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45 Abstract

46 Aim: Two main biogeographic hypotheses have been proposed to explain the Mediterranean-Turanian

- 47 disjunct distributions exhibited by numerous steppe-dwelling organisms, namely (i) dispersal during
- 48 the Messinian salinity crisis (~5.96-5.33 Ma) followed by range fragmentation and vicariance, and (ii)
- 49 Pleistocene colonization and recent processes of population subdivision (<2 Ma). Despite the two
- 50 hypotheses postulate the role of climatic alterations and changes in landmass configuration on
- 51 determining such disjunct distributions, estimates of the timing of lineage diversification have not
- 52 been complemented so far with spatially-explicit tests providing independent evidence on the
- 53 proximate processes underlying geographical patterns of population genetic
- 54 connectivity/fragmentation.
- 55 Location: Mediterranean-Turanian region
- 56 Taxon: Saltmarsh band-winged grasshopper (*Mioscirtus wagneri*)
- 57 Methods: We integrate different sources of genetic (mtDNA and ddRADseq) and spatial information
- 58 (configuration of emerged lands and niche modelling) to evaluate competing hypotheses of lineage
- 59 diversification in the saltmarsh band-winged grasshopper, a halophile species showing a classical
- 60 Mediterranean-Turanian disjunct distribution.

Results: Phylogenomic analyses reveal the presence of two North African cryptic lineages and support that trans-Mediterranean populations of the species diverged in the Pleistocene, with evidence of post-Messinian permeability of the Strait of Gibraltar to gene flow likely associated with sea level drops during glacial periods. Accordingly, spatial patterns of genetic differentiation are best explained by a scenario of population connectivity defined by the configuration of emerged landmasses and environmentally suitable habitats during glacial periods, a time when effective population sizes of the species peaked as inferred by genomic-based demographic reconstructions.

- 68 Main conclusions: Our results support post-Messinian colonization and Pleistocene diversification as
- 69 the biogeographic scenario best explaining the trans-Mediterranean disjunct distributions of
- 70 halophilous organisms.
- 71

72 KEYWORDS: bathymetry, biogeographic scenarios, ddRADseq, disjunct distributions, Messinian,
 73 phylogenomics, PleistoceneINTRODUCTION

74

75 Inferring the processes structuring genetic variation, through their effects on genetic drift and gene 76 flow, and understanding their evolutionary consequences remain a central focus of both population 77 genetics and phylogeography (Avise, 2009; Habel et al., 2015). However, despite large overlap in their 78 theoretical bodies, these disciplines have tended to evolve independently. In part, this was due to the 79 contrasting spatiotemporal scales at which the main evolutionary processes that they deal with occur 80 and whose resolution has traditionally required the usage of different molecular markers (e.g., nuclear 81 microsatellite vs. mitochondrial sequence markers; Wang, 2010). This idiosyncrasy has in part 82 hampered our capacity to understand how the microevolutionary processes operating at population 83 level ultimately drive speciation and lineage divergence at broader geographic and temporal scales 84 (Papadopoulou & Knowles, 2017). Currently, the advent of high-throughput sequencing technologies 85 and the possibility of generating vast genomic datasets has opened the door to investigate patterns of 86 genetic variation at a wide range of evolutionary scales, blurring the boundaries of population 87 genetics, phylogeography and phylogenetics (Rissler, 2016). 88 Applications of genomic data, especially when viewed through a statistical phylogeographic

Applications of genomic data, especially when viewed through a statistical phylogeographic
 framework (Knowles, 2009), are essential to studies of the divergence histories of organisms across
 geologically dynamic and environmentally heterogeneous landscapes, such as the Mediterranean. The
 modern Mediterranean basin is the legacy of complex climate and geological dynamics that have

92 taken place since the Late Miocene (Blondel & Aronson, 1999). This includes the Messinian salinity 93 crisis (MSC, ~5.96-5.33 Ma), a geological episode during which the Mediterranean-Atlantic seaways 94 progressively closed, leading to a dramatic sea-level drawdown (~1,500 m) in the Mediterranean Sea 95 and its partial desiccation (Krijgsman, Hilgen, Raffi, Sierro, & Wilson, 1999). The resulting land bridges 96 connecting regions from southern Europe and northern Africa provided a favorable geographic setting 97 for the expansion of terrestrial biotas throughout the Mediterranean region, especially for dry-98 adapted and steppe-dwelling taxa that benefited by the increased availability of open landscapes and 99 arid environments (García-Alix et al., 2016). The reopening of the Atlantic-Mediterranean marine 100 connection (Strait of Gibraltar) and refilling of the Mediterranean basin (~5.33 Ma; García-Castellanos 101 et al., 2009) led to vicariant events in numerous organism groups that had previously expanded their 102 distributions during the MSC (Chueca, Madeira, & Gómez-Moliner, 2015). The long-term persistence 103 of populations in relict patches of optimal habitat since the Messinian has been often invoked as the 104 most plausible explanation for the current disjunct distribution of many steppe or halophilous taxa 105 with populations at the western- and easternmost portions of the Mediterranean basin (Ribera & 106 Blasco-Zumeta, 1998). However, an alternative biogeographic hypothesis proposes that the origin of 107 disjunct distributions in some trans-Mediterranean organisms reflect their ability to track existing 108 suitable habitats in more recent times (i.e., post-Messinian; Allegrucci, Trucchi, & Sbordoni, 2011). 109 According to this hypothesis, disjunctions arose by colonization events and subsequent isolation 110 processes linked to Quaternary climate oscillations (Kadereit & Yaprak, 2008). Sea-level drawdown 111 (~125 m; Litcher et al., 2010) and expansions of steppe-like environments (Kajtoch et al., 2016; 112 Kirschner et al., 2020) during Pleistocene glacial periods might have facilitated the dispersal of steppe 113 species across seaways and favorable habitats and, ultimately, led to the colonization of 114 geographically distant regions that have remained isolated since then (Ortiz, Tremetsberger, Talavera, 115 Stuessy, & García-Castaño, 2007). For example, the Strait of Gibraltar, where the distance between 116 the Iberian Peninsula and Africa shortened to a few kilometers during the Pleistocene coldest stages, 117 which might have favored the exchange of fauna and flora between the two continents (Graciá et al., 118 2013). However, the east-west vicariant distributions of several Mediterranean steppe-dwelling 119 organisms have often been attributed to pre-Quaternary migration events linked to the MSC without 120 proper evaluation of alternative phylogeographic hypotheses (Ortego, Bonal, Cordero, & Aparicio, 121 2009; Ribera & Blasco-Zumeta, 1998). Clearly important insights can be gained from integrating 122 precise information on the timing of population- and lineage-level divergence, detailed 123 reconstructions of the demographic fate of the populations, and formal tests of alternative spatially-

explicit scenarios of gene flow representing expectations of population connectivity under contrastingbiogeographical hypotheses (Papadopoulou & Knowles, 2015).

126 Here, we apply an integrative and hypothesis testing framework to distinguish between 127 competing hypotheses about the Mediterranean-Turanian disjunct distribution of the saltmarsh band-128 winged grasshopper, Mioscirtus wagneri (Eversmann, 1859) (Orthoptera: Acrididae). This species is 129 the only representative of the monotypic genus *Mioscirtus* Saussure, 1888 (Cigliano, Braun, Eades, & 130 Otte, 2020) and its vast range spans from the Iberian Peninsula to central Asia (Cordero, Llorente, & 131 Aparicio, 2007). In the Mediterranean region it forms highly fragmented populations in Iberia, 132 northwestern Africa (Morocco, Algeria and Tunisia) and the Middle East (Turkey, Israel, Palestine and 133 Jordan), being absent from France, Balkan and Italian Peninsulas, northeastern Africa (Egypt and 134 Libya), and all islands with the exception of Cyprus (Cigliano et al., 2020; Cordero et al., 2007; Figure 135 1). The saltmarsh band-winged grasshopper is a specialist species intimately associated to certain 136 halophilic plants (Suaeda sp.) from the family Amaranthaceae on which it depends for feeding and 137 shelter (Cordero et al., 2007). This restricted ecological requirement limits the distribution of the 138 species to saline lowlands, including coastal marshes and inland endorheic lagoons and steppes. Three 139 subspecies of the saltmarsh band-winged grasshopper have been described so far based on subtle 140 phenotypic variations and differences in body size (Cigliano et al., 2020): M. w. wagneri (Eversmann, 141 1859) (type locality: Asia-Temperate, Middle Asia, Turkmenistan), *M. w. rogenhoferi* (Saussure, 1888) 142 (type locality: Asia-Temperate, Western Asia, Iraq), and M. w. maghrebi Fernandes, 1968 (type 143 locality: Africa, Northern Africa, Algeria). It has been hypothesized that the diversification of the 144 saltmarsh band-winged grasshopper in the Mediterranean region could be linked to the MSC. This 145 would suggest that its present-day disjunct distribution may constitute the relicts of a wider range 146 during the Late Tertiary that has since experienced gradual fragmentation (Ortego et al., 2009). 147 However, the tempo and mode of divergence between the different populations and putative 148 subspecies remain unresolved due to the lack of a comprehensive approach covering the entire 149 Mediterranean distribution range of the species and formal testing of alternative biogeographical 150 hypotheses.

151 In this study we focus on the trans-Mediterranean distribution range of the saltmarsh band-152 winged grasshopper, including populations separated >4,000 km that encompass the Mediterranean 153 and Irano-Turanian regions and comprise the three putative subspecies (Cigliano et al., 2020) (Figure 154 1). Specifically, we integrate different sources of genetic (mtDNA and ddRADSeq) and spatial 155 information (configuration of emerged lands and niche modelling) to evaluate competing 156 phylogeographic hypotheses and determine whether lineage diversification is a consequence of

157 dispersal during the MSC followed by range fragmentation and vicariance (~5.33 Ma) (Ribera & 158 Blasco-Zumeta, 1998) or, alternatively, resulted from post-Messinian colonization and Pleistocene 159 population subdivision (<2 Ma) (Graciá et al., 2013). First, we used coalescent-based methods to 160 reconstruct the phylogenetic relationships among populations, estimate the timing of lineage 161 divergence, and elucidate whether these are congruent with either Messinian or Pleistocene genetic 162 fragmentation. Second, we integrated bathymetric information and environmental niche modelling 163 (ENM) to determine the spatial configuration of emerged landmasses and climatically suitable areas at 164 different time periods (Messinian, Pleistocene and present day) and generate alternative scenarios of 165 population connectivity, the fit of which to our genetic data we evaluate based on the timing of 166 population divergence estimated in the previous step. Finally, we examined the geographical 167 distribution of genetic variation and reconstructed changes in effective population size through time 168 in order to infer the demographic responses of the different populations to past climate oscillations, 169 to identify pulses of population expansion/contraction and to link them with the historical processes 170 underlying genetic fragmentation and lineage formation.

171

172 MATERIALS AND METHODS

173

174 Sample collection

- 175 We collected 180 individuals from 20 populations of the saltmarsh band-winged grasshopper
- 176 distributed throughout its Mediterranean distribution range, including populations from Spain,

177 Portugal, Morocco, Tunisia, Turkey and Jordan (Figure 1; Table S1).

178

179 Genetic data

180 We extracted DNA and amplified and sequenced fragments of the 12S and 16S rRNA mitochondrial 181 genes as detailed in González-Serna, Ortego & Cordero (2018). After sequence alignment, trimming 182 and editing, gene fragments had 395 bp for 12S and 460 bp for 16S. From the 180 specimens, we 183 selected 7-9 individuals per population (144 samples in total; Table S1) to be processed into three 184 genomic libraries following the double-digestion restriction-fragment-based procedure (ddRADseq) 185 described in Peterson, Weber, Jay, Fisher & Hoekstra (2012), with minor modifications detailed in 186 Noguerales, Cordero & Ortego (2018). Each library was sequenced in a single-read 101-bp lane on an 187 Illumina HiSeq2500 platform at The Centre for Applied Genomics (Toronto, Canada). Raw sequences

- 188 were demultiplexed and preprocessed using STACKS 1.35 (Catchen, Hohenlohe, Bassham, Amores, &
- 189 Cresko, 2013) and assembled using PYRAD 3.0.66 (Eaton, 2014). Appendix S1 provides all details on
- 190 genomic data assembling and filtering.
- 191

192 Phylogenetic analyses: mtDNA data

193 We used mitochondrial DNA data and BEAST 1.8.0 (Drummond, Suchard, Xie, & Rambaut, 2012) to 194 reconstruct the phylogenetic relationships among the different populations and estimate the timing of 195 lineage divergence. We applied a HKY+I model of sequence evolution for both gene fragments as 196 determined in JMODELTEST2 (Darriba, Taboada, Doallo, & Posada, 2012). We assumed a normal 197 distributed substitution rate of 0.049 (SD = 0.0008) per site per million years for the 16S rRNA gene 198 fragment (Papadopoulou, Anastasiou, & Vogler, 2010) and applied a continuous-time Markov chain 199 (CTMCs) model for the 12S rRNA, as no explicit priors for the substitution rate for this gene fragment 200 are available (e.g., González-Serna et al., 2018). A strict clock and a constant demographic model was 201 used for phylogenetic reconstructions as determined by model testing using a generalized stepping-202 stone sampling approach (Baele, Lemey, & Suchard, 2016). We ran the analysis with two independent 203 MCMC chains of 100 million generations each, sampling every 10,000 generations, and discarding the 204 first 10% as burn-in.

205

206 Phylogenetic analyses: Genome-wide nuclear data

207 First, we reconstructed the phylogenetic relationships among populations using genome-wide SNP 208 data and the coalescent-based method implemented in SVDQUARTETS (Chifman & Kubatko, 2014). We 209 ran svDQUARTETS exhaustively evaluating all possible quartets and performing nonparametric 210 bootstrapping with 100 replicates for quantifying uncertainty in relationships. Second, we used BPP 4.0 211 to estimate the timing of lineage and population divergence (Flouri, Jiao, Rannala, & Yang, 2018). The 212 phylogenetic tree inferred using SVDQUARTETS was fit as the fixed topology in BPP analyses (option A00). 213 The .loci file from PYRAD was edited, converted into a BPP input file and filtered using custom R scripts 214 (J-P. Huang, https://github.com/airbugs/; Huang, Hill, Ortego, & Knowles, 2020). Due to high 215 computational burden, branch length estimation was inferred using five datasets consisting of 100, 216 200, 300, 400 and 500 randomly chosen loci (out of the 1,518 variable loci recovered) to confirm the 217 consistency of the results (e.g., Huang et al., 2020). We applied an automatic adjustment of fine-tune 218 parameters, allowing swapping rates to range between 0.30 and 0.70, and set the diploid option to

indicate that the input sequences are unphased (for further details on the diploid option in BPP, see
Flouri et al., 2018). Analyses were run for 500,000 generations, sampling every 10 generations, after a
conservative burn in of 500,000 generations.

222

223 Population genetic structure

224 We inferred population genetic structure using genome-wide SNP data and the model-based 225 clustering algorithm implemented in FASTSTRUCTURE 1.0 (Raj, Stephens, & Pritchard, 2014). We 226 performed 10 independent runs for each of the different possible K genetic clusters (from K = 1 to K = 1227 10) using the simple prior and a convergence criterion of 1×10^{-7} . We assessed the number of genetic 228 clusters that best describes our data by estimating the metrics $K_{a}^{*,c}$, the value of K that maximizes log-229 marginal likelihood lower bound (LLBO) of the data, and K_{ϵ}^{*} , the smallest number of model 230 components explaining at least 99% of cumulative ancestry contribution in our sample (Raj et al., 231 2014).

232

233 Landscape genetic analyses

234 We implemented a spatially-explicit approach based on circuit theory to model gene flow under 235 alternative hypothetical scenarios of contemporary and historical population connectivity (McRae, 236 2006). Specifically, we used CIRCUITSCAPE 4.0.5 (McRae & Beier, 2007) to calculate resistance distance 237 matrices between all pairs of populations based on isolation by resistance (IBR) scenarios defined 238 according to (i) the spatial configuration of emerged lands during the present-day, last glacial 239 maximum (LGM), and Messinian (IBR_{GEO}; see Appendix S2 for details) and (ii) the spatial configuration 240 of climatically suitable habitats as inferred from projections of an environmental niche model (ENM) 241 to present-day and LGM bioclimatic conditions (IBR_{CLI}; see Appendix S3 for details). In order to identify 242 the resistance value for sea water that best explains observed estimates of genetic differentiation, we 243 explored a range of resistance values (10-1,000,000) for this landscape feature plus a scenario that 244 considered it as an impassable barrier to dispersal. For IBR_{GEO} scenarios, pixels coded as "emerged 245 land" were assigned a resistance value of 1. For IBR_{CLI} scenarios, logistic environmental suitability 246 scores (x) yielded by projections of the ENM were transformed as 1-x, so that smaller pixel values offer 247 lower resistance to gene flow. Additionally, we tested a null model of isolation-by-distance (IBD_{NULL}) by 248 generating a completely "flat landscape" scenario based on a map in which all pixels (including sea 249 water and emerged lands) have a fixed resistance (=1) value. We used multiple matrix regressions with

- randomization (MMRR; Wang, 2013) to test all resistance matrices against matrices of population genetic differentiation (F_{ST}) calculated in ARLEQUIN 3.5 (Excoffier & Lischer, 2010) for mtDNA and genome-wide SNP data.
- 253

254



We analyzed spatial clines of genetic diversity (π , nucleotide diversity) estimated for both mtDNA and genome-wide nuclear data using DNASP 5.10/6.12 (Rozas et al., 2017). We tested the association between genetic diversity (π) and geography (latitude and longitude) using generalized linear models (GLMs) as implemented in the R package 'Im4'. GLMs were constructed using a Gaussian error distribution, an identity link function and a weighted least square (WLS) method, where weight equals the sample size for each population (Bates, Maechler, Bolker, & Walker, 2015).

261

262 Demographic inference

- 263 We inferred the demographic history of each population using STAIRWAY PLOT 2.0, which implements a
- 264 flexible multi-epoch demographic model to estimate changes in effective population size (N_e) over
- time using the site frequency spectrum (SFS) (Liu & Fu, 2015). We used the script *easySFS.py* (I.
- 266 Overcast, <u>https://github.com/isaacovercast/easySFS</u>) to calculate a folded SFS for each population. To
- avoid the effects of linkage disequilibrium and remove all missing data for the calculation of the SFS,
- 268 we considered a single SNP per locus and retained only loci present in ~66% of individuals. Populations
- with <7 individuals (PHU and MER) were excluded from STAIRWAY PLOT analyses. Demographic
- 270 reconstructions in STAIRWAY PLOT were run fitting a mutation rate of 2.8×10^{-9} per site per generation
- 271 (Keightley, Ness, Halligan, & Haddrill, 2014), considering a one-year generation time (Cordero et al.,
- 272 2007), and performing 200 bootstrap replicates to estimate 95% confidence intervals.
- 273
 - 5
- 274 RESULTS
- 275

276 Genetic data: mtDNA data

- We found 16 and 21 unique haplotypes for 12S and 16S gene fragments, respectively (Table S1). In
- 278 particular, 11 and 14 haplotypes for 12S and 16S gene fragments, respectively, were exclusively found
- in a single population (Table S1). The remaining haplotypes were shared among individuals belonging

- to nearby populations located within the same geographic region (Table S1). Remarkably, individuals
 from Iberian (TIN) and African (HOC) populations located at both sides of the Strait of Gibraltar shared
- the same 12S and 16S haplotypes (Table S1).
- 283

284 Genetic data: Genome-wide nuclear data

Four individuals (one individual from PHU and three individuals from MER) recovered a very low number of reads (<10,000) and were discarded from subsequent analyses. After filtering and assembly steps in PYRAD, each remaining individual (*n* = 140) retained on average 1.83 million sequence reads (SD = 0.66 M), which represents 92.50% (SD = 2.29%) of their initial number of reads (Figure S1). On average, we recovered 49,415 loci (SD = 9,305; range = 23,170-79,257) per individual.

290

291 Phylogenetic analyses

- 292 Phylogenetic inference based on both mitochondrial (BEAST) and genome-wide nuclear (SVDQUARTETS) 293 data consistently supported two highly divergent lineages only represented in the populations located 294 in western Morocco (ZIM) and southern Tunisia (KEB) (Figure 2). The remaining populations were 295 grouped into four clades corresponding to the Middle East, Africa along with southwestern Iberia, 296 northeastern Iberia, and central-southeastern Iberia populations (Figure 2). All analyses showed 297 consistently that Iberian populations were not reciprocally monophyletic, as populations from 298 southwestern Iberia (CAS and TIN) were placed within the northern Africa clade. We found a 299 discordance between mitochondrial and nuclear genealogies in the phylogenetic position of the 300 northeastern Iberian clade (Ebro Valley): mtDNA data supported populations from northeastern Iberia 301 as the most external lineage of the trans-Mediterranean clade, whereas nuclear data placed them as a 302 sister lineage to the central-southeastern Iberian clade (Figure 2).
- 303

304 Divergence time estimation

Estimates of divergence time based on BEAST analyses for mtDNA revealed that the ZIM+KEB clade and
 the remaining lineages diverged from a shared common ancestor during the Pliocene (~4.5 Ma;
 Figure 2). According to BPP analyses based on genome-wide nuclear data, the most ancient split was

- 308 estimated around 0.01032 τ units (Figure 3). Assuming a genomic mutation rate of 2.8 × 10⁻⁹ per site
- 309 per generation (Keightley et al., 2014) and a one-year generation time (Cordero et al., 2007), the

- 310 oldest diversification event may have taken place during the Early Pleistocene, around 1.84 Ma (τ =
- 311 $2\mu t$, being μ the mutation rate and t the divergence time). The subsequent split between ZIM and KEB
- 312 populations was dated to occur during the Early Pleistocene (mtDNA data: ~1.7 Ma; genome-wide
- 313 nuclear data: \sim 1.08 Ma, 0.00608 τ units). The diversification of the remaining clades triggered around
- 314 the Middle Pleistocene (mtDNA data: \sim 1.0 Ma; genome-wide nuclear data: \sim 0.21 Ma, 0.00117 τ
- 315 units) and finished with the split of northern Morocco and southwestern Iberia populations in the Late
- 316 Pleistocene (mtDNA data: ~100 ka; genome-wide nuclear data: ~47.5 ka, 0.00026 τ units) (Figure 3).
- 317 Inferences from BPP were consistent across analyses based on datasets considering a different number
- of loci (Figure S2).
- 319

320 Population genetic structure

321 Results from FASTSTRUCTURE analyses indicated K = 6 as the most likely number of genetic clusters $(K_{\phi}^{*c})^{c}$ 322 = 6; $K_{\varepsilon}^{*} = 6$). The inferred genetic groups were consistent with phylogenetic inferences (Figure 2b). The 323 highly divergent lineages from ZIM and KEB populations of North Africa were included in two distinct 324 genetic clusters (Figure 2b). The other four genetic clusters corresponded to the rest of North African 325 populations plus southwestern Iberian populations, and populations from northeastern Iberia, central-326 southeastern Iberia, and the Middle East (Figure 2b).

327

328 Landscape genetic analyses

329 Pairwise F_{st} matrices calculated for mitochondrial and genome-wide nuclear data (Table S2) were significantly correlated (Mantel test, r = 0.581, p < 0.001). Resistance distance matrices estimated 330 331 under most scenarios were positively correlated with population genetic differentiation (F_{sT}) at both 332 nuclear and mitochondrial levels (Table S3). However, multiple matrix regression with randomization 333 (MMRR) analyses consistently supported that the scenario of population connectivity based on the 334 spatial configuration of emerged lands and climatically suitable areas during the LGM (IBR_{CL-LGM}) was 335 the best fit to our data and the only one retained into the final model (Table 1). This result was 336 consistent across both mitochondrial and genome-wide nuclear data and when including and excluding from the analyses the highly divergent ZIM and KEB populations (Table 1). 337

338

339 Genetic diversity

340 Mitochondrial and genome-wide nuclear genetic diversity were not significantly correlated (Pearson's 341 r = -0.127, p = 0.592). The highest levels of mitochondrial genetic diversity were found in ZIM and KEB 342 populations followed by the three populations from northeastern Iberia, whereas several populations 343 from central, southeastern and southwestern Iberia, northern Morocco and Jordan exhibited no 344 mitochondrial genetic variation (π = 0; Table S1). Accordingly, we did not find any significant 345 relationship between mitochondrial genetic diversity and longitude and latitude either analyzing all 346 populations (longitude, t = -0.360, p = 0.723; latitude, t = 0.030, p = 0.977) or excluding ZIM and KEB 347 populations (longitude, t = 0.179, p = 0.860; latitude, t = 1.919, p = 0.074). With regards to genome-348 wide nuclear data, populations from the Middle East (TUZ and DEA) showed the highest levels of 349 genetic diversity, whereas populations from central and northeastern Iberia presented the lowest 350 estimates (Figure 1; Table S1). Nuclear genetic diversity was positively correlated with longitude (β = 1.19×10^{-4} , t = 8.408, p < 0.001) and negatively correlated with latitude (β = -4.11 × 10⁻⁴, t = -7.407, p 351 352 <0.001). Analyses excluding the highly divergent populations ZIM and KEB yielded analogous results 353 (longitude: $\beta = 1.16 \times 10^{-4}$, t = 8.981, p < 0.001; latitude: $\beta = -4.74 \times 10^{-4}$, t = -8.485, p < 0.001). 354

554

355 Demographic inference

356 STAIRWAY PLOT analyses revealed strong declines in N_e for all populations after the end of the last glacial 357 period (~10-20 ka; Figure 4). Phylogenetically closer populations shared more similar demographic 358 histories in terms of the trends and magnitude of population size changes. In line with analyses of 359 nuclear genetic diversity, populations from central, southeastern and northeastern Iberian Peninsula 360 exhibited on average the lowest historical estimates of N_e , Middle East populations presented the 361 highest estimates, and populations from northern Africa and southwestern Iberia had intermediate 362 values (Figure 4).

- 363
- 364 DISCUSSION
- 365

366 Our phylogenetic analyses and the estimated timing of lineage splits, complemented with evidence367 provided by spatially-explicit testing of alternative scenarios of gene flow, support a post-Messinian

368 colonization and Pleistocene genetic fragmentation as the most plausible biogeographic hypothesis to

- 369 explain the trans-Mediterranean disjunct distribution of the saltmarsh band-winged grasshopper.
- 370 Intriguingly, our analyses revealed that the species exhibits a complex phylogeographic structure

371 (Figure 2), with three main cladogenetic events and a deep genetic split between parapatric northern

- 372 Africa lineages that indicate cryptic processes of allopatric divergence and call upon a taxonomic
- revision of this monotypic genus (Cigliano et al., 2020). Shared genetic lineages between
- 374 southwestern Iberia and the Maghreb region support the recurrent permeability to gene flow of the
- 375 Strait of Gibraltar during the Pleistocene and point to the important role of sea-level fluctuations
- 376 linked to glacial cycles in promoting biotic exchanges between Europe and Africa.
- 377

378 Ancient cryptic lineages in the Maghreb region

379 One of the major questions in biogeography research on the western Palearctic realm is elucidating 380 the processes underlying the disjunct distributions of many Mediterranean organisms (Sanmartín, 381 2003). We found that genetic variation in the saltmarsh band-winged grasshopper is organized into 382 three main evolutionary lineages whose divergence most likely took place after the end of the 383 Messinian age (Figures 2-3). The first two sister lineages are distributed in western Morocco (ZIM) and 384 southern Tunisia (KEB), regions separated >1,600 km apart from each other. These lineages present a 385 parapatric distribution with respect to their geographically closest populations that belong to the third 386 and more widespread trans-Mediterranean clade. This intriguing phylogeographic structure contrasts 387 with the prevailing biogeographic pattern in the Maghreb, where the foremost genetic discontinuities 388 have been linked to the Moulouya River (northern Morocco) and the Kabylia region (northern Algeria) 389 in numerous terrestrial organisms (e.g., Beddek et al., 2018). The genetic discontinuities and disjunct 390 distributions of these lineages likely represent processes of allopatric isolation driven by inland 391 landscape changes occurring along the Pliocene and Pleistocene throughout the Maghreb region. 392 Since the Early Pliocene, the Mediterranean region experienced a progressive aridification-cooling 393 climatic shift that led to the expansion of steppe formations (Jiménez-Moreno, Fauquette, & Suc, 394 2010). This shift to a drier climate regime was particularly notable in North Africa (Griffin, 2002), which 395 could have facilitated the establishment and expansion of halophilous vegetation along vast endorheic 396 lacustrine areas (i.e., sabkhas and chotts) located across the Maghreb and whose development dates 397 back to the Late Messinian associated to local tectonic dynamics (Capella et al., 2018). These 398 geomorphological and climate alterations could have synergistically contributed to create refugial 399 areas for the species in northern Africa, leading to long-term isolation processes that might have 400 impeded gene flow (e.g., via the evolution of reproductive isolation or lack of secondary contact) with 401 its parapatric sister lineage widely distributed across the Mediterranean.

403 Pleistocene divergence throughout the Mediterranean region

404 Focusing on the trans-Mediterranean clade, our phylogenetic inferences support an east-to-west 405 divergence that led to the formation of three main lineages distributed in the Middle East, northern 406 Africa and Iberian Peninsula with a distribution gap in Egypt and Libya. As consistently estimated on 407 the basis of both mitochondrial and nuclear data (Figures 2-3), the split of these clades began around 408 the Middle Pleistocene (~1.0-0.21 Ma) and continued until the Late Pleistocene (<100 ka). These 409 findings are concordant with phylogeographic inferences (i.e., Quaternary westward expansion) for 410 other trans-Mediterranean halophilic and steppe-dwelling taxa (Escudero, Vargas, Arens, Ouborg, & 411 Luceño, 2010; Kadereit & Yaprak, 2008; Pérez-Collazos, Sánchez-Gómez, Jiménez, & Catalán, 2009; 412 Weising & Freitag, 2007). Our analyses also revealed that the three main lineages previously described 413 within the Iberian Peninsula on the basis of mitochondrial (Ortego et al., 2009) and nuclear 414 microsatellite (Ortego, Aguirre, & Cordero, 2010) data are not reciprocally monophyletic (Figures 2-3), 415 with populations from southwestern Iberia being nested within the northern Africa clade (see 416 Husemann, Schmitt, Zachos, Ulrich, & Habel, 2014). In turn, we found a mitonuclear discordance in 417 the phylogenetic position of the northeastern Iberian clade, which might have been caused by the 418 interplay between historical secondary contact and different evolutionary and demographic processes 419 (see Toews & Brelsford, 2012). These results point to two independent colonization events of the 420 Iberian Peninsula during the Pleistocene followed by allopatric divergence from their respective 421 African ancestors. Thus, our results strongly support that the Strait of Gibraltar was a permeable 422 barrier to dispersal, probably in association with sea-level changes linked to Quaternary climatic 423 oscillations (Graciá et al., 2013). The sea-level during Pleistocene glacial periods has been estimated to 424 be approximately 125 meters lower than at present in the western Mediterranean region (Rohling et 425 al., 2017), a fact that would have reduced the distance between European and African coasts to less 426 than 5-7 km at the Camariñal Sill area (Lujan et al., 2011). This contributed to the emergence of small 427 islands and shoals facilitating "steeping-stone" dispersal and the sporadic exchange of terrestrial faunas across the Strait of Gibraltar (Cosson et al., 2005; see also Ortiz et al., 2007). Accordingly, our 428 429 landscape genetic analyses supported that genetic differentiation among populations is best explained 430 by the configuration of shorelines and suitable habitats during Pleistocene glacial periods and when 431 sea surface is modelled to offer much more resistance to movement than emerged lands but without 432 acting as impassable barrier to gene flow (Tables 1 and S3).

433

434 Genetic diversity and past demographic history

435 Demographic reconstructions revealed that all analyzed populations have experienced population size 436 declines after the LGM coinciding with the beginning of the current interglacial stage (Figure 4), a 437 period during which climatic conditions likely resulted in the progressive shrink of suitable open 438 habitats for the species (Weising & Freitag, 2007). Palynological studies have evidenced that extensive 439 areas of the Mediterranean region were vegetated by steppe-like formations during glacial periods 440 that became progressively replaced by temperate forests during interglacial stages (Carrión et al., 441 2012; Sánchez-Goñi, Eynaud, Turon, & Shackleton, 1999). Consequently, the confinement of the 442 saltmarsh band-winged grasshopper in refuges of suitable habitat during unfavorable periods may 443 have led to processes of allopatric divergence along the Quaternary as reported for steppe-like and 444 halophytic species presenting similar environmental requirements (Kajtoch et al., 2016; Kirschner et 445 al., 2020). Demographic inferences based on genomic data contrast with the predictions of our ENM, 446 which suggests that the extent of climatically suitable habitats for the species have tended to increase 447 since the LGM (Figure 1f). This might reflect the difficulty of climate-based niche modelling to identify 448 refugial areas in highly specialized species tightly linked to scattered and small-size habitat patches 449 (i.e., salt-marshes, sabkhas and chotts) for which spatially-explicit information about their past 450 distribution and connectivity is not currently available (González-Serna, Cordero, & Ortego, 2019). The 451 very limited dispersal capacity of the saltmarsh band-winged grasshopper documented in previous 452 studies (Ortego, García-Navas, Noguerales, & Cordero, 2015) points to postglacial contraction of 453 suitable habitats and regional extinction in northeastern Africa, rather than long-distance dispersal, as 454 the most likely cause of the species' distribution gap in Egypt and Libya (Cigliano et al., 2020). 455 Accordingly, our data showed a general pattern of genetic erosion, the total lack of genetic variation 456 at mitochondrial level in many populations, and an east-to-west and south-to-north decline of nuclear 457 genetic diversity (Figure 1b; see also Escudero et al., 2010). This pattern might reflect the strong 458 fragmentation of most contemporary populations, historical declines revealed by demographic 459 reconstructions, and founder events resulted from serial colonization from the putatively ancestral 460 range located in the Middle East (Pérez-Collazos et al., 2009).

461

462 Taxonomic implications

463 Our study revealed cryptic diversity within the saltmarsh band-winged grasshopper, as well as
464 incongruence between current subspecies designations and the inferred phylogenetic relationships
465 (Cigliano et al., 2020). For example, the populations ZIM and KEB, putatively belonging to the
466 subspecies *M. w. maghrebi*, accumulated more genetic divergence than the remaining populations

467 assigned to the three subspecies (Figure 2). This finding highlights the need for evaluating the 468 taxonomic status of these two North Africa populations as potentially distinct taxa, which would 469 ideally require additional population sampling, the identification of potential contact zones with the 470 other Maghrebian lineage to determine whether the two are reproductively isolated, and 471 comprehensive morphological analyses that consider diagnostic phenotypic traits (e.g., Huang et al., 472 2020; Noguerales et al., 2018). Our phylogenetic analyses also shed light on the controversial 473 taxonomic status of Iberian populations (Cordero et al., 2007), supporting the existence of two 474 putative subspecies: M. w. maghrebi in southwestern Iberia (Fernandes, 1968) and M. w. wagneri in 475 central, southeastern and northeastern Iberia (Cordero et al., 2007). Finally, our study clarifies the 476 phylogenetic position of Turkish populations, where subspecies M. w. wagneri and M. w. rogenhoferi 477 have been indistinctly considered in the literature (Naskrecki & Ünal, 1995; Ünal, 2011). In this sense, 478 our analyses supported the close phylogenetic relationship between Turkish and Jordanian 479 populations (Figures 2), which is coherent from a geographical point of view and agrees with the 480 assignment of Middle East populations to the subspecies M. w. rogenhoferi, as reported in most 481 Orthoptera inventories from the region (Katbeh-Bader, 2001; Naskrecki & Ünal, 1995).

482

483 CONCLUSIONS

484

485 Our study demonstrates that the disjunct distribution of the most widespread lineage of the saltmarsh 486 band-winged grasshopper across the Mediterranean basin was shaped by its capacity to track and 487 colonize suitable habitats during the Pleistocene (Graciá et al., 2013), rather than a consequence of 488 fragmentation and long-term persistence of relict populations after range expansions linked to the 489 MSC (Ribera & Blasco-Zumeta, 1998). The cohesiveness of Moroccan and southern Iberia populations 490 evidences the post-Messinian permeability of the Strait of Gibraltar, while the presence of highly 491 divergent cryptic lineages in northern Africa highlights the pivotal role of the Maghreb as a 492 diversification area during the Pliocene-Pleistocene (Husemann et al., 2014). Overall, our study 493 emphasizes the power of combining phylogenetic inference with spatially-explicit testing of alternative 494 biogeographical scenarios to unravel hidden diversification patterns and gain insights into the 495 proximate processes underlying the origin of disjunct distributions.

496

497 DATA AVAILABILITY STATEMENT

- 498 Raw Illumina reads have been deposited at the NCBI Sequence Read Archive (SRA) under BioProject
- 499 PRJNA663622. Mitochondrial DNA sequences for the 12S and 16S gene fragments are deposited in
- 500 GenBank with accession numbers MW018492-MW018671 and MW018175-MW018354, respectively.
- 501 Input files for all analyses are available for download from the Dryad Digital Repository
- 502 (https://doi.org/10.5061/dryad.qfttdz0fk).
- 503
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- 684
- 685 BIOSKETCH
- 686 Víctor Noguerales is interested in understanding the ecological and evolutionary processes shaping
 687 spatial patterns of biological diversity. His research harnesses the power of genomic data to respond

- questions on how genetic variation arises and is maintained across different levels of biologicalorganization.
- 690
- 691 Author contributions: All authors conceived and designed the study and analyses. VN, PJC and JO
- 692 collected the samples. VN performed the lab work and analyzed the data guided by JO, who
- 693 performed the ENMs. VN wrote the manuscript with help of JO and inputs from PJC and LLK.

r Manus Author

TABLE 1 Multiple matrix regressions with randomization (MMRR) testing the relationship between mitochondrial (mtDNA) and nuclear (nDNA) genetic differentiation and distances calculated under alternative isolation-by-resistance (IBR) scenarios defined by the spatial configuration of emerged lands in the present (IBR_{GEO-CUR}), last glacial maximum (IBR_{GEO-LGM}) and Messinian (IBR_{GEO-MES}), and the spatial configuration of climatically suitable habitats in the present (IBR_{GEO-CUR}) and last glacial maximum (IBR_{GEO-LGM}). The IBD_{NULL} scenario refers to a layer in which all pixels had the same resistance value (=1). Each scenario initially considered a range of hypothetical resistance values offered by the sea water and only the one best fitting the data was included in final multivariate analyses presented in this table (see Table S3). Analyses were performed both including all populations and excluding the highly divergent ZIM and KEB populations.

701

		All populations							Without ZIM and KEB						
		mtDNA			nDNA				mtDNA			nDNA			
Ω	Variable	в	t	р	в	t	р		в	t	Р	в	t	р	
	Explanatory terms														
\geq	Intercept	0.058			0.077				0.052			0.096			
	IBR _{CLI-LGM}	0.531	8.714	0.001	0.410	6.662	0.001		0.552	8.057	0.001	0.356	4.977	0.001	
	Rejected term	าร													
\mathbf{O}	IBD _{NULL}	0.222	2.269	0.119	0.114	1.373	0.477		0.203	1.734	0.283	0.068	0.552	0.607	
	$IBR_{GEO-CUR}$	-0.218	-0.569	0.737	-0.065	-0.170	0.910		-0.420	-0.913	0.571	-0.709	-1.487	0.220	
+-	$IBR_{GEO-LGM}$	0.135	0.266	0.868	0.009	0.010	0.997		-0.241	-0.412	0.799	-0.245	-0.402	0.724	
	$IBR_{GEO-MES}$	0.226	2.092	0.204	0.130	1.601	0.467		0.202	1.652	0.310	0.052	0.407	0.715	
	IBR _{CLI-CUR}	-0.179	-0.565	0.769	0.081	0.167	0.927		-0.392	-1.106	0.519	0.029	0.195	0.877	

FIGURE 1 (a) Genetic cluster membership and (b) genetic diversity of the study populations of saltmarsh band-winged grasshopper (*Mioscirtus wagneri*) as inferred using genome-wide nuclear data.
Background maps display elevation and sea depth information. Panels c-d show the approximate spatial configuration of emerged lands during the maximum sea-level drop estimated for (c) the last glacial maximum (LGM; -125 m) and (d) the Messinian (-1,500 m). Panels e-f show the climate-based habitat suitability during the (e) present and (f) the LGM (~21 ka) as inferred by environmental niche modelling.
Yellow dots indicate the location of sampling sites (panels c and d). Population codes as in Table S1.



713 FIGURE 2 (a, c) Phylogenetic relationships and (b) genetic structure of populations of the saltmarsh 714 band-winged grasshopper (*Mioscirtus wagneri*). Panel (a) shows the maximum clade credibility tree for 715 mitochondrial data (16S and 12S gene fragments) as inferred in BEAST. Estimates of divergence time for 716 the main clades (median and lower and upper 95% highest posterior density, in brackets) and branch 717 support values (*=1.0) are indicated. Panel (b) shows population genetic structure for the most likely 718 number of clusters (K = 6) as inferred by FASTSTRUCTURE using genome-wide SNP data. Panel (c) shows 719 the species-tree inferred by SVDQUARTETS using genome-wide SNP data. Bootstrapping support values 720 are indicated on the nodes (*=100%). Putative membership of the populations to the different subspecies and population codes as in Table S1. 721

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FIGURE 3 Divergence times estimated using BPP with a subset of 500 randomly chosen loci. The topology
was fixed using the phylogenetic tree inferred using SVDQUARTETS. Bars on nodes indicate the 95% highest
posterior densities (HPD) of the estimated divergence times. Background colors represent geological
divisions as Early Pleistocene (EP, ~2.58-0.77 Ma), Middle Pleistocene (MP, ~0.77 Ma - 126 ka), Late
Pleistocene (LP, ~126-11.7 ka) and Holocene (H, ~11.7 ka to present). Population codes as in Table S1.



733 FIGURE 4 Demographic history of the studied populations inferred using STAIRWAY PLOT. Lines represent 734 the median estimate of the effective population size (N_e). Axes are logarithmically scaled and 735 populations are grouped into the different panels according to the results of phylogenetic and clustering 736 analyses. Upper panels represent temperature anomaly ($\Delta T \ ^{\circ}C$) in the Late Quaternary as estimated 737 from the EPICA (European Project for Ice Coring in Antarctica) Dome C ice core (Jouzel et al., 2007). 738 Highlighted periods show the extent of the last glacial period (~110-11.7 ka), last glacial maximum 739 (LGM: \sim 19-21 ka), and Holocene (\sim 11.7 ka to present). Population codes as in Table S1. 740



Author





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