## Comparative Phylogeography of Mexican Rock Fig (*Ficus petiolaris*) and its Fig Wasp Pollinator: Effects of Abiotic Versus Biotic Factors

Kevin Quinteros<sup>1</sup>, Finn Piatscheck<sup>2</sup>, Jordan Satler<sup>3</sup>, Tracy Heath<sup>4</sup>, and John Nason<sup>4</sup>

<sup>1</sup>University of Maryland at College Park <sup>2</sup>Smithsonian Tropical Research Institute <sup>3</sup>The Ohio State University <sup>4</sup>Iowa State University

March 30, 2024

#### Abstract

Numerous studies have tested for geographically congruent spatial genetic structures and population units in codistributed species. Yet, few have elucidated the relative importance of biogeographic influences versus ecological interactions in determining the congruence of genetic structure in coevolving species. Here, we present the first study testing for genetic codifferentiation in a widely distributed and highly-coevolved mutualism, in which symbiont gene dispersal is expected to be positively correlated. In the fig, *Ficus petiolaris*, and its host-specific *Pegoscapus* pollinating wasp, we evaluated the extent to which geographical patterns of differentiation in each species are similar and explained by shared sources of vicariance, co-dispersal, or species-dependent factors. In both species, the Trans-Mexican Volcanic Belt was a major source of vicariance differentiating southern and northern phylogroups. Within these phylogroups, however, fig and pollinator showed surprisingly different population genetic structure. In *F. petiolaris*, the Gulf of California was a strong phylogeographic break in the northern phylogroup. In contrast, within its northern phylogroup, *Pegoscapus sp.* showed no genetic structure and only weak isolation by distance over a 1500 km range. In the southern phylogroup, exceptional genetic differentiation was observed among populations separated by as little as 300 km. Despite mutual selective pressure between figs and fig wasps, and the role of fig wasps in fig gene flow, we conclude that range-wide patterns of genetic differentiation are primarily influenced by biological features unique to each species rather than by shared sources of vicariance or correlated gene dispersal.

## Comparative Phylogeography of the Mexican Rock Fig (*Ficus petiolaris*) and its Fig Wasp Pollinator: Effects of Abiotic Versus Biotic Factors

Kevin Quinteros <sup>\*1,4</sup>, Finn Piatscheck<sup>2,4</sup>, Jordan D. Satler<sup>3,4</sup>, Tracy A. Heath<sup>4</sup>, and John D. Nason<sup>4</sup>

 <sup>1</sup>Department of Biology, University of Maryland, College Park, MD
 <sup>2</sup>Smithsonian Tropical Research Institute, Panama, Panamá
 <sup>3</sup>Department of Evolution, Ecology and Organismal Biology, The Ohio State University, Columbus, OH
 <sup>4</sup>Department of Ecology, Evolution and Organismal Biology, Iowa State University, Ames, IA

<sup>\*</sup>corresponding author: Kevin Quinteros, phylofignatico@gmail.com

#### Abstract

Numerous studies have tested geographically congruent spatial genetic structures 2 and population units in codistributed species. Yet, few have elucidated the relative 3 importance of biogeographic influences versus ecological interactions in determining 4 the congruence of genetic structure in coevolving species. Here, we present the first 5 study testing for genetic codifferentiation in a widely distributed and highly-coevolved 6 mutualism, in which symbiont gene dispersal is expected to be positively correlated. In 7 the fig, *Ficus petiolaris*, and its host-specific *Peqoscapus* pollinating wasp, we evaluated 8 the extent to which the geographical patterns of differentiation in each species are sim-9 ilar and explained by shared sources of vicariance, codispersal, or species-dependent 10 factors. In both species, the Trans-Mexican Volcanic Belt was a major source of vi-11 cariance differentiating southern and northern phylogroups. Within these phylogroups, 12 however, fig and pollinator showed surprisingly different population genetic structure. 13 In F. petiolaris, the Gulf of California was a strong phylogeographic break in the north-14 ern phylogroup. In contrast, within its northern phylogroup, *Peqoscapus sp.* showed no 15 genetic structure and only weak isolation by distance over a range of 1500 km. In the 16 southern phylogroup, exceptional genetic differentiation was observed between popu-17 lations separated by as little as 300 km. Despite mutual selective pressure between figs 18 and fig wasps, and the role of fig wasps in fig gene flow, we conclude that range-wide 19 patterns of genetic differentiation are primarily influenced by biological features unique 20 to each species rather than by shared sources of vicariance or correlated gene dispersal. 21

<sup>22</sup> Keywords: pollination mutualism, coevolution, phylogeography, gene flow, fig and fig wasp

## <sup>23</sup> 1 Introduction

1

Species responses to landscape history can be complex, leading to varied phylogeographic 24 patterns within the same biota [Bowen and Avise, 1990; Lamb et al., 1992; Zink, 1996; 25 Carstens et al., 2005; Soltis et al., 2006; Moussalli et al., 2009. Nonetheless, geographical 26 features of a landscape can generate congruent phylogeographic patterns across taxa by 27 posing common physical barriers to gene flow [Hewitt, 2000; Nason et al., 2002; Pyron and 28 Burbrink, 2010; Garrick et al., 2013]. Similarities and differences in the biology of species, 29 such as dispersal ability, reproductive biology, and niche requirements, can also influence 30 their degree of phylogeographic congruence, as can the intimacy, specificity, and longevity of 31 their ecological interactions [Carstens and Richards, 2007; Moussalli et al., 2009; Smith et al., 32 2011]. Co-evolutionary patterns are often considered on macro-evolutionary timescales, but 33 they can be affected by ongoing landscape-level abiotic and biotic processes [Ehrlich and 34 Raven, 1964: Thompson, 1994, 1999, including the interplay between geographic variation 35 in reciprocal selection, the demographics of local populations, and the homogenizing effects 36 of gene flow – a predication supported by empirical studies Thompson and Cunningham. 37 2002] and theoretical models [Nuismer et al., 1999; Gomulkiewicz et al., 2000; Hochberg et al., 38 2000; Nuismer et al., 2000; Gomulkiewicz et al., 2003]. Although advances in genomic data 39 acquisition and analytical tools continue to enhance our understanding of genetic structure 40 [Knowles, 2009; Papadopoulou and Knowles, 2016; Satler and Carstens, 2017, 2019], debate 41 remains concerning the relative importance of abiotic [Hewitt, 2000; Nason et al., 2002; 42

Pyron and Burbrink, 2010; Garrick et al., 2013] versus biotic [Carstens and Richards, 2007; 43 Moussalli et al., 2009; Smith et al., 2011] in the evolution of this structure. Here we address 44 how biogeographical, ecological, and coevolutionary processes influence population genetic 45 structure within - and the spatial congruence of this structure between - symbiont species in 46 an obligate pollination mutualism. Examples of these highly specialized interactions include 47 yuccas and yucca moths, leafflowers and leafflower moths, globeflowers and globeflower flies. 48 and palms and weevils [Cruaud et al., 2012; Hembry and Althoff, 2016; de Medeiros and 49 Farrell, 2020; Pellmyr et al., 2020], as well as figs and fig wasps, which are the focus of this 50 study. 51

#### <sup>52</sup> 1.1 Co-diversification of Figs and Fig Wasps

The prolonged ( $\sim 80$  Ma) ecological interaction between figs (genus Ficus) and fig 53 wasps (family Agaonidae) has resulted in deep-time codiversification between these lineages 54 [Janzen, 1979; Ramírez, 1970; Weiblen, 2002; Anstett et al., 1997; Cook and Rasplus, 2003; 55 Rønsted et al., 2005; Machado et al., 2005; Cruaud et al., 2012]. Ficus comprises over 800 56 species employing elaborate floral fragrances to attract female fig wasps, which pollinate fig 57 flowers and oviposit in fig ovules, leading to gall formation by developing larvae [Chen and 58 Song, 2008; Wang et al., 2013]. Consequently, the mutualism has coevolved due to reciprocal 59 selective pressures for reproductive success Janzen [1979]; Weiblen [2002]. Although excep-60 tions are known [Molbo et al., 2003; Herre et al., 2008; Su et al., 2008; Wang et al., 2016; 61 Satler et al., 2019]. 62

Despite a brief adult lifespan (<60h) [Kjellberg et al., 1988; Dunn et al., 2008], pollen-63 bearing female fig wasps disperse over long distances, using wind currents to reach receptive 64 fig trees frequently located several kilometers away [Ramírez B., 1969; Nason et al., 1998; 65 Harrison and Rasplus, 2006; Ahmed et al., 2009]. While vertebrate frugivores can disperse 66 fig seeds over long distances [Galil and Neeman, 1977; Staddon et al., 2010], aiding the main-67 tenance of extensive species ranges [Hernández-Esquivel et al., 2020], rates of gene flow via 68 pollen migration are generally much higher than via seed migration in flowering plants [En-69 nos, 1994; Petit et al., 2005], including figs [Yu et al., 2010; Liu et al., 2015]. Recent studies 70 have shown that figs and fig wasps exhibit complex phylogeographical patterns influenced 71 by past climatic and geographical events shared by host and pollinator [Chen et al., 2012; 72 Honorio Coronado et al., 2014; Vieira et al., 2015; Yu and Nason, 2013; Cooper et al., 2020]. 73 Nonetheless, the co-dispersal of fig wasps and fig pollen should promote symmetries in gene 74 flow, leading to the *a priori* expectation that associated fig and wasp species should exhibit 75 spatial congruence in population structure. 76

# <sup>77</sup> 1.2 Biogeography of Baja California Peninsula and Western Mex <sup>78</sup> ico

The rock strangling fig (*F. petiolaris*) and its species-specific fig-wasp pollinator (*Pegoscapus* sp.) are endemic to the Baja California peninsula (BCP) and the western Mexican Transition Zone (MTZ) [Piedra-Malagón et al., 2011, 2019]. BCP was originally thought to have a dispersal-dominated biotic history [Savage, 1960], but our greater understanding of plate

tectonics has facilitated more complex vicariant hypotheses [Hess, 1962; Murphy, 1983; Gris-83 mer, 1994; Graham et al., 2014]. Shared signals of vicariance associated with two ancient 84 transpeninsular seaways and the formation of the Gulf of California [Sedlock, 2003] (Figure 85 1 Hypotheses A–C) have been detected in the spatial genetic structures of several animal 86 and plant species [Riddle et al., 2000a,b,c; Nason et al., 2002; Hurtado et al., 2004; Ross and 87 Markow, 2006; Pfeiler et al., 2007; Garrick et al., 2009, 2013]. Encompassing much of western 88 mainland Mexico, the MTZ includes two major highland provinces that interrupt the ranges 89 of F. petiolaris and its pollinator: the Trans-Mexican Volcanic Belt (TVB) [Ferrari et al.. 90 2012] and the Sierra Madre del Sur [Ferrari et al., 2014] (SMS, Figure 1 hyptheses D and E, 91 respectively). Numerous taxonomic and phylogeographic studies have investigated biogeo-92 graphical patterns within the TVB and SMS regions, frequently finding similar patterns of 93 dispersal and vicariance across taxa. [Halffter, 1964, 1987; Becerra, 2005; Bryson Jr et al., 94 2011; Gutiérrez-Ortega et al., 2018; Rocha-Méndez et al., 2019; Anguiano-Constante et al., 95 2021]. The distributions of F. petiolaris and its pollinator traverse several potential sources 96 of vicariance that, with correlated gene flow, could impact the spatial genetic structure of 97 both species. 98

Here we adopt a multifaceted approach to uncover how abiotic and biotic factors shape gg the genetic structure of interacting fig and fig wasp species. First, we examine the popula-100 tion genetic structure of each species to determine the extent to which they coincide with 101 biogeographical patterns hypothesized for the BCP and MTZ (Figure 1). Second, we com-102 pare the genetic structure of these two co-evolving species to determine the extent to which 103 they are geographically congruent. Finally, we evaluate the relative importance of historical 104 biogeography versus biotic factors (reproductive interactions and co-dispersal) in generating 105 shared phylogeographic patterns between host and pollinator. 106

## <sup>107</sup> 2 Materials & Methods

#### $_{108}$ 2.1 Ficus petiolaris

#### <sup>109</sup> 2.1.1 Host Fig Sampling and Sequencing

Because of the geopolitics of western mainland Mexico, our sampling of F. petiolaris 110 within this area was restricted to regions we could access safely. This resulted in sparse 111 sampling so not all our *a priori* vicariance hypotheses could be tested for the host plant. We 112 sampled 247 F. petiolaris individuals from 19 sites spanning the range of the species in the 113 BCP and mainland Mexico between 2012-2017 (Figure 1, Table 1). To provide an outgroup 114 for phylogenetic analyses, we also sampled three F. aurea individuals from Oaxaca, Mexico. 115 Leaf samples were preserved in silica gel and then stored at -80°C until DNA extraction and 116 library construction at Iowa State University. 117

To identify single nucleotide polymorphisms (SNP) for population genetic analysis, DNA was extracted from leaf tissue using a cetyltrimethylammonium bromide (CTAB) and chloroform protocol, including polyvinylpyrrolidone and proteinase K to increase DNA quality. DNA was then precipitated in 2-propanol. A modified Peterson et al. [2012] dd-RADseq protocol was used to select double digested DNA fragments cut with the restriction en-

zymes PstI and MspI (dd-RADseq). After ligation of Illumina adapters, polymerase chain 123 reaction (PCR) amplified products were size-selected to 300–800 bp using BluePippin. After 124 sequencing at the Iowa State University DNA Facility with an Illumina Hiseq 3000 raw reads 125 were processed and *de novo* assembled with Ipyrad v0.9.31 [Eaton and Overcast, 2020] using 126 these parameters: a maximum of five low-quality base calls with the quality score offset of 127 33, minimum depth for statistical base calling of six, minimum depth for the majority-rule 128 calling of six, and a maximum cluster depth within samples of 10,000. The loci were further 129 filtered to remove those found in less than 50% of the samples, having a maximum of 25%130 heterozygous sites, five or more SNPs, five or more indels or a maximum of five low quality 131 sites. 132

#### 133 2.1.2 Host Fig Population Genetic Structure

A principal component analysis (PCA) was performed to assess the clustering of individ-134 ual multilocus genotypes of F. petiolaris in multivariate space using R v4.0.3 [R Core Team, 135 2020] and the *dudi.pca* function from ade4 v1.7.19 [Dray and Dufour, 2007; Thioulouse et al., 136 2018]. For this analysis, we randomly subsampled two individuals per BCP collection site 137 for our input data set due to the greater density of collection sites there compared to other 138 regions. This subsample was used as input for all subsequent F. petiolaris analyses, unless 139 stated otherwise. Further, to reduce complexity, we limited the dataset to a single biallelic 140 SNP per locus. Allele frequencies were calculated using the R package adegenet v2.1.3 [Jom-141 bart, 2008; Jombart and Ahmed, 2011; R Core Team, 2020] and missing data were imputed 142 by using average allele frequencies within local sample sites. 143

Next, we used STRUCTURE Pritchard et al. [2000] to identify genetic clusters (K) and 144 to estimate membership coefficients of individual samples within each cluster. We applied an 145 admixture model with 500,000 iterations, preceded by a 100,000 iteration burn-in. Analyses 146 covered K = 1 - 8 with 15 replicates per K, and the optimal K was determined using the 147 maximum posterior log-likelihood and  $\Delta K$  via the Evanno method [Evanno et al., 2005] 148 in the pophelper R package v2.3.0 [Francis, 2017]. We iteratively subset the data by the 149 inferred K and re-ran independent STRUCTURE analyses to test potential sub-structuring. 150 Continuing until discrete clusters were no longer evident. To mitigate overestimation of 151 clusters due to isolation by distance (IBD), our final K determination considered all genetic 152 analysis evidence and geographical coherence of clusters. If putative hybrids were detected 153 in the PCA or STRUCTURE analyses, we utilized *snapclust* [Beugin et al., 2018] within the 154 R package adegenet to classify hybrids into either first-generation (F1) or back-cross. 155

To assess the genetic differentiation between inferred genetic clusters and populations 156 adjacent to hypothesized barriers to gene flow (as indicated in Figure 1), we estimated 157 Wright's Fst [Weir and Cockerham, 1984] and 95% confidence limits (999 permutations) 158 using the functions pairwise. WCfst and boot.ppfst in the R package hierfstat version 0.5.11 159 [Goudet, 2005]. Additionally, using hierfstat we computed site-specific estimates of  $F_{LS}$ , which 160 serve as a measure of local inbreeding among relatives. Based on a mating system analysis 161 of F. petiolaris conducted by Gates and Nason [2012], we expected substantial outcrossing 162 across and  $F_{IS}$  estimates close to zero. 163

#### <sup>164</sup> 2.1.3 Host Fig Phylogenetic Reconstruction of Phylogroups

We used a phylogenetic approach to infer the evolutionary relationships among the inferred genetic clusters. SVDquartets [Chifman and Kubatko, 2014], as implemented in PAUP\* (version 4.0a163) [Swofford, 2003], was utilized to construct a population tree based on genetic clusters identified the PCA and STRUCTURE. These analysis used the complete SNP dataset as input, exhaustively evaluating quartets. To assess the robustness of our results, we executed 100 bootstrap replicates to attain node support for the inferred trees.

#### 171 2.1.4 Host Fig Geographical Patterns of Isolation by Distance and Diversity

We conducted a Mantel test (999 iterations) to assess the correlation between genetic and 172 geographic distances to quantify the genetic differentiation resulting from IBD among host 173 populations. Within and between the genetic clusters identified by PCA and STRUCTURE 174 analyses. Population pairwise genetic distances were computed using two metrics: Nei's stan-175 dard distance [Nei et al., 1983] and Cavalli-Sforza and Edwards chord distance [Cavalli-Sforza 176 and Edwards, 1967, which make different assumptions about the roles of mutation, genetic 177 drift, and population size in population divergence through time. These genetic distances 178 were estimated using genet.dist from hierfstat [Goudet and Jombart, 2020] and great circle 179 geographic distances were determined using rdist.earth the fields R package[Douglas Ny-180 chka et al., 2017]. To explore geographical patterns in within-population genetic diversity, 181 we plotted expected heterozygosity ( $H_e$ ) against latitude for each sample population.  $H_e$  ws 182 estimated using the function basicstats in hierfstat. 183

#### <sup>184</sup> 2.1.5 Host Fig Demographic Model Selection

Using the inferred phylogeographic structure, we tested historical demographic models 185 with PHRAPL [Jackson et al., 2017]. Analyzing competing models of differentiation and 186 gene flow across major phylogeographic breaks. The models tested included isolation-only, 187 isolation with migration, and migration-only. For input, we randomly sub-sampled 400 RAD 188 loci without replacement. Per locus, we calculated maximum likelihood gene trees using 189 RAxML v8.2.11, using the GTR substitution model with gamma and proportion of invari-190 ant sites [Stamatakis, 2014]. Upon rooting each gene tree, we removed outgroup taxa and 191 then performed 20 rounds of random subsampling of empirical gene trees with replacement. 192 Subsequently, we conducted simulations of 70,000 gene trees by varying parameter values 193 for divergence time  $(\tau)$  and migration (m). Akaike weights (wAIC) were used for model 194 comparison and to calculate metrics analogous to model probabilities that go from 0 (low 195 support) to 1 (high support). 196

We opted for PHRAPL over an allele frequency spectrum (AFS) approach for demographic model selection. Unlike PHRAPL, AFS approaches cannot handle missing data, which is typical of RAD-seq datasets, necessitating further data downsampling. Moreover, a comparison of allele frequency and gene tree-based approaches in model selection accuracy has not yielded conclusive results [Ruffley et al., 2018]. Both approaches incorporate coalescent theory, and model selection accuracy may depend more on the information available in AFS or gene trees.

#### 204 2.2 The *Pegoscapus* Pollinator

#### 205 2.2.1 Pollinator Sampling and Sequencing

We collected near-mature (late interphase) and mature F. petiolaris syconia (commonly "fruit") from trees across 29 collecting sites between 2012-2019 (Figure 1 and Table 1). The syconia were placed in plastic vials and, pollinators were allowed to emerge before being preserved in 95% ethanol or RNALater. If wasps had not emerged within 24 hours, or a syconium was immature, then in some cases, galled flowers were preserved for later removal of wasp larva or pupae. In total, we genotyped 102 individual *Pegoscapus* sp. wasps, as well as one outgroup pollinator collected from *F. crocata*.

To generate genome-wide sequence data for the wasps, we used targeted enrichment of 213 ultra-conserved elements (UCEs) following the workflow outlined in Faircloth et al. [2012]. A 214 single wasp was selected per syconium to ensure independence among samples, and genomic 215 DNA was extracted with a Qiagen DNeasy Blood and Tissue Kit (Qiagen Inc., Valencia, 216 CA). Samples were fragmented to an average size range of 450 bp using a Covaris ME220 217 focused-ultrasonicator (Covaris Inc., Woburn MA) and Illumina libraries were prepared us-218 ing a KAPA HyperPrep Kit (Roche Sequencing and Life Science). After library construc-219 tion, samples were grouped into eight sets and hybridized with biotinylated RNA probes 220 to capture targeted loci. For targeting UCE loci, we used the hymenopteran probe set  $v^2$ 221 [Branstetter et al., 2017]. After probe hybridization and library amplification, we confirmed 222 size distributions using a Bioanalyzer and pooled the libraries into equimolar concentrations 223 for sequencing. Sequencing was performed by GeneWiz (South Plainfield, NJ) on two full 224 lanes of an Illumina Hiseq 3000 using 150 base pair paired-end sequencing. 225

To process raw sequence reads, we used Phyluce v1.6.7 [Faircloth, 2016] in combination 226 with SAMtools v1.10.2 [Li et al., 2009; Cock et al., 2015]. Raw sequence reads were cleaned 227 with illumiprocessor [Faircloth, 2013; Bolger et al., 2014] and then assembled into contigs us-228 ing SPAdes v3.12.0 [Bankevich et al., 2012]. Contigs were aligned to the hymenopteran probe 229 set to filter out nonspecific sequences. Generated UCE loci were aligned using MAFFT v7.407230 [Katoh and Standley, 2013], edge trimmed with trimAl Capella-Gutierrez et al. [2009], and 231 ambiguously aligned internal sites were removed using Gblocks version 0.91b [Castresana, 232 2000]. To retain a locus, a minimum of 50% sample coverage was required. 233

Omitting the outgroup sample, phased alleles were generated for downstream analysis 234 following the pipeline outline by Andermann et al. [2018]. We took our aligned loci, before 235 the use of Gblocks, and used the *phyluce\_snp\_bwa\_mulitple\_align* function to map cleaned 236 sequenced reads for each sample to the loci using BWA-MEM [Li, 2013] in bwa v0.7.17237 [Li and Durbin, 2010]. *phyluce* snp phase uces was then used to phase the mapped reads, 238 resulting in two alleles per individual per locus [Andermann et al., 2018]. The loci were then 239 realigned and cleaned as previously described, with a minimum requirement of 50% sample 240 coverage. 241

Additionally, owing to the prevalence of mitochondria in animal tissues, UCE sequence capture probes can recover maternally-inherited mitochondrial DNA sequences. The mitochondrial sequences of the cytochrome oxidase subunit I (COI) were assembled using *Novoplasty* using a reference COI (JN103329) sequenceDierckxsens et al. [2016]; Cruaud et al. <sup>246</sup> [2012]. The alignment of COI sequences was done using MAFFT.

#### 247 2.2.2 Pollinator Population Genetic Structure

We used a custom Python script to extract unlinked SNPs from our phased and unphased UCE loci as input data. We primarily used the phased data for our analyses, however phasing was not very effective for every sample. In instances where subsetting the dataset resulted in excessive missing data, we chose to utilize the unphased data. The phased dataset was used for identifying genetic clusters using PCA and STRUCTURE, the latter employing the same approach as for *F. petiolaris* Potential hybrid individuals were classified using *snapclust*, and the results were validated using COI gene trees inferred through RAxML.

In a survey of recently pollinated F. petiolaris syconia at various locations in Baja California and Coastal Sonora, we observed an average of 1.48 pollinator foundresses per fruit (based on 245 fruits, unpublished data). Given the frequent occurrence of a single foundresses and their broods in fruits, inbreeding is anticipated to be common and  $F_{IS}$  estimates should be substantially greater than zero. It is important to note that  $F_{IS}$  could not be estimated for three locations with only one sampled pollinator each (sites 214, 220, 228 in Table 1).

#### 261 2.2.3 Pollinator Phylogenetic Reconstruction of Phylogroups

We used SVDquartets to infer phylogenetic relationships among all *Pegoscapus* sp. phylogroups. For this analysis, we concatenated all recovered UCE loci into a single data matrix and exhaustively evaluated quartets, with 100 bootstrap replicates used to assess nodal support values.

#### 266 2.2.4 Pollinator Geographical Patterns of Isolation by Distance and Diversity

As for the host fig, we identified the extent of spatial genetic differentiation due to IBD by performing a Mantel test of the relationship between population pairwise estimates of genetic distance (Nei's and CavalliSforza and Edwards distances) versus geographical distance, both within and between inferred phylogroups. We also plotted population-level  $H_e$  against latitude to investigate spatial trends in genetic diversity.

#### 272 2.2.5 Pollinator Demographic Model Selection

Using the approach previously described, we assessed the fit of demographic models to 273 inferred genetic clusters using PHRAPL. For *Peqoscapus sp.* we used 767 unphased UCE 274 loci containing our outgroup taxon (pollinator of F. crocata). After estimating gene trees in 275 RAxML and trimming outgroup taxa from each rooted tree, gene trees were subsampled at 276 random with replacement 15 times to reduced computation load. After simulating 50,000 277 gene trees under a range of model  $(\tau)$  and m values, the lnL and AIC of each model were 278 calculated with respect to the empirical gene trees. As with the host fig, wAIC was used to 279 compare models and calculate metrics analogous to model probabilities. 280

#### 281 2.3 Congruence of Genetic Structure between Host and Pollinator

We used population graphs to assess the spatial symmetry of genetic structures in F. *petiolaris* and *Pegoscapus* by measuring the congruence of their population graph topologies [Dyer and Nason, 2004; Dyer, 2015]. Population graphs use graph theory to visually represent population genetic structure, creating a model-free network based on conditional genetic covariance. Populations are depicted as geometric nodes, with their size proportional to the within-population component of genetic variance. Connections between populations are represented by edges, the magnitude of which corresponds to inter-population variance [Dyer, 2015].

To assess topological congruence within the *popgraph* R package [Dyer, 2021], it is re-290 quired to generate population graphs using sites that are co-sampled in both species. We 291 assumed that nearby Oaxacan sites 209 (Pegoscapus) and 210 (F. petiolaris) are equivalent 292 for this analysis, as were the sites 214 (F. petiolaris) and 222 (Pegoscapus). We tested the 293 topological congruence of the population graphs in two ways. First, we asked if nodes that are 294 close in the F. petiolaris graph are also close together in the Pegoscapus graph by measuring 295 the correlation between the shortest path matrices of the two graphs. For this, we used the 296 test congruence function of the popgraph package to calculate the non-parametric correla-297 tion of pair-wise path distances through the graph between sites. Second, we used custom 298 code to examine the congruence of connectivity pattern between sites. Using permutation 290 (n = 10000), we tested whether the F. petiolaris and Pegoscapus population graphs share 300 more edges in common than expected by chance. Finally, we overlayed a population graph 301 of *Peqoscapus sp.* onto a raster map of Mexico to determine whether pollinators preferen-302 tially travel through lower elevations. To assess this, we used a permutation test with 999 303 iterations, randomly reassigning connections (edges) between sites (nodes). We computed 304 the mean elevation of edge set configurations by extracting elevation data along edges from 305 our raster map [Dyer et al., 2012]. Statistical significance was evaluated by comparing the 306 observed mean elevation to the distribution of mean elevation values generated through per-307 mutation. Under the null hypothesis, this approach assumes no differences in mean elevation 308 across edge set configurations. 309

### 310 **3 Results**

#### 311 3.1 Ficus petiolaris

#### 312 3.1.1 Host Fig Sampling and Sequencing

We generated 137,216,883 single-end raw reads. After *de novo* assembly, 25,997 loci were discovered. We filtered out loci found in less than 50% of samples, which removed most of the loci, indicating a large amount of missing data across individuals. To prevent the use of linked SNPs, only one SNP per locus was kept for downstream analyses. We later removed outgroups and individuals with less than 30% of genomic data across loci, which resulted in somem polymorphic loci becoming monomorphic in *F. petiolaris* only. Removing monomorphic loci resulted in a final data set of 1,192 biallelic SNPs across 247 individuals.

#### 320 3.1.2 Host Fig Population Genetic Structure

The initial two principal components of the PCA demonstrated large eigenvalues, collectively explained 52% of the variation in the data. Based on genetic differences, individual samples formed five distinct and well-separated clusters in multivariate space, which we interpret as comprising five distinct genetic clusters. PC axes one and two separated Baja California, Coastal Sonora, Inland Sonora + Central Mexico (Jalisco and Sinaloa), and Southern Oaxaca (Figure 2B), while PC one and PC three further separated Inland Sonora and Central Mexico (Figure 2C). The Coastal Sonora samples were intermediately spread between the Baja California versus Inland Sonora and Central Mexico genetic clusters in PCA space, suggesting admixture between well-differentiated Baja and northwestern Mexico populations.

The optimal number of genetic clusters for STRUCTURE analysis was K = 2 based on maximum posterior log–likelihood and  $\Delta K$  (Figure 2A). STRUCTURE supported the differentiation of Baja California from mainland Mexico and identified samples from Coastal Sonora as admixed between these two regions. However, it did not distinguish Inland Sonora or Central Mexico as separate from Oaxaca. We subset samples based on the two inferred STRUCTURE clusters and ran STRUCTURE on each subset. No geographically associated samples were further identified.

Multilocus estimates of  $F_{st}$  (Table S1) revealed significant differentiation between all 338 five regional genetic clusters of F. petiolaris represented in the PCA (Figure 2B & C). 339 Southern Oaxaca showed high genetic differentiation from all other clusters ( $F_{st} = 0.32 -$ 340 0.69). Similarly, Baja California exhibited significant differentiation from the other clusters 341  $(F_{st} = 0.28 - 0.69)$ . Despite their close geographic proximity, Coastal Sonora was strongly 342 differentiated from both Baja California ( $F_{st} = 0.28$ ) and Inland Sonora ( $F_{st} = 0.16$ ). In 343 contrast, Inland Sonora and Central Mexico showed weaker differentiation from each other, 344 with an  $F_{st}$  value of 0.07, despite their larger geographic span. 345

Given our density of sampling in northwestern Mexico, we were able to test vicariance 346 hypotheses A-C in Figure 1. Genetic differentiation between sites adjacent to the MPS (Ho 347 Figure 1A) and ILP (Ho Figure 1B) seaways was not significant ( $F_{st} = -0.019$  and 0.006, 348 respectively). In contrast, differentiation between the Baja California and Coastal Sonora 349 genetic clusters adjacent to the Gulf of California (Ho Fig.1C) was large and significant 350  $(F_{st} = 0.28)$ , as noted above). Consistent with expectations based on prior research in Baja 351 California [Gates and Nason, 2012], inbreeding in F. petiolaris populations was found to be 352 low. Averaged across sites,  $F_{IS}$  was 0.043 (range -0.069 to 0.118) with nine of 19 locations 353 having estimates significantly greater than zero. These results indicate that F. petiolaris is 354 highly outcrossing across its range, regardless of the genetic cluster or geographic region. 355

#### 356 3.1.3 Host Fig Phylogenetic Reconstruction of Phylogroups

The population tree showed that the genetic clusters of northern and central Mexico Baja 357 California, Coastal Sonora, Inland Sonora, Central Mexico formed a single monophyletic 358 clade closely sister to Southern Oaxaca (Figure 3A). The SVDquartets analysis generally 359 supported the results of the population genetic analyses. The SVD quartets population tree 360 (Figure 3A) placed Southern Oaxaca sister to all other genetic clusters, while Baja California 361 and Coastal Sonora (in north-central Mexico) formed a well-supported clade (BS = 100) sister 362 to a weakly supported Inland Sonora plus Central Mexico clade (BS = 50). However, these 363 findings should be interpreted cautiously as a bifurcating tree is not optimal for handling 364 admixture. 365

#### <sup>366</sup> 3.1.4 Host Fig Geographical Patterns of Isolation by Distance and Diversity

Sampling of F. petiolaris in mainland Mexico was relatively sparse, so we limited testing 367 of IBD to Baja California, spanning *a priori* sources of vicariance (Figure 1A and B). Using 368 all individuals from Baja California, a significant positive (R = 0.352, p = 0.029) correlation 369 was observed between Nei's genetic distance and geographical distance across the peninsula 370 (Figure 4), with similar results obtained using Cavalli-Sforza and Edwards chord distance 371 (results not shown). Though historical vicariance cannot be ruled out, the continuous IBD 372 indicates no lasting signals of vicariance associated associated with the ancient transpenin-373 sular seaways. Figure 5 shows that both Baja California  $(0.03 < H_e < 0.042)$  and Southern 374 Oaxaca  $(0.045 < H_e < 0.0475)$  genetic clusters have relatively low population genetic diver-375 sity. These clusters represent the northern and southern limits of our geographic sampling. 376 In contrast, populations in the Coastal Sonora, Inland Sonora, and Central Mexico genetic 377 clusters had substantially higher genetic diversity  $(H_e = 0.069 - 0.082)$ . 378

#### 379 3.1.5 Host Fig Demographic Model Selection

We tested 46 demographic models based on the inferred north-central and southern Mex-380 ico phylogroups of F. petiolaris indicated in the SVD quartets population tree. These models 381 included: 1) migration from the north-central to southern phylogroup, 2) migration from 382 the southern to north-central phylogroup, and 3) a coalescent event between these two phy-383 logroups. The two highest-ranked demographic models had wAIC values greater than 0.39. 384 while the remaining models all had much lower support (Table S3). These two top models 385 both indicate divergence with gene flow between the north-central and southern Mexico phy-386 logroups. More specifically, these models share the same bidirectional gene flow parameters 387 of m = 4.64 (in units of 4Nm) while differing in the timing of the coalescent event, which 388 was deeper in the top-ranked model, t = 7.691, than in the second, t = 0.300 (in units of 389 4N generations). Surprisingly, the PHRAPL models ranked 3 and 5 have the same migra-390 tion parameters and very similar coalescent times to models 1 and 2, respectively, yet have 391 substantially lower wAIC values (Table S3). 392

#### <sup>393</sup> 3.2 The *Pegoscapus* Pollinator

#### <sup>394</sup> 3.2.1 Pollinator Sampling and Sequencing

We generated 390,571,240 raw reads resulting in an average of 5,139,095 reads per indi-395 vidual  $(\pm 3, 334, 310)$ . After processing and filtering, our unphased data set with outgroup 396 taxa had a pool of 2359 loci with an average of 1940 loci per individual ( $\pm$  392.02). Requiring 397 a minimum of 50% taxon coverage to retain a locus, the final data set consisted of 2053 loci. 398 Loci had an average length of 1135 bp ( $\pm$  519.12), ranging from 262 bp to 4534 bp. For the 399 data set without outgroup taxa, we recovered a pool of 2332 loci with an average of 1952 loci 400 per individual ( $\pm 354.62$ ). Requiring a minimum of 50% taxon coverage, the unphased data 401 set consisted of 2057 loci with an average length of 1134 bp ( $\pm 522.53$ ), ranging from 302 bp 402 to 4534 bp. The phased data set (50% taxon coverage) consisted of 1886 loci with an average 403 length of 552.19 bp. One individual did not phase well (FW261) and was removed from the 404 final phased data set. After extracting unlinked, biallelic SNPs using a custom script, the 405

unphased data set had 1921 SNPs, and the phased data set had 1414 SNPs. Additionally,
we were able to recover and assemble COI sequences for 82 individuals.

#### 408 3.2.2 Pollinator Population Genetic Structure

The first two principal components of the PCA accounted for 49.95% and 7.40%, re-409 spectively, of the variation in PCA space (Figure 6B). We identified three primary and 410 well-separated genetic clusters: Northern Mexico, Morelos + Northern Oaxaca, and South-411 ern Oaxaca. The Northern Mexico cluster included individuals from Baja California, Sonora, 412 Sinaloa, Zacatecas, and Jalisco. One individual from Jalisco appeared to be a putative hybrid. 413 which was excluded from subsequent PCA. We analyzed a subset of the data to explore ge-414 netic sub-clustering within the main inferred clusters. There was no additional sub-clustering 415 revealed for individuals from Northern Mexico through PCA. For southern Mexico, PC axe 416 1 versus 2 revealed three clusters representing individuals from Morelos, Northern Oaxaca. 417 and Southern Oaxaca (Figure 6D). 418

STRUCTURE supported a strong division between northern and southern Mexico with an optimal K = 2 genetic clusters based on both the maximum posterior log-likelihood and  $\Delta K$  (Figure 6A). Evidence of admixture between these two clusters was minimal, except for the inferred hybrid individual from Jalisco. As with PCA, additional STRUCTURE runs on these two genetic clusters showed no additional sub-structure in the north, but revealed K = 3 clusters for southern Mexico, corresponding to Morelos, Northern Oaxaca, and Southern Oaxaca (Figure 6C).

Both PCA and STRUCTURE identified a pollinator individual (P342) from Jalisco as a 426 putative hybrid. Snapclust modeling inferred the putative hybrid to be an F1 hybrid between 427 north-central and southern Mexico genetic clusters The COI gene tree placed the F1 hybrid 428 within the Morelos + Northern Oaxaca clade, indicating maternal transmission from that 429 region and paternal contribution from Jalisco. However, since the maternal ancestor originally 430 dispersed from the south to Jalisco, their descendants had an equal chance of inheriting 431 the maternal haplotype or a local one. Consequently, the geographic origin of the hybrid's 432 haplotype is not highly informative. 433

To avoid inflating Wright's  $F_{st}$  estimates, the pairwise calculations (Table S4) excluded 434 the F1 hybrid from Jalisco. Within the north-central Mexico genetic cluster, genetic dif-435 ferentiation between nested regional populations was generally low, with most pairwise  $F_{st}$ 436 estimates <0.07, and only five significantly greater than zero. Low differentiation was ob-437 served between regional populations around the Gulf of California ( $F_{st} \leq 0.021$ ). Higher 438 differentiation was found between Zacatecas and Coastal Sinaloa ( $F_{st} = 0.134$ , sig.), and 439 Inland Sinaloa ( $F_{st} = 0.060$ , n.s.), as well as between Jalisco and Coastal Sinaloa ( $F_{st} =$ 440 0.064, sig.). While southern populations were geographically closer than northern ones, the 441 differentiation among the three inferred genetic clusters exhibited greater variability. For ex-442 ample,  $F_{st}$  between Morelos and Northern Oaxaca was 0.057. However,  $F_{st}$  increased to 0.577 443 and 0.558 when comparing Morelos and Northern Oaxaca with Southern Oaxaca, respec-444 tively. Differentiation was consistently high between regional populations in the north-central 445 phylogroup versus the southern phylogroups  $(F_{st} = 0.715 - 0.809)$ . 446

447 Consistent with expectations based on the observation of multifoundress fruit, deviations

from HWE in *Pegoscapus sp.* populations were found to be substantial. Averaged across the 26 sites with sample sizes greater than one,  $F_{IS}$  was 0.393 (data not shown). Further, 20 sites had estimates significantly greater than zero. The remaining six sites had an even larger mean  $F_{IS} = 0.512$  but also had very broad confidence limits on their site-specific estimates. These results indicate that the pollinator maintains a highly inbred mating system throughout its range, irrespective of genetic cluster or geographical area.

#### 454 3.2.3 Pollinator Phylogenetic Reconstruction of Phylogroups

The SVD quartets population tree supported the division between north-central and southern Mexico (BS = 100, Figure 3). Southern Mexico clade had subclades Morelos + Northern Oaxaca (BS = 93) and southern Oaxaca (BS = 100). Southern Oaxaca branch was sister to Morelos + Northern Oaxaca.

#### <sup>459</sup> 3.2.4 Pollinator Geographical Patterns of Isolation by Distance and Diversity

In north-central Mexico, there was weak but significant IBD (R = 0.281, p = 0.035). 460 Figure 7A), even though populations were separated by distances of up to 1500 km. In the 461 GOC region, no significant IBD was found, indicating genetic connectivity over a 1000 km 462 range (R = 0.158, p = 0.124, Figure 7B). In contrast, within southern Mexico, despite the 463 close proximity of populations ( $\leq 400 \text{ km}$ ), strong and significant IBD was observed (R 464 = 0.847, p = 0.021, Figure 7C) consistent with Morelos, northern and southern Oaxaca 465 comprising three distinct genetic clusters. When comparing between northern and southern 466 Mexico populations, no significant IBD was detected (R = -0.318, p = 0.735, Figure 7D), 467 indicating the TVB acts as a strong barrier to gene flow. An analysis specifically between 468 populations bracketing the TVB (Sites 215 and 221) yield inconclusive results, attributed 469 to a limited sample size. Genetic diversity in the pollinator did not show a clear latitudinal 470 pattern like the host fig with  $H_e$  varing widely among populations within and across regions 471 (Figure 5). The analysis excluded the F1 hybrid individual from Jalisco to avoid inflating 472  $H_e$ . 473

#### 474 3.2.5 Pollinator Demographic Models Selection

Using PHRAPL, we tested the same 46 demographic models as we did for F. petiolaris. 475 We initially ran PHRAPL models on populations immediately north (Jalisco and Zacate-476 cas) and south (Morelos and Northern Oaxaca) of the TVB. Due to limited sample sizes, 477 PHRAPL failed to differentiate between models. Thus, we included all northern and south-478 ern populations for the analysis. Additionally, we excluded the F1 hybrid individual from 479 Jalisco to assess whether gene flow played a significant role in our demographic models, in-480 dependent of any recent migration event's influence. The highest-ranked PHRAPL models 481 (wAIC = 0.85, Table S6) included gene flow and a coalescent event. These results indicate 482 divergence with gene flow between the northern and southern Mexico *Peaoscapus sp* phy-483 logroups. The leading model indicated an asymmetrical gene flow pattern, with estimated 484 migration from south to north being approximately twice as much as from north to south 485  $(m1_2.1 \text{ vs. } m1_1.2, \text{ Table S6}).$ 486

#### 487 3.3 Congruence of Host and Pollinator Population Structure

In the estimated population graph, *F. petiolaris* sites were grouped into four distinct clusters (Fig. 8A). These clusters represent independently evolving units, categorized as follows: (1) Baja California sites, (2) coastal Sonora sites, (3) central Mexico and inland Sonora sites, and (4) Oaxacan sites. The population graph for *Pegoscapus sp.* revealed two primary clusters (Fig. 8 B). The first consisting exclusively of Oaxacan sites and the second cluster encompassing the remaining sites (Baja California, Coastal Sonora, Inland Sonora, and Central Mexico)

Our test for topological congruence revealed a weak and statistically non-significant cor-495 relation (R=0.1457, p=0.5515, t=0.6075, df=17, 95% CI of [-0.3303, 0.5627]) between the 496 shortest path matrices of F. petiolaris and Pegoscapus sp. graphs. The weak "distance con-497 gruence" correlation between the two graphs suggests differences in their respective inter-498 population variance. The test of "structural congruence," which examined connectivity pat-499 terns between sites, did show a significance number of shared edge connections between the 500 F. petiolaris and Peqoscapus population graphs (p=0.0048, rep=10,000, total possible edges 501 = 78, edge correlation = 0.2696). Considering the node and edge count in F. petiolaris (11) 502 edges) and *Peqoscapus sp.* (15 edges) population graphs, the five shared edges between these 503 graphs are more than expected by chance. These shared edges (100-104, 158-39, 172-39, 201-504 95, and 209/210-214/222) show a distribution across geographic locations, with three located 505 in Baja California, one in Coastal Sonora, and one in Oaxaca. All shared edges connected 506 neighboring sites, with the exception of site 39 in Baja California, which is the southernmost 507 BCP site yet shares edges with the two northernmost BCP sites, 172 and 158. 508

Our last test investigated whether pollinators demonstrate a preference for traveling 509 through lower elevations compared to those encountered through out their species dis-510 tribution. Our analysis revealed a statistically significant difference between the observed 511 mean elevation and the distribution of mean elevation values generated through permuta-512 tion  $(p-value = 0.037, mean_{obs} = 1559.69 \text{ m}, n = 999)$ . This suggests that the arrangement 513 of edges we observed in the population graph of *Peqoscapus sp.* is associated with lower ele-514 vations compared to the complete range of potential edge connections between nodes. Impor-515 tantly, these results hold even when considering variations in both population geographical 516 distribution and graph structure. 517

### 518 4 Discussion

The interplay of historical gene flow and vicariance events is a key determinant of con-519 temporary patterns of genetic variation, providing insights into evolutionary processes and 520 biogeographical scenarios. Biotic associations, particularly in obligate pollination mutual-521 ists, also affect species spatial genetic structure. Here, we explored how historical gene flow, 522 vicariance, and contemporary biogeography patterns influence genetic variation by investi-523 gating these dynamics in Western Mexico's complex geography. Despite the strong selective 524 pressure of their mutualistic relationship, species-specific biological traits strongly influence 525 the population genetic structure of figs and fig wasps. 526

## 4.1 Genetic Patterns in Baja California Peninsula and Mexican Highlands

The Baja California Peninsula (BCP) and the Mexican Highlands have played pivotal 529 roles in shaping the genetic patterns and distribution of fauna and flora [Riddle et al.. 530 2000c; Mastretta-Yanes et al., 2015]. In the BCP, we focus on three hypothesized sources of 531 vicariance: the mid-peninsular seaway (MVP), the formation of the Isthmus La Paz (ILP), 532 and the Gulf of California (GOC, Fig. 1 A-C) [Upton and Murphy, 1997; Riddle et al., 533 2000c]. Within the Mexican Highlands, we focused on two prominent features as potential 534 sources of vicariance: the Trans-Mexican volcanic belt (TVB) and the Sierra Madre del Sur 535 (SMS, Fig. 1 E-F). 536

We identified four distinct genetic clusters in F. petiolaris: (1) Baja California, (2) Inland 537 Sonora, (3) Central Mexico, and (4) Southern Oaxaca – with a fifth zone of admixture, 538 Coastal Sonora, located between Baja California and Inland Sonora (Fig. 2B & C). This 539 genetic structure is unlikely to be the result of restricted gene flow due to local inbreeding, as 540 low estimates of  $F_{IS}$  (Table S2) suggest highly outcrossing populations. For Pegoscapus sp., 541 we identified four distinct genetic clusters: (1) northern Mexico (Baja California + Sonora + 542 Sinaloa + Zacatecas + Jalisco), (2) Morelos, (3) Northern Oaxaca, and (4) Southern Oaxaca 543 (Fig. 6). There is no evidence of admixture between northern Mexico and the genetic cluster 544 south of the TVB. Despite a large estimate of  $F_{is}$  (Table S5), Pegoscapus interpopulation 545 genetic connectivity remains high throughout northern and central Mexico (*i.e.*, low  $F_{st}$ 546 values; Table S4). 547

#### <sup>548</sup> 4.2 Lack of Vicariance Within the Baja California Peninsula

Long-term habitat stability influences the spatial distribution of intraspecific genetic 549 diversity [Vasconcellos et al., 2019], leading to expected phylogeographic congruence be-550 tween taxa. However, recent range expansion can obscure the signatures imposed by long-551 term habitat stability. The evolutionary history of BCP is often cryptically integrated into 552 widespread species and species-groups genetic structure[Upton and Murphy, 1997; Riddle 553 et al., 2000c]. Not all BCP taxa exhibit such shared vicariance signals, with some species 554 showing limited or no genetic structure [Vázquez-Miranda et al., 2022]. Similarly, our results 555 indicate that MVP and ILP formation has not left a vicariance signal in the genetic structure 556 of F. petiolaris and Peqoscapus Weak IBD was observed in F. petiolaris BCP populations 557 (Figure 4). High genetic connectivity was observed despite an 800 km distance between 558 sites, which was supported by the low genetic differentiation of pollinators between BCP 559 sites (Table S2 and Table S4). 560

Nason et al. [2002] proposed that frost-sensitive plant systems and their associated insects experienced a contraction of their geographical ranges into southern refugia during the last glacial maximum (LGM), followed by a subsequent northward expansion to their present distributions in the Holocene. The absence of vicariance signals in *F.petiolaris* and *Pegoscapus sp.* could be attributed to the contraction of its range into the southern BCP refugia during the LGM. However, sampling is insufficient to reach a definite conclusion. Although the BCP genetic cluster shows the lowest genetic diversity among the inferred genetic clusters (as shown in Figure 5A and Table S2), it is difficult to determine whether *F.petiolaris* expanded from a central refugee population, as southern Oaxaca also has low genetic diversity. Even if insects possess moderate frost-tolerance, their genetic structure is expected to be affected by the frost-sensitivity of their host. Consequently, glacial-interglacial cycles of the Pleistocene are likely to have impacted the genetic structure of the frost-sensitive desert flora and their associated insects [Nason et al., 2002; Garrick et al., 2009, 2013].

## 4.3 Divergent Phylogeographic Patterns and Contrasting Life His tory Strategies

Despite sharing some phylogeographic patterns, there are points of incongruence between 576 F. petiolaris and Pegoscapus sp. around the BCP. PCA (Fig. 2B & C), along with the pop-577 ulation tree from the SVD quartets (Fig. 3 A) highlight the GOC as a factor contributing to 578 the separation between F. petiolaris populations in Baja California and the mainland (Fig. 579 1C). The effectiveness of GOC as a barrier to genetic flow is limited due to detection of 580 gene admixture in coastal Sonora (Fig. 2A). Baja California had the lowest genetic diversity 581 among clusters, whereas coastal Sonora exhibited higher  $H_e$ , akin to Inland Sonora and Cen-582 tral Mexico (Figure 5A), suggesting that Inland Sonora and Central Mexico predominantly 583 contribute to the genetic variance in Coastal Sonora. 584

This signal of vicariance and admixture is absent in *Peqoscapus* (Fig. 6), and genetic 585 differentiation is relatively low in north central Mexico (Table S1). Weak IBD was detected 586 (Fig. 7A & B), but there was no genetic structuring in *Peqoscapus* (Fig. 6A & B). Differences 587 in life-history strategies between F. petiolaris and Peqoscapus sp. may explain the contrasting 588 vicariance signals. Classical coevolutionary models predict that obligately interacting species 589 adhere to the same adaptive principles and operate within similar time frames [Alvarez et al., 590 2010]. We propose considering their ecological relationships and idiosyncrasies, including two 591 life-history traits that influence distinct vicariance signals: 1) life-span and 2) gene flow. When 592 a pollinator has a shorter generation time and greater dispersal capabilities than its host, 593 signs of vicariance are expected to diminish more rapidly in the pollinator species than in its 594 host. Ficus petiolaris and its pollinator have considerably different generation times (years 595 to decades vs. 44-77 days, respectively) [Piedra-Malagón et al., 2019]. As such, the pollinator 596 *Peqoscapus* possesses traits that promote gene flow among its populations more effectively 597 than its host. 598

Pollen migration contributes to total gene flow by at least an order of magnitude more 599 than seed migration, a pattern similarly observed in the genus *Ficus* [Petit et al., 2005; Yu 600 et al., 2010; Liu et al., 2015]. Fig wasps can travel long distances exceeding 100 km, aided 601 by prevailing wind currents [McKey, 1989; Ahmed et al., 2009]. Consequently, pollinator 602 dispersal plays a substantial role in influencing the gene flow and genetic structure of host 603 fig populations. This does not ensure a congruent phylogeographic signal between the host 604 and its pollinator. Cross-pollinated F. petiolaris seeds often fail to produce viable offspring. 605 while female wasps that access fig syconia tend to have more successful reproduction [Crawley 606 and Ross, 1990; Bronstein and Hossaert-McKey, 1996]. 607

Extreme events, such as droughts, can further increase incongruence in genetic structure

by causing local fig wasp extinction while leaving host-plant populations intact [Harrison, 609 2000. High summer temperatures can reduce syconium volume, leading to smaller wasps 610 with fewer resources for long-distance pollination [Krishnan et al., 2014]. These phenomena 611 are likely to become more prevalent due to climate change. Given these factor two scenarios 612 emerge: (1) reduced genetic structure of the pollinator compared to the host due to long-613 distance pollinator dispersal or (2) pronounced genetic structure in fig wasp populations 614 due to bottlenecks and founder effects from limited colonizers. Our data, along with the 615 results from Liu et al. [2015], support the first scenario, which estimated that pollinators (W. 616 *pumilae*) introduced genetic variability (cpDNA and nuclear DNA) into their populations at 617 a rate of 14:1 compared to genetic variability introduced into the populations of the host plant 618 (F. pumila). This discrepancy suggests that generation time, dispersal, and abiotic factors 619 influences play a crucial role in shaping discordant population genetic structure patterns 620 between F. petiolaris and Pegoscapus. 621

#### <sup>622</sup> 4.4 Barriers to Gene Flow and Hybridization Dynamics

The genetic landscape of *F. petiolaris* populations across the TVB exhibit strong genetic differentiation. Minimal IBD was observed in BCP *F. petiolaris* samples, so IBD is unlikely to explain the significant genetic differentiation across the TVB. Futhermore, the genetic differentiation between inland Sonora and central Mexican *F. petiolaris* populations remains relatively low, despite a geographic distance of ~ 1000 km. The genetic differentiation observed between north-central and southern Mexico is more likely due to vicariance than IBD.

Our demographic modeling indicated strongly supported isolation with migration, sug-630 gesting ongoing gene flow between these regions (see Table S3). This gene flow was quan-631 tified at 4Nm = 4.64, which translates to an  $Fst \approx 1/(4Nm + 1) = 0.177$ , assuming an 632 infinite island model operating at migration-drift equilibrium. Typically, such Fst values 633 (ranging from 0.15 to 0.25) indicate substantial genetic differentiation [Wright, 1949]. The 634 top PHRAPL models shared identical migration parameters, but their coalescent parame-635 ter estimation varied. This variability can be attributed to the limited phylogenetic signal 636 obtained from ddRAD gene trees and PHRAPL's inherent bias towards models that empha-637 size divergence, making distinguishing between isolation-only and isolation with migration 638 models challenging. This bias persists even with an increasing number of loci [Jackson et al., 639 2017]. Considering all factors, PHRAPL results suggest a short divergence time, with lim-640 ited gene flow homogenizing the observed differentiation between F. petiolaris populations 641 separated by TVB ( $F_{st} \ge 0.292$ ; see Table S1). 642

The *Pegoscapus* populations exhibit strong genetic differentiation throughout the TVB, characterized by  $F_{st}$  values commonly seen between species. The top PHRAPL model supported an isolation-with-migration scenario, which was confirmed by the detection of an F1 hybrid. *snpclust* verified that the F1 hybrid originated from north-central or southern Mexico. Identifying this F1 hybrid within a sample of 102 wasps suggests the prevalence of long-distance dispersal and hybridization among genetic groups. However, detecting migration events does not always equate to effective gene flow between source populations.

<sup>650</sup> Since the hybrid was found in Jalisco, it suggests that a female *Pegoscapus* traveled from

Morelos or northern Oaxaca. Based on field observations, single-foundress broods (65% preva-651 lence) in F. petiolaris often produce multiple generations before admixture. When source 652 populations are deeply divergent genetic incompatibilities result in infertile first-generation 653 hybrids [Orr, 1995]. Therefore, the absence of second-generation hybrids in our 102 samples 654 is unsurprising. The divergence between north-central and southern Mexico highlights their 655 restricted effective gene flow. Genetic differentiation is less pronounced north of the TVB, 656 possibly due to the Sierra Madre Occidental orientation along the western coast, which allows 657 for migration corridors aided by seasonal wind patterns [Adams and Comrie, 1997]. 658

Although Morelos and Oaxaca are less than 300 m apart, there is a high level of ge-659 netic differentiation between their populations, similar to the pairwise  $F_{st}$  values seen in 660 populations separated by the TVB. The variation in genetic connectivity may be due to 661 physiological limitations experienced during traversal of environmental gradients. Heat and 662 humidity constrain fig wasps [Jevanandam et al., 2013; van Kolfschoten et al., 2022; Aung 663 et al., 2022], limiting their navigation abilities. Altitude gradients on Mount Wilhelm in 664 Papua New Guinea have been shown to affect the composition and diversity of wasp com-665 munities [SoutoVilarós et al., 2020]. Elevation changes hinder gene flow between Ficus and 666 pollinator populations, causing genetic divergence with a 500m elevation shift over 4km. 667 [Segar et al., 2017; Souto-Vilarós et al., 2019]. F. petiolaris pollinators in high-altitude en-668 vironments experience lower humidity levels, which increase fig wasp mortality rates [Dunn 669 et al., 2008; Jevanandam et al., 2013]. Our data shows that pollinators prefer lower elevations 670 (Figure S1) despite the altitude range available within their environment. 671

#### <sup>672</sup> 4.5 The Spatial Scale of Gene Flow

We found differences in the spatial symmetry of genetic structures between F. petiolaris 673 and *Peqoscapus* based on a topological test that identified significant disparities in inter-674 population variance (Figure 8). Furthermore, a structural test of population connectivity 675 patterns revealed a significant number of shared edge connections between the host and 676 pollinator population graphs. Considering the population graph results along with PCA 677 and STRUCTURE, there is significant population genetic structure differences between F. 678 petiolaris and Peqoscapus. However, our results also suggest localized genetic connectivity 679 similarities between the host and pollinator. These observed phylogeographic patterns are 680 not unique to our system. The absence of IBD and the lack of spatial genetic structure over 681 extensive geographical ranges are seen in other *Ficus* host-pollinator pairs Molbo et al., 682 2004; Zavodna et al., 2005; Yu et al., 2010; Kobmoo et al., 2010; Lin et al., 2008; Yu and 683 Nason, 2013; Tian et al., 2015; Heer et al., 2015; Bain et al., 2016; Honorio Coronado et al., 684 2019; Wilde et al., 2021]. 685

Various factors contribute to the differences in broad-scale population genetic structure between host and pollinator, These include specific traits and their environment. Life-history traits, physiology, and environment influence fig wasp dispersal, longevity, and community composition. Behavioral factors, such as flight height and emergence times can exert strong influences on the dispersal capabilities of pollinators. Some fig wasp species have adopted a nocturnal flight strategy to avoid high air temperatures during the day, thereby expanding their dispersal range [Warren et al., 2010]. Fig wasps also exhibit behaviors that optimize resource allocation to enhance their fitness [Greeff and Kjellberg, 2022]. Lack of pollination harms fig wasp larvae since figs entered by pollen-free wasps are more likely to abort [Jousselin et al., 2003; Jansen-González et al., 2012; Borges, 2021]. Pollen collection and deposition have an energy cost, so balancing oviposition with pollination is necessary [Kjellberg et al., 2001; Anstett et al., 1997]. Long-distance dispersal should lead to increased allocation of energy resources towards oviposition instead of pollination. Therefore, pollinators are more likely than their host plant to introduce genes within their own populations. [Liu et al., 2015].

The growth patterns of fig tree species can also contribute to the incongruent host and pol-700 linator genetic structure. [Chen et al., 2011]. Dioecious figs are typically small trees or shrubs 701 that are sparsely distributed across the landscape, yet they frequently form relatively dense 702 local populations [Wang et al., 2009]. Additionally, individual plants of dioecious species 703 tend to bear fruit more frequently compared to their monoecious counterparts [Harrison. 704 2003]. As a result, dioecious figs pollinators do not need to disperse as far, leading to a 705 strong population structure among dioecious figs [Wang et al., 2009; Chen et al., 2011; Dev 706 et al., 2011]. In contrast, monoecious figs are thinly scattered (<1 individual per hectare) 707 and predominantly depend on pollinators that rely on wind currents to disperse their pollen. 708 Consequently, monoecious figs tend to exhibit limited or no genetic structure across their 709 species distribution [Kobmoo et al., 2010; Bain et al., 2016; Honorio Coronado et al., 2019; 710 Wilde et al., 2021]. 711

At a local level, we observed similar population connectivity patterns between *F. petiolaris* 712 and *Peqoscapus* likely driven by seed dispersal Heer et al. [2015]. In the case of F. petiolaris, 713 the dispersion of seeds is facilitated by frugivorous bats, which are commonly recognized as 714 effective long-distance seed dispersers [Piedra-Malagón et al., 2019; Thornton et al., 1996]. 715 It is unlikely that seeds ingested by bats are dispersed over long distances as they typically 716 spend less than 30 minutes in the digestive tracts of bats [Morrison, 1980; Laska, 1990]. Most 717 fig seeds are expected to be dropped close to their maternal fig tree [Heer et al., 2015]. It is 718 possible that some *Peqoscapus* pollinators disperse shorter distances, which could contribute 719 to the similar localized population connectivity between host and pollinator. This signal is 720 likely to be very weak in comparison to the gene flow occurring at broader scales. 721

#### <sup>722</sup> 4.6 Implications for Host Fig and Pollinator Classification

Initially described as a single species, ongoing taxonomic debates resulted in the recog-723 nition of a species complex consisting of four morphologically distinct species, including F. 724 petiolaris, F. jaliscana, F palmeri, and F. brandegei. Piedra-Malagón et al. [2011] found only 725 gradual morphological variations among these species, indicating a potential single species. 726 Our results support the classification of F. petiolaris subsp. petiolaris and palmeri, demon-727 strating significant genetic differentiation between Baja California and mainland Mexico 728 clusters. The classification of subsp. *brandeqii* is not supported by our findings. We recognize 729 three distinct allopatric subspecies of F. petiolaris: one in Southern Oaxaca, another in BCP. 730 and a third widespread across northern and central mainland Mexico. 731

There has been no formal taxonomic analysis of its associated *Pegoscapus* pollinator. Similar to the host, genetic differentiation across TVB indicates the presence of distinct pollinator subspecies. Fig populations in Baja California, Inland Sonora, and Central Mexico comprise two subspecies, while minimal pollinator population genetic differentiation indicates a single subspecies. We recognize three subspecies of the *F. petiolaris* pollinator: one
widespread in Baja California and northern+central mainland Mexico, a second in Morelos
and northern Oaxaca, and a third southern Oaxaca.

#### 739 4.7 Conclusion

The phylogeography of *F. petiolaris*, and its pollinator, *Pegoscapus sp.*, demonstrates that biological traits, such as life history strategies, dispersal capabilities, and physiological constraints, are important factors that shape their respective population genetic structure. This study contributes to our understanding of the complex dynamics that shape the genetic variation in obligate mutualistic interactions and highlights the need to consider historical factors along side biological traits to unravel their complex genetic patterns.

### 746 5 Acknowledgments

Field collection was supported by grants from the Center for Global & Regional En-747 vironmental Research at the University of Iowa and Finch Funds from the Department of 748 Ecology, Evolution, and Organismal Biology at Iowa State University. Sequencing and library 749 preparation were funded through a National Science Foundation grant (DEB-1556853). The 750 ResearchIT and High-Performance Computing facility at Iowa State University provided 751 computational resources. We thank Jose Lopez for his assistance with field collection and 752 DNA extractions. Special thanks to the Nason Lab and Heath Lab for their insightful com-753 ments and discussions that greatly enhanced the quality of this manuscript. Lastly, we would 754 like to express our deep appreciation to the communities in Mexico that graciously assisted 755 us in collecting samples. 756

## 757 6 Author Contributions

KQ, FP and JDN conceived of the study. KQ, FP and JDN collected the fig and fig
wasp samples. KQ, JDS and FP generated and processed the sequence data. KQ and JDN
conducted all analyses. KQ, FP, JDS, TAH and JDN wrote the paper, and all authors
contributed to revised versions of the manuscript and approved of the final version.

## 762 7 Data Availability Statement and Benefit-sharing

Raw sequence data are available from the NCBI Sequence Read Archive (SRA) under BioProject ID: XXX (BioSample accessions: XXX). NCBI BioSample accession numbers for individual wasps, Ipyrad and Phyluce assemblies, along with their related script datasets are available on Dryad (https://doi.org/10.5061/dryad.fbg79cnwk). Scripts for figures and data analysis are available in a GitHub repository (https://github.com/kquinteros/ ficus-phylogeography.git).

## 769 8 Benefit-Sharing

Benefits Generated: Benefits from this research accrue from the sharing of our data and results on public databases as described above.

## 772 **References**

Adams, D. K. and A. C. Comrie, 1997. The North American Monsoon. Bulletin of the
 American Meteorological Society 78:2197–2213.

Ahmed, S., S. G. Compton, R. K. Butlin, and P. M. Gilmartin, 2009. Wind-borne insects mediate directional pollen transfer between desert fig trees 160 kilometers apart. Proceedings
of the National Academy of Sciences 106.

Alvarez, N., D. McKey, F. Kjellberg, and M. Hossaert-McKey, 2010. Phylogeography and
historical biogeography of obligate specific mutualisms. Pp. 31–39, *in* The Biogeography
of Host-Parasite Interactions. Oxford University Press.

Andermann, T., A. M. Fernandes, U. Olsson, M. Töpel, B. Pfeil, B. Oxelman, A. Aleixo,
B. C. Faircloth, and A. Antonelli, 2018. Allele phasing greatly improves the phylogenetic
utility of ultraconserved elements. Systematic Biology 68:32–46.

Anguiano-Constante, M. A., P. Zamora-Tavares, E. Ruiz-Sanchez, E. Dean, A. Rodríguez,
and G. Munguía-Lino, 2021. Population differentiation and phylogeography in Lycianthes
moziniana (Solanaceae: Capsiceae), a perennial herb endemic to the Mexican transition
zone. Biological Journal of the Linnean Society 132:359–373.

Anstett, M. C., M. Hossaert-McKey, and F. Kjellberg, 1997. Figs and fig pollinators: evolutionary conflicts in a coevoled mutualism. Trends in Ecology & Evolution 12:94–99.

Aung, K. M. M., H.-H. Chen, S. T. Segar, B.-G. Miao, Y.-Q. Peng, and C. Liu, 2022.
Changes in temperature alter competitive interactions and overall structure of fig wasp communities. Journal of Animal Ecology 91:1303–1315.

Bain, A., R. M. Borges, M. H. Chevallier, H. Vignes, N. Kobmoo, Y. Q. Peng, A. Cruaud,
J. Y. Rasplus, F. Kjellberg, and M. Hossaert-Mckey, 2016. Geographic structuring into
vicariant species-pairs in a wide-ranging, high-dispersal plantinsect mutualism: the case
of Ficus racemosa and its pollinating wasps. Evolutionary Ecology 30:663–684.

Bankevich, A., S. Nurk, D. Antipov, A. A. Gurevich, M. Dvorkin, A. S. Kulikov, V. M. Lesin,
a. S. P. Sergey I. Nikolenko, A. D. Prjibelski, A. V. Pyshkin, A. V. Sirotkin, N. Vyahhi,
G. Tesler, M. A. Alekseyev, and P. A. Pevzner, 2012. SPAdes: a new genome assembly
algorithm and its applications to single-cell sequencing. Journal of Computational Biology
19:455–477.

Becerra, J. X., 2005. Timing the origin and expansion of the Mexican tropical dry forest.
 Proceedings of the National Academy of Sciences 102:10919–10923.

- Beugin, M., T. Gayet, D. Pontier, S. Devillard, and T. Jombart, 2018. A fast likelihood
  solution to the genetic clustering problem. Methods in Ecology and Evolution 9:1006–
  1016.
- Bolger, A. M., M. Lohse, and B. Usadel, 2014. Trimmomatic: A flexible trimmer for Illumina
   Sequence Data. Bioinformatics .
- Borges, R. M., 2021. Interactions between figs and gall-inducing fig wasps: adaptations,
   constraints, and unanswered questions. Frontiers in Ecology and Evolution 9.
- Bowen, B. W. and J. C. Avise, 1990. Genetic structure of atlantic and Gulf of Mexico populations of sea bass, menhaden, and sturgeon: influence of zoogeographic factors and life-history patterns. Marine Biology 107:371–381.
- Branstetter, M. G., J. T. Longino, P. S. Ward, and B. C. Faircloth, 2017. Enriching the
  ant tree of life: enhanced UCE bait set for genomescale phylogenetics of ants and other
  Hymenoptera. Methods in Ecology and Evolution 8:768–776.
- Bronstein, J. L. and M. Hossaert-McKey, 1996. Variation in reproductive success within a
   subtropical fig/pollinator mutualism. Journal of Biogeography 23:433–446.
- Bryson Jr, R. W., R. W. Murphy, A. Lathrop, and D. Lazcano-Villareal, 2011. Evolutionary drivers of phylogeographical diversity in the highlands of Mexico: a case study of
  the Crotalus triseriatus species group of montane rattlesnakes. Journal of Biogeography
  38:697-710.
- Capella-Gutierrez, S., J. M. Silla-Martinez, and T. Gabaldon, 2009. trimAl: a tool for auto mated alignment trimming in large-scale phylogenetic analyses. Bioinformatics 25:1972–
   1973.
- Carstens, B. C., S. J. Brunsfeld, J. R. Demboski, J. M. Good, and J. Sullivan, 2005. Investigating the evolutionary history of the pacific northwest mesic forest ecosystem: hypothesis
  testing within a comparative phylogeographic framework. Evolution 59:15.
- Carstens, B. C. and C. L. Richards, 2007. Integrating coalescent and ecological niche mod eling in comparative phylogeography. Evolution 61:1439–1454.
- Castresana, J., 2000. Selection of conserved blocks from multiple alignments for their use in
   phylogenetic analysis. Molecular Biology and Evolution 17:540–552.
- Cavalli-Sforza, L. L. and A. W. F. Edwards, 1967. Phylogenetic analysis. models and esti mation procedures. American Journal of Human Genetics 19:233–257.
- <sup>835</sup> Chen, C. and Q. Song, 2008. Responses of the pollinating wasp ceratosolen solmsi marchali
   to odor variation between two floral stages of ficus hispida. Journal of Chemical Ecology .

- <sup>837</sup> Chen, Y., S. G. Compton, M. Liu, and X.-Y. Chen, 2012. Fig trees at the northern limit
   of their range: the distributions of cryptic pollinators indicate multiple glacial refugia.
   <sup>839</sup> Molecular Ecology 21:1687–1701.
- Chen, Y., Z.-X. Jiang, S. G. Compton, M. Liu, and X.-Y. Chen, 2011. Genetic diversity and
  differentiation of the extremely dwarf Ficus tikoua in Southwestern China. Biochemical
  Systematics and Ecology 39:441–448.
- Chifman, J. and L. Kubatko, 2014. Quartet inference from SNP data under the coalescent
   model. Bioinformatics 30:3317–3324.
- <sup>845</sup> Cock, P. J. A., J. K. Bonfield, B. Chevreux, and H. Li, 2015. SAM/BAM format v1.5
  <sup>846</sup> extensions for de novo assemblies. Tech. rep., Bioinformatics.
- <sup>847</sup> Cook, J. M. and J.-Y. Rasplus, 2003. Mutualists with attitude: coevolving fig wasps and
   <sup>848</sup> figs. Trends in Ecology & Evolution 18:241–248.
- <sup>849</sup> Cooper, L., L. Bunnefeld, J. Hearn, J. M. Cook, K. Lohse, and G. N. Stone, 2020. Low<sup>850</sup> coverage genomic data resolve the population divergence and gene flow history of an
  <sup>851</sup> Australian rain forest fig wasp. Molecular Ecology 29:3649–3666.
- <sup>852</sup> Crawley, M. J. and G. J. S. Ross, 1990. The population dynamics of plants. Philosophical
   <sup>853</sup> Transactions: Biological Sciences 330:125–140.
- Cruaud, A., N. Rønsted, B. Chantarasuwan, L. S. Chou, W. L. Clement, A. Couloux,
  B. Cousins, G. Genson, R. D. Harrison, P. E. Hanson, M. Hossaert-Mckey, R. JabbourZahab, E. Jousselin, C. Kerdelhué, F. Kjellberg, C. Lopez-Vaamonde, J. Peebles, Y.-Q.
  Peng, R. A. S. Pereira, T. Schramm, R. Ubaidillah, S. van Noort, G. D. Weiblen, D.-R.
  Yang, A. Yodpinyanee, R. Libeskind-Hadas, J. M. Cook, J.-Y. Rasplus, and V. Savolainen,
  2012. An extreme case of plantinsect codiversification: figs and fig-pollinating wasps. Systematic Biology 61:1029–1047.
- <sup>861</sup> Dev, S. A., F. Kjellberg, M. Hossaert-McKey, and R. M. Borges, 2011. Fine-scale Population
   <sup>862</sup> Genetic Structure of Two Dioecious Indian Keystone Species, Ficus hispida and Ficus
   <sup>863</sup> exasperata (Moraceae). Biotropica 43:309–316.
- <sup>864</sup> Dierckxsens, N., P. Mardulyn, and G. Smits, 2016. NOVOPlasty: de novo assembly of
   <sup>865</sup> organelle genomes from whole genome data. Nucleic Acids Research P. gkw955.
- <sup>866</sup> Douglas Nychka, Reinhard Furrer, John Paige, and Stephan Sain, 2017. Fields: tools for
   <sup>867</sup> spatial data.
- Dray, S. and A.-B. Dufour, 2007. The ade4 package: implementing the duality diagram for
   ecologists. Journal of Statistical Software 22:1–20.

- Dunn, D. W., D. W. Yu, J. Ridley, and J. M. Cook, 2008. Longevity, early emergence and 870 body size in a pollinating fig wasp - implications for stability in a fig-pollinator mutualism. 871 Journal of Animal Ecology 77:927–935. 872
- Dyer, R. J., 2015. Population Graphs and Landscape Genetics. Annual Review of Ecology, 873 Evolution, and Systematics 46:327–342.

874

- -, 2021. popgraph: This is an R package that constructs and manipulates population 875 graphs. 876
- Dyer, R. J., D. M. Chan, V. A. Gardiakos, and C. A. Meadows, 2012. Pollination graphs: 877 quantifying pollen pool covariance networks and the influence of intervening landscape on 878 genetic connectivity in the North American understory tree, Cornus florida L. Landscape 879 Ecology 27:239–251. 880
- Dyer, R. J. and J. D. Nason, 2004. Population Graphs: the graph theoretic shape of genetic 881 structure. Molecular Ecology 13:1713–1727. 882
- Eaton, D. A. R. and I. Overcast, 2020. Ipyrad: Interactive assembly and analysis of RADseq 883 datasets. Bioinformatics 36:2592–2594. 884
- Ehrlich, P. R. and P. H. Raven, 1964. Butterflies and plants: a study in coevolution. Evolution 885 18:586-608. 886
- Ennos, R. A., 1994. Estimating the relative rates of pollen and seed migration among plant 887 populations. Heredity 72:250–259. 888
- Evanno, G., S. Regnaut, and J. Goudet, 2005. Detecting the number of clusters of individuals 889 using the software structure: a simulation study. Molecular Ecology 14:2611–2620. 890
- Faircloth, B. C., 2013. Illumiprocessor: a trimmomatic wrapper for parallel adapter and 891 quality trimming. http://dx.doi.org/10.6079/J9ILL. . 892
- , 2016. PHYLUCE is a software package for the analysis of conserved genomic loci. 893 Phylogenetics 32:786–788. 894
- Faircloth, B. C., J. E. McCormack, N. C. Crawford, M. G. Harvey, R. T. Brumfield, and 895 T. C. Glenn, 2012. Ultraconserved elements anchor thousands of genetic markers spanning 896 multiple evolutionary timescales. Systematic Biology 61:717–726. 897
- Ferrari, L., M. Bergomi, M. Martini, A. Tunesi, T. Orozco-Esquivel, and M. López-898 Martínez, 2014. Late Cretaceous-Oligocene magmatic record in southern Mex-899 for a temporal slab window along ico: The case the evolving Caribbean-900 North America-Farallon triple boundary. Tectonics 33:1738-1765. eprint: 901 https://onlinelibrary.wiley.com/doi/pdf/10.1002/2014TC003525. 902

- Ferrari, L., T. Orozco-Esquivel, V. Manea, and M. Manea, 2012. The dynamic history of
   the Trans-Mexican Volcanic Belt and the Mexico subduction zone. Tectonophysics 522 523:122-149.
- Francis, R. M., 2017. POPHELPER: an R package and web app to analyse and visualize
   population structure. Molecular Ecology Resources 17:27–32.
- Galil, J. and G. Neeman, 1977. Pollen transfer and pollination in the common fig (Ficus carica l.). New Phytologist 79:163–171.
- Garrick, R. C., J. D. Nason, J. F. Fernández-Manjarrés, and R. J. Dyer, 2013. Ecological
  coassociations influence species' responses to past climatic change: an example from a
  Sonoran Desert bark beetle. Molecular Ecology 22:3345–3361.
- Garrick, R. C., J. D. Nason, C. A. Meadows, and R. J. Dyer, 2009. Not just vicariance:
  phylogeography of a Sonoran Desert euphorb indicates a major role of range expansion
  along the Baja peninsula. Molecular Ecology 18:1916–1931.
- Gates, D. J. and J. D. Nason, 2012. Flowering asynchrony and mating system effects on
  reproductive assurance and mutualism persistence in fragmented fig-fig wasp populations.
  American Journal of Botany 99:757–768.
- Gomulkiewicz, R., S. Nuismer, and J. Thompson, 2003. Coevolution in variable mutualisms.
   The American Naturalist 162:80–93.
- Gomulkiewicz, R., J. N. Thompson, R. D. Holt, S. L. Nuismer, and M. E. Hochberg, 2000.
  Hot spots, cold spots, and the geographic mosaic theory of coevolution. The American
  Naturalist 156:156–174.
- Goudet, J., 2005. hierfstat, a package for R to compute and test hierarchical F-statistics.
  Molecular Ecology Notes 5:184–186.
- <sup>926</sup> Goudet, J. and T. Jombart, 2020. Hierfstat: estimation and tests of hierarchical F-statistics.

Graham, M. R., R. W. Bryson, and B. R. Riddle, 2014. Late Pleistocene to Holocene distributional stasis in scorpions along the Baja California peninsula: Distributional Stasis in Baja California Scorpions. Biological Journal of the Linnean Society 111:450–461.

- Greeff, J. M. and F. Kjellberg, 2022. Pollinating fig wasps simple solutions to complex sex
   ratio problems: a review. Frontiers in Zoology 19:3.
- Grismer, L. L., 1994. The origin and evolution of the peninsular herpetofauna of Baja
  California, México. Herpetological Natural History 2:51–106.
- Gutiérrez-Ortega, J. S., M. M. Salinas-Rodríguez, J. F. Martínez, F. Molina-Freaner, M. A.
  Pérez-Farrera, A. P. Vovides, Y. Matsuki, Y. Suyama, T. A. Ohsawa, Y. Watano, and
  T. Kajita, 2018. The phylogeography of the cycad genus Dioon (Zamiaceae) clarifies its

- <sup>937</sup> Cenozoic expansion and diversification in the Mexican transition zone. Annals of Botany
   <sup>938</sup> 121:535-548.
- Halffter, G., 1964. La entomofauna americana, ideas acerca de su origen y distribución.
  Sociedad Mexicana de Entomología.
- 941 —, 1987. Biogeography of the montane entomofauna of Mexico and Central America.
   942 Annual review of entomology 32:95–114.
- Harrison, R. D., 2000. Repercussions of El Niño: drought causes extinction and the breakdown of mutualism in Borneo. Proceedings of the Royal Society B: Biological Sciences
  267:911–915.
- 946 —, 2003. Fig wasp dispersal and the stability of a keystone plant resource in Borneo.
   947 Proceedings of the Royal Society of London. Series B: Biological Sciences 270.
- Harrison, R. D. and J.-Y. Rasplus, 2006. Dispersal of fig pollinators in Asian tropical rain
  forests. Journal of Tropical Ecology 22:631–639.
- <sup>950</sup> Heer, K., E. K. V. Kalko, L. Albrecht, R. García-Villacorta, F. C. Staeps, E. A. Herre, and
- 951 C. W. Dick, 2015. Spatial scales of genetic structure in free-standing and strangler figs
- (Ficus, Moraceae) inhabiting neotropical forests. PLOS ONE 10:e0133581.
- Hembry, D. H. and D. M. Althoff, 2016. Diversification and coevolution in brood pollination
  mutualisms: Windows into the role of biotic interactions in generating biological diversity.
  American journal of botany 103:1783–1792.
- Hernández-Esquivel, K. B., E. M. Piedra-Malagón, G. Cornejo-Tenorio, L. Mendoza-Cuenca,
  A. González-Rodríguez, E. Ruíz-Sánchez, and a. G. Ibarra-Manríquez, 2020. Unraveling
  the extreme morphological variation in the neotropical Ficus aurea complex (subg. Spherosuke, sect. Americanae, Moraceae). Journal of Systematics and Evolution 58:263–281.
- Herre, E. A., K. C. Jandér, and C. A. Machado, 2008. Evolutionary ecology of figs and their
  associates: recent progress and outstanding puzzles. Annual Review of Ecology, Evolution,
  and Systematics 39:439–458.
- Hess, H. H., 1962. History of Ocean Basins. in Petrologic Studies. Geological Society of
   America.
- <sup>965</sup> Hewitt, G., 2000. The genetic legacy of the Quaternary ice ages. Nature 405:907–913.
- <sup>966</sup> Hochberg, Gomulkiewicz, Holt, and Thompson, 2000. Weak Sinks Could Cradle Mutualistic
- <sup>967</sup> Symbioses Strong Sources Should Harbour Parasitic Symbioses. Journal of Evolutionary
- <sup>968</sup> Biology 13:213–222.
- <sup>969</sup> Honorio Coronado, E. N., K. G. Dexter, M. L. Hart, O. L. Phillips, and R. T. Pennington,
  <sup>970</sup> 2019. Comparative phylogeography of five widespread tree species: Insights into the history
  <sup>971</sup> of western Amazonia. Ecology and Evolution 9:7333–7345.

<sup>972</sup> Honorio Coronado, E. N., K. G. Dexter, M. F. Poelchau, P. M. Hollingsworth, O. L. Phillips,

- R. T. Pennington, and M. Carine, 2014. Ficus insipida subsp. insipida (Moraceae) reveals
- <sup>974</sup> the role of ecology in the phylogeography of widespread Neotropical rain forest tree species.
- Journal of Biogeography 41:1697–1709.
- <sup>976</sup> Hurtado, L., T. Erez, S. Castrezana, and T. Markow, 2004. Contrasting population ge<sup>977</sup> netic patterns and evolutionary histories among sympatric Sonoran Desert cactophilic
  <sup>978</sup> Drosophila. Molecular Ecology 13:1365–1375.
- Jackson, N. D., A. E. Morales, B. C. Carstens, and B. C. OMeara, 2017. PHRAPL: phylogeographic inference using approximate likelihoods. Systematic Biology 66:1045–1053.
- Jansen-González, S., S. d. P. Teixeira, and R. A. S. Pereira, 2012. Mutualism from the
  inside: coordinated development of plant and insect in an active pollinating fig wasp.
  Arthropod-Plant Interactions 6:601–609.
- Janzen, D. H., 1979. How to be a Fig. Annual Review of Ecology and Systematics 10:13–51.
- Jevanandam, N., A. G. R. Goh, and R. T. Corlett, 2013. Climate warming and the potential extinction of fig wasps, the obligate pollinators of figs. Biology Letters 9:20130041.
- Jombart, T., 2008. adegenet: a R package for the multivariate analysis of genetic markers.
  Bioinformatics 24:1403–1405.
- Jombart, T. and I. Ahmed, 2011. adegenet 1.3-1: new tools for the analysis of genome-wide SNP data. Bioinformatics .
- Jousselin, E., J.-Y. Rasplus, and F. Kjellberg, 2003. Convergence and coevolution in a
   mutualism: evidence from a molecular phylogeny of Ficus. Evolution; International Journal
   of Organic Evolution 57:1255–1269.
- Katoh, K. and D. M. Standley, 2013. MAFFT multiple sequence alignment software version
  7: improvements in performance and usability. Molecular Biology and Evolution 30:772–
  780.
- <sup>997</sup> Kjellberg, F., B. Doumesche, and J. L. Bronstein, 1988. Longevity of a fig wasp (Blastophaga
   <sup>998</sup> psenes). Proceedings of the Koninklijke Nederlandse Akademie van Wetenschappen : Series
   <sup>999</sup> C : Biological and medical sciences .
- Kjellberg, F., E. Jousselin, J. L. Bronstein, A. Patel, J. Yokoyama, and J.-Y. Rasplus, 2001.
   Pollination mode in fig wasps: the predictive power of correlated traits. Proceedings of
   the Royal Society of London. Series B: Biological Sciences 268:1113–1121.
- Knowles, L. L., 2009. Statistical phylogeography. Annual Review of Ecology, Evolution, and
   Systematics 40:593–612.

Kobmoo, N., M. Hossaert-Mckey, J. Y. Rasplus, and F. Kjellberg, 2010. Ficus racemosa is
 pollinated by a single population of a single agaonid wasp species in continental south-east
 asia: Ficus racemosa have a single agaonid species. Molecular Ecology 19:2700–2712.

van Kolfschoten, L., L. Dück, M. I. Lind, and K. C. Jandér, 2022. Rising temperatures
 threaten pollinators of fig treesKeystone resources of tropical forests. Ecology and Evolution 12:e9311.

Krishnan, A., G. K. Pramanik, S. V. Revadi, V. Venkateswaran, and R. M. Borges, 2014.
High Temperatures Result in Smaller Nurseries which Lower Reproduction of Pollinators and Parasites in a Brood Site Pollination Mutualism. PLOS ONE 9:e115118. Publisher:
Public Library of Science.

 Lamb, T., T. R. Jones, and J. C. Avise, 1992. Phylogeographic histories of representative herpetofauna of the southwestern U.S.: mitochondrial DNA variation in the desert iguana (Dipsosaurus dorsalis) and the chuckwalla (Sauromalus obesus). Journal of Evolutionary Biology 5:465–480.

Laska, M., 1990. Food transit times and carbohydrate use in three Phyllostomid bat species.
 Z Säeugetierkunde 50:49–54.

Li, H., 2013. Aligning sequence reads, clone sequences and assembly contigs with bwa-mem. Genomics preprint:arXiv:1303.3997.

Li, H. and R. Durbin, 2010. Fast and accurate long-read alignment with BurrowsWheeler transform. Bioinformatics 26:589–595.

Li, H., B. Handsaker, A. Wysoker, T. Fennell, J. Ruan, N. Homer, G. Marth, G. Abecasis, R. Durbin, and 1000 Genome Project Data Processing Subgroup, 2009. The sequence alignment/map format and samtools. Bioinformatics 25:2078–2079.

Lin, R.-C., C. K.-L. Yeung, and S.-H. Li, 2008. Drastic Post-LGM expansion and lack of historical genetic structure of a subtropical fig-pollinating wasp (Ceratosolen sp. 1) of Ficus septica in Taiwan. Molecular Ecology 17:5008–5022.

Liu, M., S. G. Compton, F.-E. Peng, J. Zhang, and X.-Y. Chen, 2015. Movements of genes between populations: are pollinators more effective at transferring their own or plant genetic markers? Proceedings of the Royal Society B: Biological Sciences 282.

Machado, C. A., N. Robbins, M. T. P. Gilbert, and E. A. Herre, 2005. Critical review of host
 specificity and its coevolutionary implications in the fig/fig-wasp mutualism. Proceedings
 of the National Academy of Sciences 102:6558–6565.

Mastretta-Yanes, A., A. Moreno-Letelier, D. Piñero, T. H. Jorgensen, and B. C. Emerson,
 2015. Biodiversity in the Mexican highlands and the interaction of geology, geography and
 climate within the Trans-Mexican Volcanic Belt. Journal of Biogeography 42:1586–1600.

- McKey, D., 1989. Population biology of figs: Applications for conservation. Experientia
   45:661–673.
- de Medeiros, B. A. S. and B. D. Farrell, 2020. Evaluating insect-host interactions as a driver of species divergence in palm flower weevils. Communications Biology 3:1–9.
- Molbo, D., C. A. Machado, E. A. Herre, and L. Keller, 2004. Inbreeding and population
   structure in two pairs of cryptic fig wasp species: inbreeding in cryptic fig wasps. Molecular
   Ecology 13:1613–1623.
- Molbo, D., C. A. Machado, J. G. Sevenster, L. Keller, and E. A. Herre, 2003. Cryptic
  species of fig-pollinating wasps: Implications for the evolution of the fig-wasp mutualism,
  sex allocation, and precision of adaptation. Proceedings of the National Academy of
  Sciences 100:5867–5872.
- <sup>1051</sup> Morrison, D. W., 1980. Foraging and Day-Roosting Dynamics of Canopy Fruit Bats in
   <sup>1052</sup> Panama. Journal of Mammalogy 61:20–29.
- Moussalli, A., C. Moritz, S. E. Williams, and A. C. Carnaval, 2009. Variable responses of
   skinks to a common history of rainforest fluctuation: concordance between phylogeography
   and palaeo-distribution models. Molecular Ecology 18:483–499.
- Murphy, R., 1983. Paleobiogeography and genetic differentiation of the Baja California
   herpetofauna. Occasional Papers of the California Academy of Sciences 137:1–48.
- Nason, J. D., J. L. Hamrick, and T. H. Fleming, 2002. Historical vicariance and postglacial
   colonization effects on the evolution of genetic structure in Lophocereus, a Sonoran Desert
   columnar cactus. Evolution 56:2214–2226.
- Nason, J. D., E. A. Herre, and J. L. Hamrick, 1998. The breeding structure of a tropical
   keystone plant resource. Nature 391:685–687.
- Nei, M., F. Tajima, and Y. Tateno, 1983. Accuracy of estimated phylogenetic trees from
   molecular data. Journal of Molecular Evolution 19:153–170.
- Nuismer, S. L., J. N. Thompson, and R. Gomulkiewicz, 1999. Gene flow and geographically
   structured coevolution. Proceedings of the Royal Society B: Biological Sciences 266:605.
- 1067 —, 2000. Coevolutionary clines across selection mosaics. Evolution 54:1102–1115. Pub 1068 lisher: The Society for the Study of Evolution.
- Orr, H. A., 1995. The population genetics of speciation: the evolution of hybrid incompati bilities. Genetics 139:1805–1813.
- Papadopoulou, A. and L. L. Knowles, 2016. Toward a Paradigm Shift in Comparative
   Phylogeography Driven by Trait-Based Hypotheses. Proceedings of the National Academy
   of Sciences 113:8018–8024.

Pellmyr, O., F. Kjellberg, E. A. Herre, A. Kawakita, D. H. Hembry, J. N. Holland, T. Terrazas, W. Clement, K. A. Segraves, and D. M. Althoff, 2020. Active pollination drives selection for reduced pollenovule ratios. American Journal of Botany 107:164–170.

Peterson, B. K., J. N. Weber, E. H. Kay, and H. H. H.S. Fisher, 2012. Double digest RADseq:
an inexpensive method for de novo SNP discovery and genotyping in model and non-model
species. PLoS One .

- Petit, R. J., J. Duminil, S. Fineschi, A. Hampe, D. Salvini, and G. G. Vendramin, 2005.
   Invited review: comparative organization of chloroplast, mitochondrial and nuclear diversity in plant populations: organization of genetic diversity in plants. Molecular Ecology 14:689–701.
- Pfeiler, E., T. Erez, L. Hurtado, and T. Markow, 2007. Genetic differentiation and demo graphic history in Drosophila pachea from the Sonoran Desert. Hereditas 144:63–74.

Piedra-Malagón, E. M., B. Hernández-Ramos, A. Mirón-Monterrosas, G. Cornejo-Tenorio,
 A. Navarrete-Segueda, and G. Ibarra-Manríquez, 2019. Syconium development in *Ficus petiolaris* (*Ficus*, sect. americanae, moraceae) and the relationship with pollinator and
 parasitic wasps. Botany 97:190–203.

- Piedra-Malagón, E. M., V. Sosa, and G. Ibarra-Manríquez, 2011. Clinal variation and species
   boundaries in the *Ficus petiolaris* complex (Moraceae). Systematic Botany 36:80–87.
- Pritchard, J. K., M. Stephens, N. A. Rosenberg, and P. Donnelly, 2000. Association mapping
   in structured populations. The American Journal of Human Genetics 67:170–181.
- Pyron, A. R. and F. T. Burbrink, 2010. Hard and soft allopatry: physically and ecologi cally mediated modes of geographic speciation: modes of allopatric speciation. Journal of
   Biogeography .
- <sup>1097</sup> R Core Team, 2020. R: a language and environment for statistical computing.
- <sup>1098</sup> Ramírez, W. B., 1970. Host specificity of fig wasps (Agaonidae). Evolution 24:13.
- <sup>1099</sup> Ramírez B., W., 1969. Fig wasps: mechanism of pollen transfer. Science 163:580–581.
- Riddle, B. R., D. J. Hafner, and L. F. Alexander, 2000a. Comparative Phylogeography of
  Baileys' Pocket Mouse (Chaetodipus baileyi) and the Peromyscus eremicus Species Group:
  Historical Vicariance of the Baja California Peninsular Desert. Molecular Phylogenetics
  and Evolution 17:161–172.
- -----, 2000b. Phylogeography and Systematics of the Peromyscus eremicus Species Group
   and the Historical Biogeography of North American Warm Regional Deserts. Molecular
   Phylogenetics and Evolution 17:145–160.

Riddle, B. R., D. J. Hafner, L. F. Alexander, and J. R. Jaeger, 2000c. Cryptic vicariance in
the historical assembly of a Baja California Peninsular Desert biota. Proceedings of the
National Academy of Sciences 97:14438–14443.

Rocha-Méndez, A., L. A. Sánchez-González, C. González, and A. G. Navarro-Sigüenza, 2019.
The geography of evolutionary divergence in the highly endemic avifauna from the Sierra Madre del Sur, Mexico. BMC Evolutionary Biology 19:237.

Ross, C. and T. Markow, 2006. Microsatellite variation among diverging populations of
 Drosophila mojavensis. Journal of Evolutionary Biology 19:1691–1700.

Ruffley, M., M. L. Smith, A. Espíndola, B. C. Carstens, J. Sullivan, and D. C. Tank, 2018.
Combining allele frequency and tree-based approaches improves phylogeographic inference
from natural history collections. Molecular Ecology 27:1012–1024.

Rønsted, N., G. D. Weiblen, J. M. Cook, N. Salamin, C. A. Machado, and V. Savolainen,
2005. 60 million years of co-divergence in the figwasp symbiosis. Proceedings of the Royal
Society B: Biological Sciences 272:2593–2599.

Satler, J. D. and B. C. Carstens, 2017. Do ecological communities disperse across biogeographic barriers as a unit? Molecular Ecology 26:3533–3545.

1123 —, 2019. The *Sarracenia alata* pitcher plant system and obligate arthropod inquilines 1124 should be considered an evolutionary community. Journal of Biogeography 46:485–496.

Satler, J. D., E. A. Herre, K. C. Jandér, D. A. R. Eaton, C. A. Machado, T. A. Heath, and
 J. D. Nason, 2019. Inferring processes of coevolutionary diversification in a community of
 Panamanian strangler figs and associated pollinating wasps\*. Evolution 73:2295–2311.

<sup>1128</sup> Savage, J. M., 1960. Evolution of a peninsular herpetofauna. Systematic Zoology 9:184–212.

Sedlock, R. L., 2003. Geology and tectonics of the Baja California Peninsula and adjacent
areas. *in* Tectonic evolution of northwestern Mexico and the Southwestern USA. Geological
Society of America.

Segar, S. T., M. Volf, J. Zima Jnr, B. Isua, M. Sisol, L. Sam, K. Sam, D. Souto-Vilarós,
and V. Novotny, 2017. Speciation in a keystone plant genus is driven by elevation: a case
study in New Guinean Ficus. Journal of Evolutionary Biology 30:512–523.

Smith, C. I., S. Tank, W. Godsoe, J. Levenick, E. Strand, T. Esque, and O. Pellmyr, 2011.
Comparative phylogeography of a coevolved community: concerted population expansions
in joshua trees and four Yucca moths. PLoS ONE 6.

Soltis, D. E., A. B. Morris, J. S. McLachlan, P. S. Manos, and P. S. Soltis, 2006. Comparative phylogeography of unglaciated eastern North America: phylogeography of unglaciated eastern north america phylogeography in a pitcher plant system. Molecular Ecology 15:4261–4293.

- Souto-Vilarós, D., A. Machac, J. Michalek, C. T. Darwell, M. Sisol, T. Kuyaiva, B. Isua,
  G. D. Weiblen, V. Novotny, and S. T. Segar, 2019. Faster speciation of fig-wasps than
  their host figs leads to decoupled speciation dynamics: Snapshots across the speciation
  continuum. Molecular Ecology 28:3958–3976.
- SoutoVilarós, D., M. Houadria, J. Michalek, M. Sisol, B. Isua, T. Kuyaiva, G. D. Weiblen,
  V. Novotny, and S. T. Segar, 2020. Contrasting patterns of fig wasp communities along
  Mt. Wilhelm, Papua New Guinea. Biotropica 52:323–334.
- Staddon, S. C., S. G. Compton, and A. Portch, 2010. Dispersal of fig seeds in the Cook
  Islands: introduced frugivores are no substitutes for natives. Biodiversity and 19:1905–
  1916.
- Stamatakis, A., 2014. RAxML version 8: a tool for phylogenetic analysis and post-analysis
   of large phylogenies. Bioinformatics 30:1312–1313.
- Su, Z.-H., H. Iino, K. Nakamura, A. Serrato, and K. Oyama, 2008. Breakdown of the one to-one rule in Mexican fig-wasp associations inferred by molecular phylogenetic analysis.
   Symbiosis 45:9.
- <sup>1157</sup> Swofford, D., 2003. PAUP\*: phylogenetic analysis using parsimony (and other <sup>1158</sup> method)(Version 4.0 bl0) EM/OL] Sinauer Associates, Sunderland-Massachusetts.
- Thioulouse, J., S. Dray, A.-B. Dufour, A. Siberchicot, T. Jombart, and S. Pavoine, 2018.
   Multivariate Analysis of Ecological Data with ade4.
- Thompson, J. N., 1994. The Coevolutionary Process. University of Chicago Press, Chicago,
   IL.
- <sup>1163</sup> —, 1999. Specific hypotheses on the geographic gosaic of coevolution. The American <sup>1164</sup> Naturalist 153:S1–S14.
- Thompson, J. N. and B. M. Cunningham, 2002. Geographic structure and dynamics of coevolutionary selection. Nature 417:735–738.
- Thornton, I. W. B., S. G. Compton, and C. N. Wilson, 1996. The role of animals in the
   colonization of the Krakatau Islands by fig trees (Ficus species). Journal of Biogeography
   23:577-592.
- Tian, E., J. D. Nason, C. A. Machado, L. Zheng, H. Yu, and F. Kjellberg, 2015. Lack of genetic isolation by distance, similar genetic structuring but different demographic histories
  in a fig-pollinating wasp mutualism. Molecular Ecology 24:5976–5991.
- <sup>1173</sup> Upton, D. E. and R. W. Murphy, 1997. Phylogeny of the Side-Blotched Lizards (Phrynoso-<sup>1174</sup> matidae:Uta) Based on mtDNA Sequences: Support for a Midpeninsular Seaway in Baja <sup>1175</sup> California. Molecular Phylogenetics and Evolution 8:104–113.

Vasconcellos, M. M., G. R. Colli, J. N. Weber, E. M. Ortiz, M. T. Rodrigues, and D. C.
Cannatella, 2019. Isolation by instability: Historical climate change shapes population
structure and genomic divergence of treefrogs in the Neotropical Cerrado savanna. Molecular Ecology 28:1748–1764.

Vieira, F. d. A., R. M. L. Novaes, C. G. Fajardo, R. M. d. Santos, H. d. S. Almeida, D. d. Carvalho, and M. B. Lovato, 2015. Holocene southward expansion in seasonally dry tropical forests in South America: phylogeography of *Ficus bonijesulapensis* (Moraceae): Phylogeography and Niche Modelling of a Fig Tree. Botanical Journal of the Linnean Society 177:189–201.

Vázquez-Miranda, H., R. M. Zink, and B. J. Pinto, 2022. Comparative phylogenomic patterns
in the Baja California avifauna, their conservation implications, and the stages in lineage
divergence. Molecular Phylogenetics and Evolution 171:107466.

Wang, G., C. H. Cannon, and J. Chen, 2016. Pollinator sharing and gene flow among closely
 related sympatric dioecious fig taxa. Proceedings of the Royal Society B 283.

Wang, G., S. G. Compton, and J. Chen, 2013. The mechanism of pollinator specificity
between two sympatric fig varieties: a combination of olfactory signals and contact cues.
Annals of Botany 111.

<sup>1193</sup> Wang, R., B. Ai, B.-Q. Gao, S. Yu, Y.-Y. Li, and X.-Y. Chen, 2009. Spatial genetic structure and restricted gene flow in a functionally dioecious fig, Ficus pumila L. var. pumila (Moraceae). Population Ecology 51:307–315.

<sup>1196</sup> Warren, M., M. P. Robertson, and J. M. Greeff, 2010. A comparative approach to understanding factors limiting abundance patterns and distributions in a fig treefig wasp <sup>1198</sup> mutualism. Ecography 33:148–158.

<sup>1199</sup> Weiblen, G. D., 2002. How to be a fig wasp. Annual Review Entomology 61:299–330.

Weir, B. S. and C. C. Cockerham, 1984. Estimating F-statistics for the analysis of population
 structure. Evolution 38:1358–1370.

Wilde, B. C., S. Rutherford, J.-Y. S. Yap, and M. Rossetto, 2021. Allele Surfing and Holocene
Expansion of an Australian Fig (FicusMoraceae). Diversity 13:250. Number: 6 Publisher:
Multidisciplinary Digital Publishing Institute.

<sup>1205</sup> Wright, S., 1949. The genetical structure of populations. Annals of Eugenics 15:323–354.

Yu, H. and J. D. Nason, 2013. Nuclear and chloroplast DNA phylogeography of Ficus hirta:
obligate pollination mutualism and constraints on range expansion in response to climate
change. New Phytologist 197:276–289.

Yu, H., J. D. Nason, X. Ge, and J. Zeng, 2010. Slatkins Paradox: when direct observation and realized gene flow disagree. A case study in Ficus. Molecular Ecology 19:4441–4453. Zavodna, M., P. Arens, P. J. Van Dijk, T. Partomihardjo, B. Vosman, and J. M. M.
Van Damme, 2005. Pollinating fig waSPS: genetic consequences of island recolonization.
Journal of Evolutionary Biology 18:1234–1243.

Zink, R. M., 1996. Comparative phylogeography in North American birds. Evolution 50:308–
 317.

## 1216 9 Figures



Figure 1: *Ficus petiolaris* and *Pegoscapus sp.* sample sites and known biogeographical barriers in Mexico. The geographical distribution of the fig and its obligately-associated pollinator are indicated by the green shading. *A priori* vicariance hypotheses are based on biographical barriers: (A) ancient mid-peninsular seaway (MPS, ca. 1 Mya), (B) formation of the Isthmus of La Paz (ILP, ca. 3 Mya), (C) the Gulf of California (GOC, ca. 5 Mya), (D) the Trans-Mexican Volcanic Belt (TVB, ca. 11-3 Mya), and (E) the Sierra Madre del Sur (SMS, ca. 48-23 Mya). Ficus petiolaris geographical distribution is based on Global Biodiversity Information Facility (GBIF) occurrence data.



1217

Figure 2: Population genetic structure of *Ficus petiolaris* individuals, colored according to the region in which they were sampled. (A) STRUCTURE analysis revealed K = 2 genetic clusters corresponding to Baja California and Inland Sonora + Central Sonora + Oaxaca, with coastal Sonora admixed between them. Subsequent STRUCTURE analyses of each genetic cluster revealed no additional subdivision. Principal component analysis (PCA) revealed five distinct genetic clusters with Baja California, coastal Sonora, inland Sonora + central Mexico, and Oaxaca separated by PC 1 versus PC 2, which together explained more than 50% of the variation among samples (B), and inland Sonora and central Mexico further differentiated by PC 1 versus PC 3 (C). As in the STRUCTURE analysis, coastal Sonora is intermediate between Baja California and mainland clusters, specifically inland Sonora and Central Mexico. Both STRUCTURE and the PCA support the hypothesis that the Gulf of California is a significant source of vicariance in this species (Figure 1C). The PCA further identifies the Trans-Mexican Volcanic Belt (Figure 1D) as a significant source of vicariance. Pairwise analyses using Wright's  $F_{st}$  subsequently showed the five clusters identified by the PCA to be significantly genetically differentiated from each other (Table S1).



Figure 3: SVDquartets population trees for *Ficus petiolaris* and its *Pegoscapus sp.* pollinator, in which individuals are grouped into populations corresponding to genetic clusters inferred from PCA and STRUCTURE analyses (Figures 2 and 3, respectively). Bootstrap values are shown at the nodes. In the host fig tree (top), consistent with SNAPP results (Figure 4), southern and north-central Mexico form monophyletic clades and, within the latter, coastal Sonora is sister to Baja California while inland Sonora is sister to central Mexico. In the pollinator (bottom), all northern and central Mexico clades were grouped into a single North (Mexico) population based on our PCA and STRUCTURE results The North Mexico phylogroup was sister to all southern Mexico phylogroups. Among the southern Mexico phylogroups, Oaxaca was sister to Northern Oaxaca and Morelos. SVDquartets also provide high support for Northern Oaxaca being sister to Morelos.



Figure 4: In *F. petiolaris*, weak but significant isolation by distance (IBD) was observed across Baja California. This result does not support hypotheses A and B in Figure 1, that two ancient trans-peninsular seaways have been lasting sources of vicariance in this region.



Figure 5: Expected heterozygosity of host fig and pollinator populations as a function of latitude. (A) In *F. petiolaris*, Baja California and Oaxaca populations, which are peninsular and located at the southern limits of the species range, respectively, had similarly low genetic diversity. In contrast, Coastal Sonora, Inland Sonora, and Central Mexico had substantially higher genetic diversity. (B) In *Pegoscapus sp.*, there were no discernable regional differences in genetic diversity and expected heterozygosity varied widely among populations within each region and between regions. Main clusters distinguishes between sites from inferred genetic clusters located north and south of the Trans-Mexico Volcanic Belt.



Figure 6: Population genetic structure of *Peqoscapus sp.* individuals, which are colored according to the region in which they were sampled.(A) Two genetic clusters (K = 2) were inferred by STRUCTURE, with individuals from Morelos and Oaxaca, in the south, forming one cluster and all more northern samples forming the other. One individual from Jalisco had equal admixture from these two genetic clusters. (B) In a principal component analysis (PCA), axis PC 1 separated individuals into two major genetic clusters, northern and central Mexico versus southern Mexico, and explained nearly 50% of variation among samples. The second PCA axis separated the southern samples into three clusters, Morelos, Northern Oaxaca, and Southern Oaxaca, for four genetic clusters total. As in the STRUCTURE analysis, the PCA identifies an apparently admixed individual from Jalisco, subsequently shown using *snapclust* to be a F1 hybrid between northern-central Mexico and southern Mexico clusters. (C) STRUCTURE was subsequently run on each of the two previously inferred clusters (northern + central Mexico; southern Mexico), but only southern Mexico showed further subdivision, with K = 3 clusters corresponding to Morelos, Northern Oaxaca, and Southern Oaxaca. Consistent the PCA analysis, the STRUCTURE analyses revealed K = 4distinct genetic clusters total. (D) PCA analysis of samples from southern Mexico further differentiated genetic clusters from Morelos, Northern Oaxaca, and Southern Oaxaca. Pairwise analyses using Wright's  $F_{st}$  subsequently showed the four clusters identified by both STRUCTURE and PCA analyses to be significantly genetically differentiated from each other (Table S4).



Figure 7: Isolation by distance (IBD) analysis of landscape-level genetic connectivity regressing population-pairwise Nei's genetic distance versus geographic distance for *Pegoscapus sp.* pollinator. Each plot shows the best fit regression line, correlation, Mantel p-value, and slope. Weak but significant IBD was detected across the inferred phylogroup comprising Baja California and northern and central mainland Mexico (A), whereas a nested analysis of Baja, Sonora, and Sinaloa samples and spanning the Gulf of California was not significant (B). In southern Mexico (C), strong and significant IBD was detected across samples from Morelos, northern Oaxaca, and southern Oaxaca, despite their having been identified as distinct phylogroups by STRUCTURE and PCA analyses. (D)Analysis of IBD between *Pegoscapus sp.* populations located north and south of the Trans-Mexican Volcanic Belt, a hypothetical source of vicariance (Figure 1D). IBD between these two regions was not significant, indicating that this area of active volcanos and high plateaus is a strong barrier to dispersal and gene flow in this species.



Figure 8: Population graphs representing the genetic relationships among F. petiolaris (A) and Pegoscapus (B) regional sites. Each node represent sampling sites, and the color of each node corresponds to the regional site where the sampling site is located. Node sizes reflect within-population genetic variability, whereas the edge lengths represent the among-population component of genetic variation due to the connecting nodes. A significant number (permutation test p=0.0048, edge correlation=0.2696, reps=10,000) of edges (colored in red) were shared than expected by chance between F. petiolaris and Pegoscapus population graphs. All shared edges are between neighboring sites, except those involving site 39. Site 39 is the southernmost Baja site yet it shares edges with the two most northerly Baja sites, 172 and 158. These results imply a broad difference in genetic structure between F. petiolaris and Pegoscapus with localized similarities in population genetic structure.

## 1219 10 Supplemental Figures



Figure S1: Distribution of Permutated Elevation (p = 0.037,  $mean_{obs} = 1559.69$  m, 999 permutations). The data shows that our observed values consistently fall towards the lower end of the elevation spectrum. This trend remains even when accounting for the geographical distribution of populations and the graph's structure.

Decion	Sample	Trees	Wasps	ا ماناندا م	Longitudo	Elevation
Region	Site	Sampled	Sampled	Latitude	Longitude	Range (m)
Baja California	39	7	2	23.13187	-109.75406	188
	70	23	2	23.73801	-109.81612	3089
	95	17	2	26.36298	-111.80387	263
	96	23	2	24.02943	-110.13009	750
	112	27		27.56675	-113.07373	864
	113	19		27.08696	-112.51638	644
	158	20	2	29.26645	-114.02414	899
	172	19	2	28.29053	-113.11295	697
	179	22		25.91582	-111.34837	19
	201	21	2	25.37728	-111.31542	288
	204	6		24.83127	-110.80179	384
	205	7		23.05125	-110.09175	9
Coastal Sonora	36		5	28.89523	-112.00658	192
	100		5	28.00606	-111.04223	84
	100T	6	2	27.94121	-111.08463	33
	104	6	9	27.08341	-109.68225	52
Inland Sonora	103	4	6	26.94181	-108.88808	254
	106		5	29.47434	-110.25687	423
	107		5	28.58991	-109.55771	224
	108		6	28.07264	-109.32043	224
Coastal Sinaloa	226		5	23.17888	-106.42636	101
	230		5	25.58337	-109.11327	5
	231		3	26.02424	-109.03569	98
Inland Sinaloa	228		1	23.61378	-106.33288	146
	229		2	24.16040	-106.74877	188
Zacatecas	217	6	3	21.31893	-103.14842	1271
Jalisco	215	6	2	20.75497	-103.32047	1414
	218		6	19.48764	-103.46098	1055
Morelos	219		5	18.67604	-98.77137	1368
	220		1	18.91518	-99.20843	1457
Northern Oaxaca	221		3	17.68726	-97.94190	1341
Southern Oaxaca	209		2	16.64853	-96.07933	784
	210	3		16.39097	-95.38333	201
	214	5	1	16.63315	-96.05944	805
	222		6	16.67925	-96.55636	1746

Table 1: *F. petiolaris* and *Pegoscapus* sp. sample site information and the number of trees (n = 247) and wasps (n = 102) sampled per site for phylogeographic analysis.

E notiolaria	Baja	Coastal	Inland	Central	Southern
	California	Sonora	Sonora	Mexico	Oaxaca
Baja	ΝΛ	0.2803	0.5904	0.5564	0.694
California	n/A	(0.2282 - 0.3255)	(0.5272 - 0.6455)	(0.4941 - 0.6072)	(0.6412 - 0.7344)
Coastal		ΔΙΔ	0.1622	0.1246	0.4049
Sonora		MA	(0.1254 - 0.2047)	(0.0981 - 0.1512)	(0.3582 - 0.4456)
Inland			ΝΑ	0.0703	0.4312
Sonora			1VA	(0.0405 - 0.1039)	(0.3794 - 0.482)
Central				ΔιΔ	0.3215
Mexico					(0.2725 - 0.3653)
Southern					ΝΔ
Oaxaca					N/A

Table S1: Estimates of Wright  $F_{st}$  between pairs of inferred phylogroups of F. petiolaris. Values in parentheses are bootstrapped 95% confidence intervals and significant positive  $F_{st}$  estimates are denoted in bold.

			<b>F</b> .	petiolaris	5			
Regional Sites	Sample Site	Sampled Trees	Ν	n_ests	Fis	II	hl	Hs
	39	7	7	75	0.0781	0.0257	0.2622	0.0263
	70	23	17	107	0.0449	0.0196	0.1836	0.0242
	95	17	16	119	0.0588	0.0468	0.2123	0.0279
	96	23	14	115	0.0869	0.0533	0.2209	0.0279
	112	27	20	135	0.0442	0.0300	0.1748	0.0261
Baja	113	19	15	117	0.0613	0.025	0.2329	0.0246
California	158	20	16	107	0.0549	0.0373	0.1892	0.0236
	172	19	16	119	0.0262	-0.0111	0.1253	0.0263
	179	22	19	121	0.0447	0.0469	0.2014	0.0243
	201	21	19	114	0.0436	-0.0100	0.1507	0.0254
	204	6	6	75	0.0641	0.0231	0.2636	0.0254
	205	7	7	71	0.0477	-0.0208	0.2104	0.0237
	100T	6	5	147	0.0351	-0.0020	0.1603	0.0641
Sonora	103	4	4	109	0.0341	-0.0209	0.197	0.0564
	104	6	6	148	-0.0209	-0.0987	0.0687	0.0687
Central	215	6	6	155	0.0016	-0.0581	0.0814	0.0688
Mexico	217	6	6	163	0.0663	0.0284	0.1949	0.0687
Southern	210	3	3	68	0.1177	-0.0047	0.3350	0.0448
Oaxaca	214	5	5	74	-0.0691	-0.1883	0.0950	0.0393

Table S2: Estimates of inbreeding coefficient  $(F_{IS})$  of *F. petiolaris* regional sites. Columns indicate the Regional sites, Sampling site, the number of sampled *F. petiolaris* individuals per site (Sampled Trees), the number of *F. petiolaris* samples after filtering (N), the number of loci used to calculate  $F_{IS}$  (n\_ests), The point estimate of  $F_{IS}$ , the lower confidence limit of  $F_{IS}$  estimate (ll), the upper confidence limit of  $F_{IS}$  estimate (ul), and expected heterozygosity (HS).

	F. petiolaris PHRAPL Results									
Rank	AIC	dAIC	wAIC	Parameters	Migration	Migration	Coalesce			
					$(m_{1\rightarrow 2})$	$(m_{2\to 1})$	$(t_{1.2})$			
1	15576.996	0	0.568	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	4.64	4.64	7.690			
2	15577.738	0.742	0.392	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	4.64	4.64	0.300			
3	15582.284	5.288	0.040	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	4.64	4.64	7.720			
4	15597.388	20.392	$2.12E{-5}$	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	4.64	4.64	2.131			
5	15599.090	22.094	9.05 E - 6	$m_{1.2}, t_{1.2}$	4.64	4.64	0.300			
6	15604.526	27.530	$5.97 E{-7}$	$m_{1.2}$	4.64	4.64	NA			
7	15604.567	27.571	$5.85 E{-7}$	$m_{1 \to 2}, m_{2 \to 1}, t_{1.2}$	4.64	4.64	1.415			
8	5609.904	32.908	4.06E - 8	$m_{1.2}$ , $t_{1.2}$	4.64	4.64	0.353			
9	15613.998	37.002	5.24E - 9	$m_{1.2}, t_{1.2}$	4.64	4.64	9.611			
10	15668.925	91.929	6.20E-21	$m_{1\to 2}, m_{2\to 1}, t_{1.2}$	2.150	4.639	1.110			

Table S3: The top ten ranked *Ficus petiolaris* demographic models from PHRAPL. Columns indicate values of the model rank, Akiake Information Criterion (AIC), difference in AIC from the top-ranked model (dAIC), AIC weights (wAIC), and the vector of demographic parameters (parameter vector, with northern Mexico denoted as 1 and southern Mexico as 2) corresponding to migration from population 1 to 2  $(m_{1\rightarrow 2})$ , migration from population 2 to 1  $(m_{1\rightarrow 2})$ , single migration parameter representing asymmetric or symmetric migration  $(m_{1,2})$ , and a coalescent event of population 1 and 2  $(t_{1,2})$ . The last three columns correspond to the estimate migration parameter value for migration from population 1 to 2  $(m_{1\rightarrow 2})$ , migration parameter value for migration from population 2 to 1  $(m_{1\rightarrow 2})$ , and coalescent event between population  $(t_{1,2})$ . Migration rates are given in units of 4Nm and time parameters are given in units of 4N.

Dorocosta	Baja	Coastal	Inland	Coastal	Inland	7 acretocre	Inlien	Moroloc	Northern	Southern
r egostapus	California	Sonora	Sonora	Sinaloa	Sinaloa	Tarateras	nalise		Оахаса	Оахаса
Baja	VIV	0.0043	0.0266	0.0282	0.0159	0.0263	0.0508	0.7339	0.7408	0.7624
California	1-/N	(-0.0034 - 0.0115)	(0.0125 - 0.0427)	(0.0062 - 0.0536)	(-0.0711 - 0.1082)	(-0.0539 - 0.1098)	(0.0063 - 0.1098)	(0.6755 - 0.7813)	(0.6855 - 0.7842)	(0.7181 - 0.7997)
Coastal		111	0.0147	0.0066	0.0357	0.0544	0.0498	0.7554	0.7566	0.7718
Sonora		L/ / /	(0.002 - 0.0296)	(-0.0046 - 0.018)	(-0.0408 - 0.1113)	(-0.0455 - 0.157)	(0.015 - 0.0917)	(0.7046 - 0.7991)	(0.7071 - 0.797)	(0.7285 - 0.8083)
Inland			V 1V	0.0119	-0.0127	0.0605	0.0342	0.7443	0.7459	0.7625
Sonora			14/A/	(-1e-04 - 0.0252)	(-0.0706 - 0.046)	(-0.0364 - 0.1594)	(0.0081 - 0.0622)	(0.6894 - 0.7893)	(0.6927 - 0.7861)	(0.7179 - 0.8009)
Coastal				V/V	0.0721	0.134	0.0642	0.8094	0.8056	0.8064
Sinaloa				F/V1	(-0.0052 - 0.1486)	(0.0196 - 0.2359)	(0.0304 - 0.0979)	(0.7616 - 0.8492)	(0.7602 - 0.8438)	(0.7684 - 0.8393)
Inland					111	0.1597	-0.0132	0.8017	0.7813	0.792
Sinaloa					HAN	(0.0274 - 0.2986)	(-0.0751 - 0.0617)	(0.747 - 0.8514)	(0.728 - 0.8329)	(0.7464 - 0.8323)
Tootooo						777	0.0449	0.8092	0.7977	0.7971
Zdraferas						1-/A/	(-0.0372 - 0.1371)	(0.7509 - 0.8587)	(0.7475 - 0.8433)	(0.7509 - 0.8366)
alieco							VIV	0.7157	0.7267	0.7624
nalisco							F/V/	(0.6547 - 0.7687)	(0.6712 - 0.7764)	(0.7166 - 0.8007)
Morelos								VIV	0.0577	0.5779
									(-0.0809 - 0.1999)	(0.4671 - 0.6636)
Northern									VV	0.5582
Оахаса									1.77.7	(0.4558 - 0.6488)
Southern										D//A
Оахаса										1.787
עס יוקים	. D <sub>at</sub> :	MY JO DO + O	ν. μ+λα Γ Γ μ <sub>α</sub>		امتصنيمه ا	in to solow of L		nollinoton	ب: ممنامينامين	found about

ing inferred phy-	95% confidence	
sp. pollinator, includ	ses are bootstrappe	
amples of <i>Pegoscapus</i>	ca. Values in parenth	bold.
in pairs of regional s	and Southern Oaxa	ates are denoted in
Wright's $F_{st}$ betwee	i, Northern Oaxaca,	nt positive $F_{st}$ estim
able S4: Estimates of	groups from Morelos	itervals and significal

		Pego	osca	pus				
Regional Site	Sampling Site	Sampled Wasps	Ν	n_ests	Fis	II	hl	Hs
	39	2	2	27	0.5185	0.3111	0.8039	0.0199
	70	2	2	29	0.2069	0	0.5476	0.0184
	95	2	2	23	0.2609	-0.1155	0.5791	0.0162
Baja California	96	2	0	NA	NA	NA	NA	NA
	158	2	2	21	0.0952	-0.0801	0.5	0.0126
	172	2	2	31	0.1935	-0.1668	0.4002	0.0213
	201	2	2	34	0.4412	0.1702	0.6668	0.026
	36	5	5	77	0.495	0.5368	0.7457	0.0264
Coastal Sanara	100	5	4	62	0.5244	0.5667	0.7755	0.0248
COASTAL 2011014	104	9	9	104	0.3773	0.4084	0.6067	0.0209
	100T	2	2	21	0.4762	0.4242	0.8276	0.0133
	103	6	6	51	0.553	0.4687	0.7753	0.0169
Inland Sonora	106	5	5	62	0.7505	0.7023	0.9013	0.026
	107	5	4	57	0.5002	0.5229	0.7624	0.0249
	108	6	6	76	0.4051	0.432	0.6675	0.022
	226	5	5	55	0.4082	0.4336	0.6767	0.018
Coastal Sinaloa	230	5	5	51	0.4793	0.4487	0.7584	0.017
	231	3	3	23	0.2826	0.1723	0.613	0.0112
Juland Charles	228	1	0	NA	NA	NA	NA	NA
Inland Sinaloa	229	2	2	26	0.2692	-0.0345	0.5882	0.018
Zacatecas	217	3	3	29	0.2931	0.2222	0.6192	0.0137
	215	2	2	28	0.3571	0.1111	0.6452	0.0193
Jalisco	218	6	4	68	0.3618	0.3401	0.6481	0.029
Maralaa	219	5	0	NA	NA	NA	NA	NA
ivioreios	220	1	0	NA	NA	NA	NA	NA
Northern Oaxaca	221	3	3	38	0.5263	0.4527	0.7934	0.0205
	209	2	2	8	0.5	-0.2512	0.9476	0.0066
Southern Oaxaca	214	1	0	NA	NA	NA	NA	NA
	222	6	6	77	0.4148	0.4352	0.6813	0.0235

Table S5: Estimates of inbreeding coefficient  $(F_{IS})$  of *Pegoscapus* regional sites. Columns indicate the Regional sites, Sampling site, the number of sampled *Pegoscapus* individuals per site (Sampled Wasps), the number of *Pegoscapus* samples after filtering (N, individuals with less than 50% phased data), the number of loci used to calculate  $F_{IS}$  (n\_ests), The point estimate of  $F_{IS}$ , the lower confidence limit of  $F_{IS}$  estimate (ll), the upper confidence limit of  $F_{IS}$  estimate (ul), and expected heterozygosity (HS).

Pegoscapus PHRAPL Results								
Rank	AIC	dAIC	wAIC	Parameters	Migration	Migration	Coalesce	
					$(m_{1\to 2})$	$(m_{2\to 1})$	$(t_{1.2})$	
1	25265.173	0	0.849	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	0.46	1	4.07	
2	25269.128	3.954	0.117	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	0.220	1.000	4.069	
3	25271.699	6.526	0.032	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	1	1	2.12	
4	25288.654	23.481	6.77 E - 6	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	0.46	2.149	7.808	
5	25290.092	24.919	3.30E - 6	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	0.220	2.149	2.120	
6	25294.179	29.006	$4.27E{-7}$	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	1	1	7.809	
7	25306.558	41.385	8.76E-10	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	0.455	1.073	1.299	
8	25312.562	47.389	$4.35E{-}11$	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	0.459	2.149	2.119	
9	25314.244	49.071	1.88E-11	$m_{1 ightarrow 2}$ , $m_{2 ightarrow 1}$ , $t_{1.2}$	0.458	2.144	3.829	
10	25319.975	54.802	1.07E - 12	$m_{1 \rightarrow 2}, m_{2 \rightarrow 1}$	0.22	2.15	NA	

Table S6: The top ten ranked *Pegoscapus sp.* demographic models from PHRAPL. Columns headings are as in Table S3.