238 Natural selection is widely acknowledged as the most important force driving the 239 evolution of biological diversity, yet we still have much to learn about how this process acts 240 across micro- and macro-evolutionary scales. At one end of this scale, populations within a 241 species can locally adapt in response to variation in selection across space, generating diversity 242 within a species; while at the other end, response to selection can drive adaptive divergence 243 between taxa and even cause significant reproductive isolation, thus contributing to speciation. 244 Many studies have characterized the response to selection at a given scale, yet there are few 245 studies that integrate across scales to compare how selection drives divergence within and 246 between species.

247 Here we have characterized adaptive divergence between closely related species and 248 local adaptation within one of these species. Furthermore, we show that selection driving 249 divergence between species is distinct from, and orthogonal to, selection driving divergence 250 within species. Our results suggest a broadly applicable explanation of how a species can both 251 maintain extensive adaptive phenotypic variation across broad ecological habitats while 252 simultaneously maintaining distinct adaptive divergence from recently diverged taxa. Selection 253 acts along many axes and the axis correlated with reproductive isolation and species interaction 254 may be entirely different from the axis allowing populations to locally adapt across a species' 255 range.

Our results suggest that natural selection drove adaptive divergence between the three co occurring species of *Phlox* – pilosa, amoena, and deamii. The wide-spread species – pilosa and

amoena – show broadly sympatric ranges with statistically significant yet minimal niche
divergence. Specifically, after reducing dimensionality of multiple environmental and soil
variables, we find that amoena inhabits a distinct subset of the broader environmental tolerance
of pilosa. This likely reflects the more northern range limit of pilosa compared to amoena.
Nevertheless, all three species show significant overlap in both geographic and environmental
space. While our niche modeling suggests minimal adaptive divergence between species, our
experimental gardens reveal extensive fitness differences.

265 Across the five proxies for fitness we measured, we found that the local species 266 generally does better in its local habitat garden as would be predicted by adaptive divergence 267 between species. Because we measured five traits in three gardens across three species, we 268 performed abundant statistical tests to identify patterns of differential success, which likely have 269 led to some false positives. We focus not on the results from any specific test but instead on the robust pattern that, for each species, we found evidence for natural selection favoring the local 270 271 species. The specific patterns of adaptive divergence are different for each species, which is 272 consistent with other studies that find that different lineages locally adapt in different ways (15). 273 For example, in the amoena garden we found amoena had the highest survival across the three 274 year experiment. In this garden there was extensive large-mammal herbivory with nearly 50% of 275 the plants showing signs of severe damage, but amoena plants suffered the least damage. The 276 pilosa garden had the greatest sun exposure and the pilosa plants seem to exploit this light to 277 have the highest survival and set the most fruits. Yet, pilosa individuals also suffered the most 278 when grown in the other gardens, potentially in response to low-light availability. Although our 279 conclusions of adaptive differentiation are strongly supported, this work inspires future 280 investigations to untangle the specific selective agents and traits underlying this pattern.

The support for adaptive differentiation between species may be particularly surprising given that the three common gardens were geographically close (within 120 km of each other) but the individuals in the garden were sourced from across the country, spanning 900 km. The patterns of adaptation were robust to the extensive geographic sampling and the breadth of source environmental conditions. This suggests that the traits that adaptively differentiate the species are shared across populations within their ranges and could therefore contribute to ecological reproductive isolation between species.

As is often observed for widespread species, one of our *Phlox* species also shows strong 288 289 patterns of local adaptation among populations. Two of our *Phlox* species span extensive 290 environmental gradients with large (and overlapping) geographic ranges. This presents the 291 opportunity for selection to favor different trait values between, for example the warm and dry 292 habitats in northern Florida and the cooler and wetter populations in western Kentucky. If local 293 adaptation within species is driven by these ecological gradients across their ranges, then we 294 predict that as distance increases between population source and an experimental garden, fitness 295 will decrease. This is precisely the pattern we documented across pilosa populations. Individuals 296 sourced from populations near the pilosa experimental garden grew bigger, had more flowers, 297 and set more fruits than individuals from populations farther way from the experimental garden. 298 This signal was robust to various measures of distance including geographic distance, 299 environmental distance, and genetic distance. Interestingly, this negative relationship between 300 fitness measures and distance from garden is also strong for pilosa individuals grown in the 301 amoena-habitat garden. As noted above, all pilosa individuals do worse in the amoena garden 302 than in the pilosa garden, but pilosa individuals sourced from farther away do even worse in the 303 amoena garden than individuals sourced from nearby. We interpret this pattern to mean that there

are traits that make pilosa individuals more fit in the northern part of the range, where the
common gardens were located, as compared to the southern part and that this difference in
success is independent of precise location (i.e., adjacent to a wild pilosa population or a wild
amoena population).

308 Interestingly, we found little to no signal of local adaptation in amoena populations. We 309 hypothesize that this difference in degree of adaptation within pilosa and amoena species could 310 reflect either differences in migration or selection. High migration between populations of 311 amoena could cause homogenization of genetic variation across the range and swamping of 312 locally adapted alleles. This is unlikely to explain the difference in pattern between species since 313 the range of genetic distances ( $F_{ST}$ ) represented in our experiment were similar for amoena (0 – 314 (0.44) and pilosa (0 - 0.46) and for a given distance between populations,  $F_{ST}$  is actually higher 315 for amoena than pilosa (Goulet-Scott et al. 2021; Fig. 4B (26)). In contrast, the range of 316 environmental distances (based on a PCA of environmental variables) represented in our 317 experiment was significantly less for amoena (0 - 3.48) than for pilosa (0 - 5.53). Therefore, 318 pilosa populations may face stronger selection throughout their range to adapt to local ecological conditions. 319

By characterizing adaptive divergence between species and local adaptation within a species, we can compare and contrast how natural selection generates diversity across these scales. We quantify diversity in leaf morphological and physiological traits across species and find that orthogonal axes of diversity predict fitness between species versus within species. These three species of *Phlox* grow in close geographic proximity, share pollinators, and have similar flower shape, size, color and timing. Therefore, their major phenotypic axis of diversity is in vegetative traits such as leaf morphology. Pilosa plants have long narrow leaves whereas

amoena plants have shorter, wider leaves. Both species show extensive variation in the overallsize and mass of the leaf.

Our results demonstrate that the major leaf-trait differences between species strongly predict fitness variation across species in our common-garden experiments. Plants with wider, shorter leaves do better in the amoena garden and plants with longer narrower leaves do better in the pilosa garden. It is perhaps unsurprising, that the traits that phenotypically differentiate species also predict fitness differences across the species' habitats. We have highlighted the link between key traits that define and differentiate closely related species, and fitness differences between species in their respective habitats.

This axis of phenotypic variation differentiating species (PC1) does not predict fitness variation within a species; instead, orthogonal trait variation (PC2) predicts within-species relative success. We found evidence of local adaptation across multiple proxies of fitness in pilosa. For all of these proxies we find that success of an individual is strongly predicted by size of the leaf (PC2) but not the shape of the leaf (PC1). We also find that leaf-trait PC2 value predicts fitness in amoena plants, further supporting the notion that different traits predict success within and between species.

The observation that adaptation within and between species operates along different axes of selection, might seem surprising given the perspective of a continuum of divergence between locally adapted populations and ecologically isolated species. The ecological speciation hypothesis suggests that populations within a species diverge ecologically until those populations evolve sufficient reproductive isolation and become distinct species. What we have demonstrated is that the process is multi-dimensional: if ecological divergence along one axis leads to reproductive isolation and a signature of local adaptation between lineages, then local adaptation

350 between populations within each lineage may persist or develop along other ecological axes. As has been articulated by others (24), the process of speciation is complex and not linear; similarly, 351 352 the role of selection in driving divergence is also complex and multi-dimensional. 353 Further research is needed to determine if different ecological factors are more or less 354 likely to drive between or within species divergence. For example, adaptive divergence driven by 355 ecological factors with discrete or step-like variation may be more likely to contribute to 356 reproductive isolation between species due to the absence of intermediate habitat that could be 357 suitable for hybrids (9, 34). In contrast, local adaptation to ecological factors that vary more 358 continuously may be less likely to lead to reproductive isolation and therefore act among 359 populations within species. Selective landscapes are clearly multifaceted; our study showcases 360 this by demonstrating that different ecological forces generate divergence between closely 361 related species than among populations within a species. 362

#### 363 Methods

364 *Ecological niche modeling* 

We used ecological niche modeling to assess environments occupied by our *Phlox* species. We combined coordinates from our field collections and occurrence data from the Global Biodiversity Information Facility (GBIF; https://www.gbif.org/) and the Southeast Regional Network of Expertise and Collections (SERNEC; https://sernecportal.org/portal/), including records within the native ranges that were identified to subspecies (*Phlox amoena* subsp. *amoena*, *Phlox pilosa* subsp. *deamii*, and *Phlox pilosa* subsp. *pilosa*). We thinned occurrences to one within 20km using the R package 'spThin' (35) and retained 33 amoena, 87 pilosa, and only 5 deamii (Table S2). We could not perform ecological niche modeling analysesfor deamii due to low occurrences.

- 374 We extracted bioclimatic variables from the WorldClim dataset
- 375 (https://www.worldclim.org/data/bioclim.html) and soil composition and chemistry variables
- 376 from the Unified North America Soil Map
- 377 (https://daac.ornl.gov/NACP/guides/NACP\_MsTMIP\_Unified\_NA\_Soil\_Map.html) at each
- 378 occurrence location for amoena and pilosa. We reduced collinearity between variables to retain
- 379 11 variables with correlation coefficients <0.8 (Table S1). With these variables, we constructed
- 380 Maxent ecological niche models for amoena and pilosa using the R package 'dismo' (36)

following established protocols (37, 38). Model performance was evaluated using a repeated

382 cross-fold approach in which 90% of the data were sampled to train a Maxent ENM before

testing the model with the remaining 10% of the occurrence points. For both amoena (median

testing AUC = 0.942) and pilosa (median testing AUC = 0.889), we were able to construct robust

385 niche models (Figure S1).

We performed a principal components analysis on all environmental and soil variables used in our niche models. We assessed if the niches of the two species differed by comparing the empirical differences between species in median and breadth (difference between 5<sup>th</sup> and 95<sup>th</sup> percentile) along PC1 and PC2 to a null distribution defined by bootstrap resampling 1000 times the pooled and randomly reassigned occurrence points across both species (37, 39, 40) (Fig S1).

391 This PCA was later used to calculate environmental distances between populations.

392

393 *Plant propagation* 

394	We propagated collections of 122 genotypes of Phlox amoena amoena (eight				
395	populations), 125 genotypes of Phlox pilosa pilosa (nine populations), and 37 genotypes of				
396	Phlox pilosa deamii (three populations) from throughout their native ranges for our common				
397	garden experiment (Table S5). Wild plants were collected as cuttings of vegetative shoots and				
398	rooted and grown in the greenhouse facilities at the Arnold Arboretum of Harvard University.				
399	After growing for nine months replicate cuttings, each four inches in length, were taken from				
400	vegetative shoots on each plant and rooted and grown in fine potting media for one month before				
401	being transplanted into experimental gardens. To increase the sample size for deamii, we				
402	included two individuals per genotype in each garden, while only one individual per genotype				
403	was planted in each garden for the other two taxa.				
404					
405	Experimental gardens and fitness measurements				
406	We established three experimental gardens adjacent to one native population of each				
407	taxon (Table S6). Each garden site contained four cleared plots into which the 321 plants were				
408	assigned a randomized position. Clonal cuttings from the greenhouse were planted in their				
409	assigned position, which was marked by an aluminum tag. Each plot was protected from large				
410	herbivores by PVC and chicken wire cages for one year after planting. The gardens were watered				
411	immediately after planting and then weekly for a month at which point supplemental watering				
412	stopped.				
413	We monitored fitness-related traits in the gardens over the course of three growing				
414	seasons between planting in April 2018 and final data collection in September 2020.				
415	Survivorship across all three gardens in the first year was high (92.5% in amoena garden, 91.9%				
416	in deamii garden, 92.8% in pilosa garden, 92.4% total). At the end of the first winter in early				

417 2019, we removed the wire cages and returned regularly to record traits throughout spring and summer. We recorded damage from large vertebrate herbivores as a binary trait (0 = herbivore 418 419 damage, 1 = no herbivore damage). We counted the total number of open flowers on each plant 420 on a weekly basis from beginning of April through beginning of June 2019. Flowers on these 421 taxa remain open and fresh for about one week, so our timing minimized double counting or 422 missing flowers. We counted the total number of fruits set by each plant including both mature 423 fruits that remained on the plant as well as open calyces where fruits had already shattered. In 424 October 2019, we harvested all aboveground biomass for each plant, leaving root systems and 425 the stem at the base of each plant intact consistent with the annual aboveground die-back that 426 these taxa experience each winter. We dried this tissue in a drying oven at 60° C for 48 hours 427 before measuring the mass with an electronic scale. We were not able to return to the gardens 428 again until September 2020 when we recorded final survival.

429

#### 430 *Between species adaptive divergence analyses*

431 To test the hypothesis of adaptive differentiation between taxa, we used a generalized linear mixed model (GLMM) approach implemented in the R package 'lme4' (41, 42). For each 432 433 fitness-related trait measured in the gardens, we modeled trait value with fixed effects of taxon, 434 garden, and taxon-by-garden interaction and a random effect of genotype nested within 435 population. Each genotype occurred at least once as a clonal replicate in each garden. For 436 herbivory and survival we used a binomial link function in our models, while for number of flowers and fruits we used a Poisson link function. For biomass, we transformed the raw data by 437 438 taking the natural logarithm and modelled this trait using a linear mixed model. After fitting each 439 model, we evaluated them using ANOVA as implemented in the R package 'car' (43).

Local adaptation between the species is expected to result in a significant taxon-bygarden interaction effect. Specifically, we predict the local taxon to outperform the two foreign taxa in its home garden (local vs. foreign comparisons) and/or for each taxon to perform better in its home garden than in other two habitats (home vs. away comparison) (7). To test these predictions we performed post-hoc contrasts using Tukey's Test as implemented in the R package 'multcomp' (44).

- 446
- 447 Within species local adaptation analyses

448 We implemented a provenance trial analysis to test for local adaptation and thus expected a negative correlation between a plant's performance and the distance between its source and the 449 450 experimental garden in which it was measured. We calculated distance between experimental 451 garden and source population in three ways: geographic, genetic, and environmental (Table S3). 452 We calculated geographic distance with longitude/latitude of each population's wild collection 453 site and each experimental garden using the Haversine formula as implemented in the R package 454 'geosphere' (38). We calculated the genetic distance as F<sub>ST</sub> between each wild source population and an intraspecific population adjacent to each experimental garden site. DNA sequencing and 455 456 F<sub>ST</sub> calculations among these populations are detailed and reported in Goulet-Scott et al. 2021. 457 Briefly, five individuals from each wild population were sequenced using double digest 458 restriction-site associated DNA sequencing (ddRADseq), and all pairwise Weir-Cockerham F<sub>ST</sub> 459 values between populations were calculated using VCFtools (45, 46). Finally, we calculated 460 environmental distance as the Euclidean distance between each population's wild collection site 461 and each experimental garden site in PC1 vs. PC2 space of the environmental PCA that 462 accompanied ecological niche modeling detailed above.

463 To quantify the contribution of the source population to the fitness of each clone in the experimental gardens, we used a GLMM. For each species, we modeled fitness trait value with a 464 465 random effect of population nested within garden, using the same link functions for each trait as 466 described previously. These models yielded "population random effects" for each garden that 467 estimated the average effect on the fitness trait value in that garden attributable to being from a 468 given population. To test for local adaptation, we regressed population random effects for each trait/taxon combination against each measure of distance using linear models as implemented in 469 470 base R (47). For each linear model, we recorded the coefficient associated with the distance 471 predictor, the coefficient of determination  $(R^2)$ , and associated *p*-value.

472

#### 473 *Between and within species trait selection analyses*

Finally, we evaluated how morphological and physiological trait variation predicted 474 fitness both between and within species. We measured a standard suite of morphological and 475 476 physiological traits on a clonal replicate of each experimental individual growing in the Arnold 477 Arboretum greenhouse. These trait measurements required destructive sampling and were 478 therefore not able to be measured on the plants growing in the field without compromising the 479 experiment. From each plant, the most recently fully expanded leaf was collected and the following measurements taken: fresh mass, relative chlorophyll content using an atLeaf 480 481 chlorophyll meter (FT Green, Wilmington, DE, USA), and dry mass. Each fresh leaf was 482 scanned and we used ImageJ to measure leaf length, width, and area. We calculated specific leaf area (SLA) as area (cm<sup>2</sup>) divided by dry mass (g). We summarized variation in leaf traits by 483 484 performing principal component analysis (PCA) on leaf length, width, length/width ratio, area, 485 relative chlorophyll content, and SLA using the correlation matrix. Together the first two

486 principal components described over 70% of the phenotypic variation and were thus used in subsequent analyzes (PC1 = 45.0% of variation explained, PC2 = 27.8%). We used a linear 487 model in R to determine the extent to which species identity explains variation on PC1 and PC2. 488 489 Because we were interested in understanding fitness variation both within and between 490 species we focused our analyses on plants in the pilosa garden and the three fitness traits that 491 showed both adaptive divergence between species and local adaptation within pilosa (flower number, fruit number, and biomass). For these analyzes, fitness traits and PC axes were 492 493 normalized with a mean of 0 and standard deviation of 1. First, we implemented two linear 494 models in R, one for each of the first two PCs, to ask how PC of trait variation, taxon identity, 495 and the interaction between these two main effects predicted fitness trait variation across all three 496 species. Second, we implemented a series of simple linear models in R to specifically ask how 497 PC1 and PC2 predicted fitness variation in four data sets: all species combined, only pilosa, only 498 amoena, and only deamii. By comparing the results of these models for each fitness-trait we 499 could assess whether the same dimension(s) of leaf trait variation predicted fitness within a 500 species versus across all species together. 501

## 502 Funding

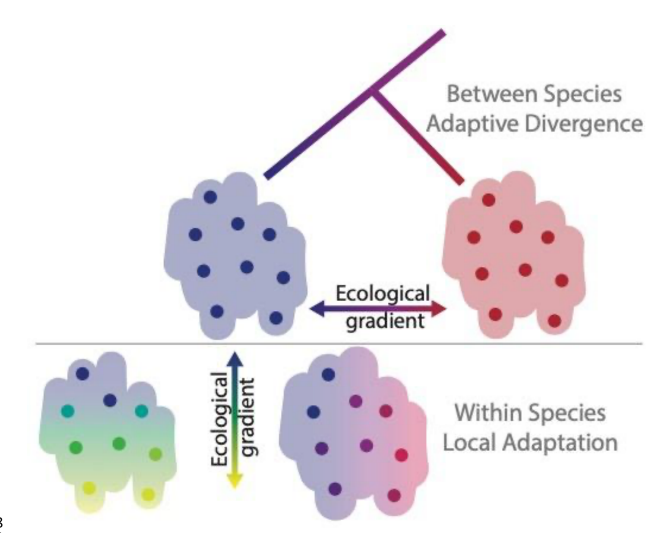
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Figure 1: Conceptual schematic representing divergence across scales of biological diversity in
 response to selection along axes of ecological variation. Top panel represents adaptive

612 divergence between populations of two species shown as blue and red dots on different

613 ecological habitats denoted by red and blue backgrounds. Bottom panel represents alternative

scenarios of within species local adaptation. Each colored point is a population adapted to the

615 gradient of ecological conditions in the habitat represented by color across the background. In

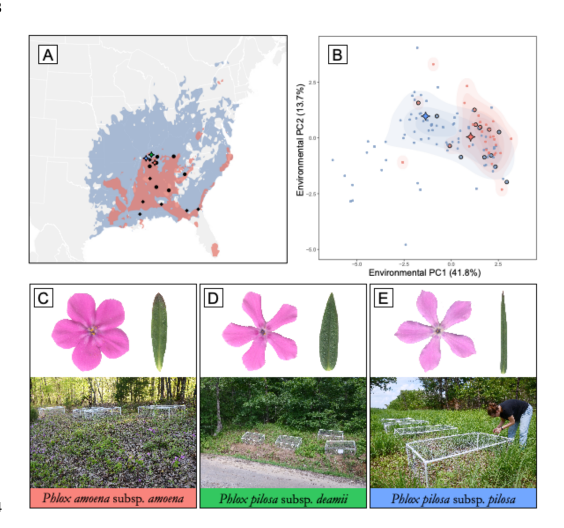
616 the scenario shown at the right (blue to red), the ecological gradient driving within species local

617 adaptation is parallel to the ecological gradient driving between species adaptive divergence. In

- the left scenario (blue to yellow) the gradient of within species adaptation is orthogonal to the
- 619 gradient driving divergence between species (blue to yellow).
- 620



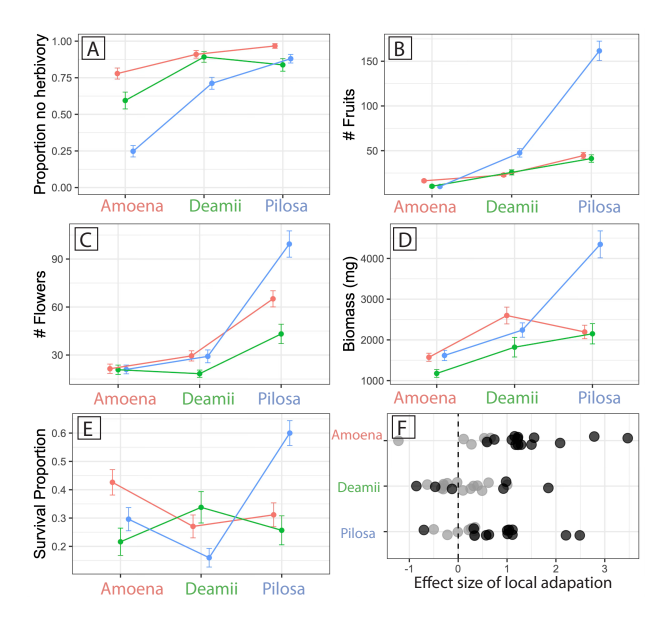
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626 Figure 2: Geographic and environmental variation of broadly sympatric *Phlox* species. (A) 627 Ecological niche modeling predicts the geographic distributions of *P. pilosa pilosa* (pilosa; blue) 628 and *P. amoena amoena* (amoena; red) with sampling locations indicated as black diamonds (pilosa) and black circles (amoena). Locations of the common gardens are indicated by colored 629 diamonds. (B) Environmental variation of pilosa and amoena summarized with a principal 630 component analysis. Blue and red points indicate conditions of known populations of pilosa and 631 632 amoena respectively. Black outlined points are populations sampled for transplant experiment 633 and diamonds are the common garden sites. Representative flowers and leaves (not to scale) and 634 pictures of local common garden site, of amoena (C), P. pilosa deamii (deamii) (D), and pilosa 635 (E). 636

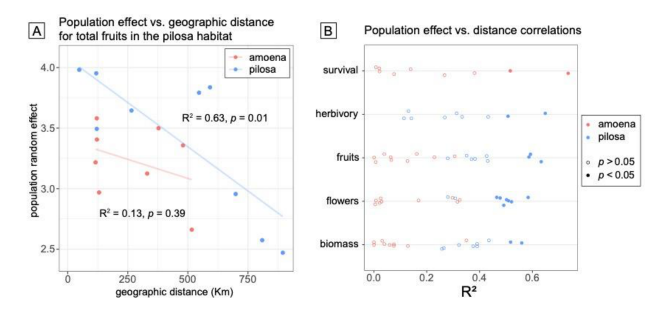


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Figure 3: Performance of each taxon across three garden environments in A) proportion of 639 640 plants without herbivore damage, B) total number of flowers, C) total number of fruits, D) 641 aboveground biomass, and E) proportion survived to the end of the experiment. Values plotted 642 are taxon means +/- standard error in each garden. The ANOVA evaluation of a mixed model analysis for each trait revealed a significant taxon on by garden interaction for all traits. F) 643 644 Summary of effect size of post-hoc contrasts evaluating local adaptation and home-garden 645 advantage for each species. Positive values indicate local species performed superior while 646 negative values indicate local species performs worse. Black points indicate effect sizes 647 significant at p > 0.05. See Table 1 for full model results.

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**Figure 4:** Distance from common garden predicts *P. pilosa pilosa* (pilosa) success indicating

local adaptation. As an example, (A) the relationship between population effect on total fruit set
success and geographic distance for *P. amoena amoena* (amoena; red) and pilosa (blue;

populations with  $R^2$  values indicated. (B) Distribution of  $R^2$  values for population effect vs.

distance measures among populations of amoena (red) and pilosa (blue) grown in all three

experimental gardens. Solid points indicate significant evidence of local adaptation with full

657 model details in Table 2.

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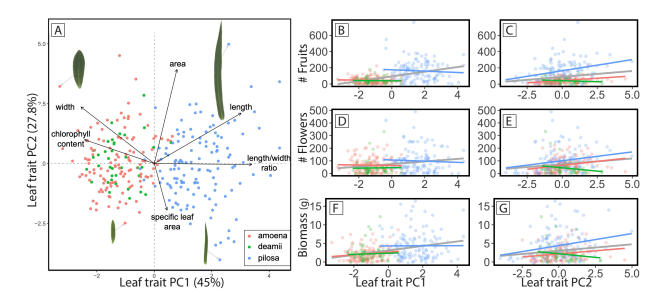




Figure 5: Principal components analysis describing phenotypic variation across three *Phlox* 660 species (A). Points indicate values from individuals grown in the common garden experiment 661 and black arrows indicate loadings of specific traits on the axes of variation. Images of four 662 leaves (to scale) are connected to their points with gray arrows to demonstrate variation in shape 663 along PC1 and area along PC2. (B-G) Relationship between fitness traits and leaf trait variation 664 along PC1 and PC2. Colored points indicate fitness values of individuals grown in the pilosa 665 666 habitat garden. Gray lines show significant relationship across all species and colored lines show relationship within each species (pilosa in blue, amoena in red and deamii in green). For PC1, no 667 668 relationships within species are significant while all within pilosa and amoena relationships are significant between PC2 and fitness traits. Full results in Table S8. 669

**Table 1.** Model details for five fitness-related traits measured in a three-garden reciprocal transplant experiment. The  $X^2$  and *p*-values reported for taxon\*garden interactions were determined by ANOVA on generalized linear mixed models as described in the methods. The contrasts reported for local vs. foreign and home vs. away comparisons were determined using Tukey's Test. Gray shading indicates a contrast showing evidence of adaptive divergence with local species doing better than foreign or a species doing better in the home versus away garden. Dea indicates the deamii home garden site, Pil indicates the pilosa home garden site, and Amo indicates the amoena home garden site.

	4	amoena		deamii		pilosa	
	taxon*garden	local vs. foreign	home vs. away	local vs. foreign	home vs. away	local vs. foreign	home vs. away
herbivory	$X^2 = 17.87$	vs. deamii, p = 0.037	vs. Dea, p = 0.005	vs. amoena, p = 0.625	vs. Amo, p < 0.001	vs. amoena, p = 0.06	vs. Amo, p < 0.001
	p = 0.001	vs. pilosa, p < 0.001	vs. Pil, p < 0.001	vs. pilosa, p = 0.082	vs. Pil, p = 0.328	vs. deamii, p = 0.25	vs. Dea, p < 0.001
flower #	$X^2 = 685.89$	vs. deamii, p = 0.703	vs. Dea p < 0.001	vs. amoena, p = 0.698	vs. Amo, p < 0.001	vs. amoena, p = 0.789	vs. Amo, p < 0.001
	p < 0.001	vs. pilosa, p = 0.422	vs. Pil, p < 0.001	vs. pilosa, p = 0.864	vs. Pil, p < 0.001	vs. deamii, p = 0.634	vs. Dea, p < 0.001
fruit #	$X^2 = 2583.5$	vs. deamii, p = 0.307	vs. Dea, p < 0.001	vs. amoena, p = 0.439	vs. Amo, p < 0.001	vs. amoena, p < 0.001	vs. Amo, p < 0.001
	p < 0.001	vs. pilosa, p = 0.007	vs. Pil, p < 0.001	vs. pilosa, p = 0.288	vs. Pil, p < 0.001	vs. deamii, p < 0.001	vs. Dea, p < 0.001
biomass	$X^2 = 22.03$	vs. deamii, p = 0.384	vs. Dea, p = 0.068	vs. amoena, p = 0.110	vs. Amo, p = 0.91	vs. amoena, p = 0.07	vs. Amo, p < 0.001
DIOIIIASS	p <0.001	vs. pilosa, p = 0.466	vs. Pil, p = 0.955	vs. pilosa, p = 0.4425	vs. Pil, p = 0.103	vs. deamii, p = 0.135	vs. Dea, p = 0.001
survival	$X^2 = 43.05$	vs. deamii, p = 0.004	vs. Dea, p = 0.011	vs. amoena, p = 0.355	vs. Amo, p = 0.099	vs. amoena, p < 0.001	vs. Amo, p < 0.001
	p < 0.001	vs. pilosa, p = 0.041	vs. Pil, p = 0.063	vs. pilosa, p = 0.007	vs. Pil, p = 0.279	vs. deamii, p < 0.001	vs. Dea, p < 0.001