INTRODUCTION

Natural populations change in size and range in response to biotic and abiotic factors over both ecological and evolutionary timescales. Given that sequence variation is the only source of information about the evolutionary past of many taxa, there is considerable interest in using genetic data to reconstruct the demographic history of populations (see Beichman et al., 2018, for a recent review). A key question is to what extent past demographic changes have been shaped by climatic events and whether such responses are concordant across co-distributed taxa, or vary with species traits (Hickerson et al., 2010; Papadopoulou & Knowles, 2016; Stone et al., 2012).

A major aim of phylogeographic studies has been to understand the origins and determinants of spatial structure in natural populations (Avise et al., 1987; Hickerson et al., 2010). Systematic comparisons of demographic histories across co-distributed taxa offer the chance to identify shared abiotic triggers of demographic events (Bermingham & Avise, 1986; Hewitt, 2000; Hickerson et al., 2010). If divergence and admixture between populations are largely the outcome of shared climatic and geological history, one would expect co-distributed and ecologically similar species to show concordant
For example, many temperate plant and animal taxa (including humans) have colonized the Western Palaearctic in a series of westwards range expansions. Overlaid onto this older longitudinal colonization history is a cyclical history of latitudinal range changes (Hewitt, 2000; Hofreiter & Stewart, 2009). Starting with the onset of major glacial cycles in the mid Pleistocene, the ranges of temperate taxa in Europe were restricted to southern refugia during glacial maxima, primarily in Iberia, Italy and the Balkans (Feliner, 2011; Hewitt, 2000; Hofreiter & Stewart, 2009) and expanded northwards during interglacial periods. As a consequence, modern refugial populations commonly harbour higher genetic diversity than northern populations founded by postglacial range expansion (e.g. Hewitt, 2000; Rokas et al., 2003; Tison et al., 2014; Vitales et al., 2016) (but see Comps et al., 2001). While the structuring of genetic diversity by refugia is ubiquitous (Hewitt, 1996), recent comparative studies of the population and speciation history of European taxa have, perhaps unsurprisingly, shown no support for temporal concordance (Bunnefeld et al., 2018; Ebdon et al., 2021).

Bunnefeld et al. (2018) used whole-genome sequence data to compare the timing of divergence and admixture events between three southern refugial populations across 13 co-distributed species in a tritrophic Western Palaearctic community comprising oak (Quercus) host plants, cynipid gall wasp herbivores and chalcid parasitoid natural enemies (Stone et al., 2002). The latter are obligate specialists of cynipid galls, allowing the guild of parasitoids to be considered in ecological isolation. Many of the component species show broad longitudinal distributions, extending from Iberia in the West to Iran in the East, and all species studied to date are genetically structured into major southern refugial populations (Lohse et al., 2010, 2012; Nicholls et al., 2010, 2012; Petit et al., 2002; Stone et al., 2012, 2017). Bunnefeld et al. (2018) showed that seven out of the 13 community members studied colonized refugia in the Balkans and Iberia through westwards range expansion from an eastern origin one or more glacial cycles in the past. However, four species had a discordant, Western origin and two showed no clear signal of a longitudinal history. Bunnefeld et al. (2018) also found little evidence for temporal concordance: although divergence and admixture between refugial populations are broadly concentrated in the late Pleistocene, the timing of divergence and admixture between refugia is to a large extent idiosyncratic across species. However, in order to explore a plausible space of necessarily complex demographic scenarios which capture aspects of both the longitudinal colonization history (divergence between refugia) and the more recent history of latitudinal range shifts (admixture between refugia), Bunnefeld et al. (2018) had to make drastic simplifying assumptions about the second major axis of demographic history: population size change. Specifically, their analyses assumed a single fixed effective population size per refugium. This immediately raises the question of whether the lack of concordance of species’ histories found by Bunnefeld et al. (2018) is unique to the broad spatial and deep temporal scale captured by their models, or a more general property of the history of ecological communities which holds over a range of spatio-temporal scales. There are several reasons to expect the degree of temporal concordance to depend on spatial and temporal scales (Papadopoulou & Knowles, 2016). In particular, colonization and admixture histories may be more temporally discordant than population size changes, simply because the former depend on long-distance dispersal events which are both a function of a species’ dispersal ability and inherently random (Smith et al., 2014).

Here, we use whole-genome sequence (WGS) data to quantify and compare the histories of population size change within a single glacial refugium across seven chalcid parasitoid species in the oak gall wasp community previously studied by Bunnefeld et al. (2018). We focus on the Iberian refugium, which for most species in the oak gall wasp parasitoid community—including four out of the seven parasitoids studied here (Table 1)—represents the end point of a longitudinal expansion history: Iberian populations have lower genetic diversity than more eastern refugia and show little evidence of population structure (Nicholls et al., 2010; Rokas et al., 2003; Stone et al., 2018).

**TABLE 1** Maximum-composite likelihood estimates (MCLE) of parameters under the single-step-change model for Iberian populations of seven species of chalcid parasitoid wasps under a model of a single-step change. Estimates for the time of \( N_e \) change (\( T \)) are given in thousands of years ago (kya). 95% CIs are shown in brackets. The maximum \( N_e \) is scaled relative to \( \kappa \) as \( \text{Max}[N_e]/\kappa \). The second column gives the origin of the longitudinal expansion history as inferred by Bunnefeld et al. (2018).

| Species          | Origin  | \( \kappa \) | Current \( N_e \times 10^3 \) | Ancestral \( N_e \times 10^3 \) | \( T \) (kya) | Relative Max \( [N_e] \) |
|------------------|---------|-------------|-------------------------------|-------------------------------|-------------|------------------------|
| Torymus auratus  | n/a     | 0.00509     | 1,142 (1,126–1,157)           | 422 (418–425)                | 100 (94–106)| 2.14                   |
| Megastigmus stigmatizans | West    | 0.00054     | 18 (14–21)                   | 61 (60–61)                    | 10 (9.5–10.4)| 1.55                   |
| Megastigmus dorsalis | East   | 0.00142     | 67 (59–75)                   | 144 (143–145)                | 12.5 (12.0–13.0)| 1.40                   |
| Ormyrus nitidulus | East    | 0.00126     | 47 (45–49)                   | 194 (192–197)                | 33 (31–35)   | 2.13                   |
| Ormyrus pomaceus  | West    | 0.00230     | 207 (207–208)                | n/a                          | n/a         | n/a                    |
| Eurytoma brunniventris | East    | 0.00569     | 523 (521–525)                | n/a                          | n/a         | n/a                    |
| Cecidostiba fungosa | East   | 0.00258     | 234 (233–235)                | n/a                          | n/a         | n/a                    |
2007) (for an exception, see Rokas et al. (2001)). Crucially, Bunnefeld et al. (2018) show that Iberian populations are not affected by post-colonization gene flow from other refugia. This allows for a meaningful comparison of Iberian population size histories in isolation from other refugia. Our sampling targets males (five from Iberia, one from the Balkans in each species), whose haploid genome facilitates analysis (Bunnefeld et al., 2018; Hearn et al., 2014). WGS data allow for powerful demographic inference even for non-model organisms. However, given fragmented reference genomes and limited sample sizes available for these species, inference methods must be chosen carefully and, ideally, should use linkage as well as allele frequency information. In particular, inference based only on the site frequency spectrum (SFS) (e.g. Gutenkunst et al., 2009) requires larger samples for accurate inference of recent population history.

We investigate the history of population size change in Iberia using two approaches that use the signal contained in genome-wide variation of a small sample of individuals in different ways (Bunnefeld et al., 2018; Li & Durbin, 2011; Lohse et al., 2011). We use a parametric maximum-composite likelihood (MCL) method based on a blockwise summary of sequence variation (Lohse et al., 2011) to fit a model of a single instantaneous step change in \( N_e \) (hereafter referred to as the ‘single-step-change analyses’). Using analytic likelihood calculations to fit such fully specified but minimally complex histories utilizes both allele frequency and linkage information and facilitates statistical comparison between species for the timing and magnitude of \( N_e \) change. We also applied a non-parametric method commonly used to visualize population size change, the pairwise sequentially Markovian coalescent (PSMC) (Li & Durbin, 2011), which is based on the distribution of pairwise differences in minimal samples of two haploid genomes. PSMC generates a more resolved picture of \( N_e \) change through time, including a qualitative assessment of the time at which populations have diverged. As the two methods make opposing assumptions, we will not attempt any formal comparison between them but rather compare inferences qualitatively. Both methods use different data properties and sampling strategies and, as a consequence, have contrasting limitations: while the single-step-change analyses—by design—cannot detect gradual \( N_e \) changes or resolve population histories involving multiple changes, PSMC is known to smooth out very sudden changes. PSMC is based on fewer samples which contain less information (i.e. fewer coalescence events) about \( N_e \) in the recent past (Li & Durbin, 2011). The two methods therefore complement each other and together provide a comprehensive picture of population history.

We address the following questions: (i) To what extent are the direction and timing of changes in \( N_e \) concordant across members of the oak gall parasitoid guild? Do species show evidence for simultaneous \( N_e \) change, suggesting concordant responses to a shared underlying driver, or are their demographic histories largely idiosyncratic as shown by Bunnefeld et al. (2018) for the timing of divergence and admixture between refugia? (ii) Do sudden changes in population size coincide with specific glacial or interglacial periods in the Pleistocene? (iii) In species that have an Eastern origin, do the most drastic demographic changes occur after the colonization of Iberia, or are they shared with other refugial populations, suggesting that genetic diversity in Iberia largely reflects the size of the ancestral source populations?

2 | MATERIALS AND METHODS

2.1 | Samples and sequencing

We analysed whole-genome Illumina paired end resequencing data for six (haploid) male individuals in each of seven species of chalcid parasitoids (spanning five families): *Megastigmus dorsalis* and *M. stigmatizans* (*Megastigmidae*), *Torymus auratus* (*Torymidae*), *Ormyrus nitidulus* and *O. pomaceus* (*Ormyridae*), *Eurytoma brunnenensis* (Eurytomidae) and *Cecidostiba fungosa* (Pteromalidae) (Table S1). The *M. dorsalis* individuals correspond to cryptic species 1 for this complex, as defined by Nicholls et al. (2010). For each species, we sampled five individuals from Iberia (the focal refugium) and one from Hungary (the Balkan refugium). Data for the Hungarian and two Iberian samples of each species were generated previously by Bunnefeld et al. (2018). We generated analogous data for the remaining three Iberian individuals using the protocols described by Bunnefeld et al. (2018). DNA was extracted from individual wasps using the Qiagen DNeasy kit. Individual Nextera genomic libraries were generated and sequenced on an Illumina HiSeq 2000 by the NERC Edinburgh Genomics facility, UK. Raw reads were deposited at the SRA (PRJEB20883). Mean coverage per haploid individual ranged from 3.5x to 18.5x. Raw reads were mapped back to reference genomes assembled by Bunnefeld et al. (2018) using BWA (0.7.15-r1140) (Li & Durbin, 2009), duplicates were marked with samtools picard (V2.9.0) MarkDuplicates (broadinstitute.github.io/picard/), and variants were called using Freebayes (v1.1.0-3-g961e5f3) (https://github.com/KLohse/BunnefeldEtAL_2018/block_cutter_vcf.py) with a minimum base quality of 10 and a minimum mapping quality of 20 (see Bunnefeld et al. (2018) for further details).

2.2 | Inferring single-step-change models in population size

We fitted a model of a single, instantaneous step change in \( N_e \) using the framework for blockwise likelihood calculations developed by Lohse et al. (2011). The single-step-change model includes three parameters: the scaled mutation rate \( \theta = 4N_e\mu l \) (where \( N_0 \) is the current \( N_e \) and \( l \) is block length); \( T \), the time of \( N_e \) change measured in \( 2N_e \) generations; and \( \lambda = N_0/N_e \), the relative magnitude of the \( N_e \) change (where \( N_1 \) denotes the \( N_e \) prior to the step change).

Following Bunnefeld et al. (2015), we summarized sequence variation in short blocks of a fixed length \( l \) by the (folded) blockwise site frequency spectrum (bSFS) using the script block_cutter_vcf.py (https://github.com/KLohse/BunnefeldEtAL_2018/block_cutter_vcf.py). Given our sampling scheme of \( n = 5 \) haploid males, the bSFS consists of counts of two types of variants: those for which the
minor allele occurs once or twice in the sample. The probability of observing a particular set of mutations in a block can be computed analytically as a higher-order derivative of the generating function of genealogies (Lohse et al., 2011). The product of probabilities of bSFS configurations across blocks can be interpreted as the composite likelihood (CL) of the model. We maximized lnCL, i.e. model support, in Mathematica (Wolfram Research, 2016) using the function NMaximise (Data S1).

To generate blockwise data for each species, we applied the same quality filters used for calling SNPs to all sites; that is, we identified regions of the genome with a base quality >10 and mapping quality >20 in each individual from bam files by the CallableLoci walker of GATK (v3.4) (Van der Auwera et al., 2013). Only regions meeting these criteria in all five Iberian individuals were included in further analyses. Custom scripts were used to partition the data into blocks of a fixed length of callable sites. l was chosen to be inversely proportional to the pairwise genetic diversity of each species (Table 2) such that blocks, each capturing a random configuration of closely linked variants had an average two pairwise differences between samples. This ensures that the information content per block, as well as any effect of intra-block recombination, is consistent across species. Blocks with a physical span (including non-callable sites) of >2l or more than five ‘None’ (uncalled) sites were removed.

We assessed support for a step change in \( N_e \) relative to the (nested) null model of constant \( N_e \) by generating parametric bootstrap replicates with the coalescent simulator msprime (Kelleher et al., 2016). For each species, 100 replicate data sets were simulated under the null model assuming estimates of recombination inferred by Bunnefeld et al. (2018) (Table S3). Each data set had the same total length as the real data (after filtering) and was partitioned into 5,000 windows of sequences (for computational efficiency). Both a null model of constant \( N_e \) and a step-change model were fitted to each replicate, and the 95% quantile of the difference in support between models was compared to that of the real data. Confidence intervals (CIs) of parameter estimates were obtained via an analogous parametric bootstrapping procedure: we simulated 100 data sets with recombination under the inferred step-change model and fitted that model to each simulation replicate. CIs were obtained as 2.5% and 97.5% quantiles of the distribution of parameter estimates and were centred around the point estimates of parameters obtained from the real data.

### 2.3 | Reconstructing ancestral population size with PSMC

The pairwise sequentially Markovian coalescent (PSMC) (Li & Durbin, 2011) was used to infer a history of population size change for each species. PSMC is a non-parametric method that reconstructs a trajectory of past \( N_e \) from the density of pairwise differences along the genome via a hidden Markov model in which the hidden states are pairwise coalescence times, the distribution of which is used to estimate \( N_e \) in discrete time intervals. In each species, the two Iberian individuals with the greatest average read depth were used as the focal pair. Pileup files were generated using samtools mpileup (version 0.1.19) (Li et al., 2009), and a consensus sequence (fastq file) per individual was generated using the bcftools utility vcf2fq. Regions covered in both individuals were combined into fastq files using seqtk mergefq (https://github.com/lh3/seqtk) and converted into PSMC input files. PSMC, by default, discretizes pairwise alignments into blocks of 100 bp which are encoded as variant if they contain at least one variant. While this makes analyses of large genomes with low diversity (e.g. humans) computationally efficient, this discretization is too coarse when considering more diverse genomes where the chance of several pairwise differences occurring in the same 100 bp block is non-negligible. We investigated the effect of varying block length (100, 50, 25 and 1 bp) on \( N_e \) inference for the most diverse (E. brunniIVentris; \( \pi = 0.0071 \)) and the least diverse (M. stigmatizans; \( \pi = 0.00067 \)) species in our set. As expected, population trajectories showed higher \( N_e \) and were pushed back in time with smaller block lengths (Figure S1). We chose a block length of 25 bp for all analyses which minimizes these biases without excessive computational cost.

We inferred 30 free interval parameters across 64 time intervals (with the option `-p ‘28’2+3+5’). The maximum coalescence time (–t) was set to 5, the initial value of \( \theta/\rho \) (–t) to 1. 100 bootstraps were performed for each run. Times of peak \( N_e \) and values of \( N_e \) in each time interval were considered to be different between species if

#### TABLE 2 | Summaries of PSMC trajectories for Iberian populations of seven species of chalcid parasitoid wasps. Times are given in thousands of years ago (kya). 95% CIs of peak \( N_e \) times are shown in brackets. The maximum \( N_e \) is scaled relative to \( \pi \) as \( \text{Max}(N_e)/N_e^{\pi} \). |

| Species            | \( \pi \) | Peak \( N_e \) \( \times 10^3 \) | Peak \( N_e \) time (kya) | Split time (kya) | Relative Max \( N_e \) |
|--------------------|----------|-------------------------------|--------------------------|-----------------|---------------------|
| Torymus auratus    | 0.0064   | 835                           | 78 (74–82)               | n/a             | 1.81                |
| Megastigmus stigmatizans | 0.00067 | 65                            | 47 (42–52)              | 37              | 1.34                |
| Megastigmus dorsalis | 0.0018  | 139                           | 35 (31–39)              | 128             | 1.07                |
| Ormyrus nitidulus  | 0.0016   | 157                           | 114 (104–124             | 132             | 1.36                |
| Ormyrus pomeaeus    | 0.0028   | 275                           | 35 (30–40)              | 681             | 1.36                |
| Eurytoma brunniVentris | 0.0071 | 831                            | 46 (36–56)              | n/a             | 1.62                |
| Cecidostiba fungosa | 0.0032   | 213                           | 25 (12–38)              | 267             | 0.92                |
there was no overlap in bootstrap replicates. To be able to compare the magnitude of inferred \( N_e \) change between PSMC and single-step-change analyses, we normalized the maximum \( N_e \) inferred by each method by the long-term average \( N_e \) estimated from \( \pi \) Nei, 1972 (Table 2) as \( \bar{N}_e = \pi / (4\mu) \) and computed the following measure of \( N_e \) change: \( \max(\bar{N}_e) / \bar{N}_e \). Unlike the size of the step change (\( \lambda \)), this measures the maximum \( N_e \) relative to the average over the entire history of a sample and is therefore expected to be smaller than \( \lambda \).

### 2.4 Calibrating the timing of events

Time estimates were converted into years assuming a mutation rate of \( 3.46 \times 10^{-9} \) mutations per base per generation estimated for *Drosophila melanogaster* (Keightley et al., 2009). We assumed two generations per year for all species with the exception of *M. stigmatis* which has a single generation per year (Stone et al., 2012). While this calibration allows comparisons with the estimates obtained by Bunnefeld et al. (2018) (calibrated in the same way), these absolute times are likely underestimates and should be treated with caution (given that we use a spontaneous mutation rate estimate and assume selective neutrality). We stress, however, that comparing the relative timing of demographic events across this set of parasitoids only relies on the assumption of the same per generation mutation rate across species rather than any absolute calibration.

### 2.5 Cross-population PSMC analyses

To test whether population size changes in each species occurred before or after the divergence between the Iberian and Balkan refuge populations, we compared the PSMC trajectories of Iberian pairs and cross-population pairs (one Iberian and one Hungarian individual). Any divergence between Iberian and the Balkan refuge populations should be visible as a separation between within-population (Iberian pair) and cross-population (Iberia-Balkan pair) PSMC trajectories. Specifically, in the absence of significant post-divergence gene flow, we expect \( N_e \) estimates for the cross-population pairs to increase from the time of the population split due to the accumulation of genetic differences between the populations. Thus, population size changes that occurred in an ancestral population (potentially outside Iberia) should predate the divergence of within and cross-population trajectories. In contrast, we would expect demographic events unique to the Iberian population to happen after divergence of the within- and cross-population PSMC trajectories. We also compared our PSMC divergence time estimates to those made by Bunnefeld et al. (2018) under explicit models of divergence and admixture.

### 2.6 Potential population structure in Iberia

Population structure within any assumed panmictic population can confound inferences of past demography (Bunnefeld et al., 2015; Gatetpaille et al., 2013). In particular, \( N_e \) estimates may be inflated due to divergence between demes (Gatetpaille et al., 2013). Likewise, when samples are taken from the same deme, the (structured) coalescent generates a mixture of very recent within-deme ancestry and older ancestry resulting from migration between demes (Wakeley, 2008), which can mimic the signatures of a bottleneck (Bunnefeld et al., 2015). To test for population structure within Iberia, we repeated PSMC analyses on every pairwise combination of our five individuals. In a structured population, and given that the sampling locations were widely spaced across Iberia (Table S1), \( N_e \) trajectories are expected to differ between pairs of haploid males sampled from the same versus different sub-populations (analogous to the within- and between-population analyses involving Iberian and Hungarian samples).

### 2.7 Sensitivity analyses

The assumption of no recombination within blocks (required to make the composite likelihood estimation of the single-step-change model tractable) potentially biases parameter estimates. Specifically, recombination may lead to a downward bias of \( N_e \) estimates (Bunnefeld et al., 2015; Wall, 2003). We used simulations in msprime (Kelleher et al., 2016) to quantify the effect of recombination on parameter estimates (Table S2). One million unlinked blocks of 586 bp (corresponding to the block length used for *O. nitidulus*) were simulated under a step-change model with different recombination rates and step sizes.

It is well known that both selective sweeps (Smith & Haigh, 1974) and background selection (Charlesworth et al., 1993) affect variation at neutral sites in the genome which, in turn, can bias demographic inference (Ewing & Jensen, 2016; Schrider et al., 2016). To explore the effect of selection on the single-step-change analyses, we fitted step-change models separately to blockwise data generated from genic and intergenic regions of the *O. nitidulus* genome. Genes were predicted ab initio with Augustus (Stanke & Morgenstern, 2005). *Nasonia vitripennis*, a chalcid parasitoid (family Pteromalidae), was used as a training set. If selection has a strong effect on demographic inference, estimates of \( \theta \) are expected to be lower for genic compared to intergenic regions, as most selection tends to reduce diversity both at selective targets and linked regions of the genome (Smith & Haigh, 1974). Similarly, we would expect estimates for the time of the step change, \( T \), to be biased towards the present in genic compared to intergenic regions.

### 3 RESULTS

After filtering, the blockwise data sets ranged in total length from 54 to 151 Mb (Table S3). The pairwise alignments used as input for PSMC spanned a total length of 161 to 379 Mb (Table S3). Despite this difference in overall length, which is mainly due to missing data and the difference in sample size (two versus five individuals),
estimates of average pairwise diversity, as measured by \( \pi \) (Nei, 1972), agreed well between the two data sets (Table 1). Across species, \( \pi \) estimates spanned an order of magnitude, from 0.00067 in \( M. stigmatizans \) to 0.0071 in \( E. brunniventris \).

### 3.1 Large changes in \( N_e \) detected in four species

Taking the results of the single-step-change and PSMC analyses across all seven parasitoid species together, four species (\( M. stigmatizans \), \( M. dorsalis \), \( O. nitidulus \) and \( T. auratus \)) show evidence for large (at least a factor of three) changes in population size during the Quaternary period (Figure 1). In these species, an instantaneous step change in \( N_e \) fits the blockwise data significantly better than a null model of a constant \( N_e \). We infer a decrease in \( N_e \) towards the present (\( \lambda < 1 \)) in three species (\( M. stigmatizans \), \( M. dorsalis \) and \( O. nitidulus \)) and an increase in one species (\( T. auratus \) (\( \lambda > 1 \)) (Figure 1 and Table 1). In all four species, the change in \( N_e \) in the PSMC trajectory (Figure 1) agrees both in direction and in coarse timescale with the inference under the single-step-change model. However, the \( N_e \) changes visible in the PSMC trajectory for these species are not equally abrupt. For example, \( T. auratus \) shows a gradual increase of \( N_e \) over a period of more than 200 ky, while the decreases in the PSMC trajectories of \( M. dorsalis \) and \( M. stigmatizans \) are comparatively sudden. PSMC trajectories and single-step-change analyses are also broadly similar for \( O. pomaceus \) and \( C. fungosa \): a step change in \( N_e \) is supported in neither species, and PSMC suggests comparatively small population size changes for both (Table 1 and Table 2).

However, while we have resisted the temptation to attempt any formal comparison between PSMC trajectories and single-step-change analyses which would necessarily be post hoc and potentially misleading (given the opposing assumptions of these methods, see

![Figure 1](image.png)

**Figure 1** PSMC and maximum-composite likelihood estimates (MCLE) under the single-step-change model for Iberian populations of seven species of chalcid parasitoid wasps. PSMC estimates and bootstrap replicates are shown in dark red and pale red, respectively. Population sizes estimated using the composite likelihood step-change model are shown in blue with 95% CIs in grey. The three species with green plot frames are those for which a step-change model is not supported. Results for \( M. stigmatizans \) are also plotted on an alternative timescale to reveal recent \( N_e \) change (bottom right).
Discussion}, inferences under the two methods disagrees qualitatively in at least two obvious ways:

First, with the exception of *M. stigmatizans*, the magnitude of $N_e$ change inferred under the step-change model is greater than the relative magnitude of peak $N_e$ in the PSMC trajectories (Figure 1, Table 1 and Table 2). Second, in one species, *E. brunniventris*, single-step-change and PSMC analyses are hard to reconcile: we found no significant support for a step change in this species, that is, we cannot reject a null model of a single fixed $N_e$. Yet, the PSMC trajectory indicates a substantial (but gradual) increase in $N_e$. Additional analyses (see Discussion) suggest that demographic inferences for *E. brunniventris* may be affected by genetic structure and/or the low contiguity of its reference genome.

### 3.2  No support for temporal concordance of population size change

The four species with support for a step change in $N_e$ (*M. stigmatizans*, *M. dorsalis*, *O. nitidulus* and *T. auratus*) show no overlap in the estimated times of $N_e$ change (Table 1). Using an insect spontaneous mutation rate (Keightley et al., 2009) to calibrate $T$ estimates (see Methods for caveats), Iberian populations of *M. stigmatizans* and *M. dorsalis* most likely decreased in size at the start of the current interglacial (10 and 12.5 kya, respectively). In contrast, the decrease in $N_e$ inferred for *O. nitidulus* most likely dates to the last glacial period (33 kya) and the increase in $N_e$ in *T. auratus* at 100 kya falls to just after the end of the (Eemian) interglacial (Table 1). Formalizing this comparison, we conclude that a maximally complex model that assumes a unique time of $N_e$ change for each of the four species fits significantly better than any scenario involving concordant/clustered times of $N_e$ change.

Comparing times of $N_e$ change inferred via PSMC across species is inherently problematic (see Discussion). However, when comparing the time of maximum $N_e$ by computing the overlap in peak $N_e$ intervals across bootstrap replicates, we find that the four taxa *M. stigmatizans*, *M. dorsalis*, *O. nitidulus* and *T. auratus* which show support for a single-step change at unique times also have non-overlapping peak $N_e$ times (Table 1). Interestingly, the three species with no support for a single-step change (*O. pomaceus*, *C. fungosa* and *E. brunniventris*) and *M. dorsalis* show highly similar (overlapping CIs) times of peak $N_e$ (Figure 2 and Table 2).

### 3.3  Changes in $N_e$ occur after the divergence of Iberian populations

In five species, the PSMC trajectories of within-population (Iberia) pairs diverge clearly from the cross-population (Iberia versus Hungarian) trajectories (Figure 3). In contrast, little divergence between within and cross-population PSMC trajectories is visible for *E. brunniventris* and none for *T. auratus* (Figure 3). In all three species for which the single-step-change analyses support a decline in population size, the inferred time of $N_e$ change is more recent than the divergence of within- and cross-population PSMC trajectories (Figure 3 and Table 2) and so must have occurred after Iberian and Balkan populations split. For *O. pomaceus*, *C. fungosa* and *M. stigmatizans*, the split times between Iberian and the Balkan populations inferred here post hoc by comparing PSMC trajectories are broadly compatible with the divergence estimates obtained by Bunnefeld et al. (2018) under explicit models of population divergence (Figure 3). In contrast, for both *M. dorsalis* and *O. nitidulus*, the divergence of within and between refuge PSMC trajectories substantially predates the split times inferred by Bunnefeld et al. (2018). In both cases, cross-population PSMC trajectories decrease after divergence between the Iberian and Hungarian populations, suggesting that these populations have been connected by a period of gene flow immediately following divergence which may not be detectable given the admixture pulse scenarios fitted by Bunnefeld et al. (2018).

### 3.4  No evidence for structure within Iberia

We find little variation in PSMC trajectories between different Iberian pairs in almost all species (Figure S2), suggesting that our demographic inferences are unlikely to be influenced by population structure within Iberia. For all species except *E. brunniventris*, the
differences in PSMC trajectories between different Iberian sample pairs are similar to the differences across bootstrap replicates for the focal pair (Figure 1 and Figure S2) and so likely reflect coverage variation between individuals. *E. brunniventris* is the only species that showed clear variation in PSMC trajectories between Iberian pairs. While our additional analyses for *E. brunniventris* suggest other potential reasons for this finding (see Discussion), we cannot rule out that this variation is in part driven by population structure.

### 3.5 Sensitivity analyses

Both demographic inference methods used here assume selective neutrality but make different simplifying assumptions about recombination: while PSMC approximates recombination as a Markov process along the genome, the single-step-change analyses based on the bSFS assumes no recombination within blocks. To check the extent to which recombination and selection may bias parameter estimates, we fitted histories of a step change to data simulated with recombination. Our exploration of simulated data shows that both \( N_e \) and \( \lambda \) are underestimated with increasing recombination rate whereas the time of step change (\( T \)) is biased downwards when \( N_e \) declines towards the present (\( \lambda < 1 \)) and upwards when \( N_e \) increases (\( \lambda > 1 \)) (Table S2). These biases are an expected consequence of the shuffling of genealogical histories within blocks in the presence of recombination, which reduces the variance in bSFS configurations. Given a recombination rate of \( r = 3 \times 10^{-9} \) (Table S3), our simulation check suggests that while \( T \), the time of the step change may be underestimated by up to a factor of two, the ability to accurately estimate \( \lambda \), the magnitude of the step change, is little affected.
To investigate the potential effect of selective constraint on the single-step-change analyses, we inferred single-step-change histories separately for genic and intergenic regions. Parameter estimates based on intergenic data for O. nitidulus were broadly similar to those obtained from the full data set (Table S2): while estimates of $\theta$ and $T$ were slightly lower and higher, respectively, for the full data set, as would be expected as a result of selective constraint, the estimate of $\lambda$ was little affected.

4 | DISCUSSION

We analysed genome-wide sequence variation using two contrasting inference approaches to test whether and how demographic histories vary within a guild of insect parasitoids in a single Pleistocene refugium. We find evidence for drastic declines in population size in three out of seven species (M. stigmatizans, M. dorsalis, O. nitidulus) and a large increase in population size in one (T. auratus). Interestingly, these four species encompass a mixture of longitudinal expansion histories: while Bunnefeld et al. (2018) inferred that M. dorsalis and O. nitidulus expanded into Iberia from the East, M. stigmatizans most likely expanded out of Iberia and T. auratus showed no longitudinal expansion signal. This suggests that the history of population size change in Iberia is not determined in any obvious or simple way by whether Iberia was the end point of an initial expansion into Europe or not. Contrary to any simple scenario of guild-wide temporal concordance in responses to Pleistocene climatic events, we find that population size changes of species in this guild differ markedly both in direction and in timescale. In fact, our single-step-change analyses reveal significant support for maximal temporal discordance; that is, each of the four species in this parasitoid guild that show evidence for a sudden change in $N_e$ has a unique time (Table 1).

Thus, our main result of temporally discordant $N_e$ change within Iberia mirrors the finding of temporal discordance by Bunnefeld et al. (2018) in the timing of divergence and admixture for this guild on a continental scale. One could argue that the signal of temporal discordance may simply be an artefact of applying an oversimple step-change model, which is unlikely to capture the subtleties of real population size change. However, the fact that we also find evidence for temporal discordance when visualizing $N_e$ change using PSMC, which makes minimal simplifying assumptions about the shape of past demographic change, suggests a genuine lack of signal for temporal concordance in this parasitoid guild across a range of spatio-temporal scales. This general finding mirrors results of other comparative studies on sets of co-distributed taxa in Europe (Ebdon et al., 2021) and the Americas (Burbrink et al., 2016; Dasmahapatra et al., 2010).

It is interesting that the only potential signal of temporal concordance we find is for four species that show the smallest change in past population sizes and which overlap in the time of peak $N_e$ (Figure 2). While this apparent congruence may be due to a shared background demography, the signatures of which are masked in species with more drastic changes in $N_e$, we cannot rule out alternative explanations. In particular, the fact that PSMC assumes selective neutrality is problematic given that insect genomes are more compact than mammalian genomes (Li & Graur, 1991), and so more susceptible to the effects of linked selection. Schrider et al. (2016) have shown that selective sweeps can generate troughs in PSMC trajectories while background selection has been shown to lead to erroneous inference of population growth (Ewing & Jensen, 2016). We therefore cannot rule out the possibility that congruent peaks in $N_e$ are an artefact of selective effects which, assuming similar genome composition, mutation and recombination rates, may lead to similarly distorted PSMC inferences. However, the fact that we have reconstructed strikingly different PSMC trajectories, most of which differ markedly from the selection-induced PSMC trajectories of Schrider et al. (2016) (in that they show large declines rather than increases in $N_e$ towards the present), suggests that it is unlikely that linked selection is the main driver of the inferred $N_e$ changes. For the single-step-change analyses, we find little difference in parameter estimates when analysing all data or just intergenic regions (Table S2), which again argues against a major effect of selection on our inferences. Furthermore, one would expect genomes with a shorter map length (physical length × recombination rate, see Table S3) to be disproportionately affected by linked selection (Mackintosh et al., 2019; Smith & Haigh, 1974). However, if anything, we observe the opposite pattern: the two species with shortest map lengths (O. nitidulus and C. fungosa) show less pronounced $N_e$ change than the two species with the longest map length (M. stigmatizans and M. dorsalis).

4.1 | Reconciling single-step-change and PSMC analyses

PSMC and single-step-change analyses exploit different aspects of the data and differ drastically in sampling schemes (contiguous pairwise alignments vs short blocks sampled across five individuals) and underlying assumptions: while the blockwise composite likelihood framework fits a single instantaneous step change in $N_e$ and infers the timing of this event, PSMC does not estimate a time parameter per se, but rather imposes an arbitrary discretization of time and reconstructs population size change as a continuous trajectory. It is reassuring that despite these fundamental differences, both methods yield broadly congruent conclusions: the four species for which the single-step-change analyses diagnose an abrupt change in $N_e$ also show PSMC trajectories with large $N_e$ changes in the same direction and at similar times. The greater magnitude of $N_e$ change for the single-step-change compared to the PSMC analyses may be a consequence of the fact that larger samples used to fit the single-step-change model contain more information about recent demography than a pair of lineages.

In general, one may view the fact that PSMC is essentially assumption-free as an advantage over model-based inference, because it enables a straightforward visualization of past $N_e$ change. Similarly, comparing PSMC trajectories between pairs of individuals and populations allows qualitative assessment of likely periods of
dissolved and admixture (Figure 3). However, the flip-side of this
flexibility is that PSMC provides no obvious route for quantitatively
testing (necessarily) simple demographic hypotheses across species.
Furthermore, cross-species comparisons of PSMC trajectories are
problematic even when they are focused on a single clearly defined
summary such as the interval of peak $N_e$ (as we have done here),
because the time discretization PSMC imposes depends on the rate
of coalescence, and so differs between species.

4.2 | Changes in population size coincide with late
Pleistocene climatic transitions

While we emphasize that absolute time estimates need to be inter-
preted with caution, our estimates of the timing of drastic $N_e$ changes
coincide broadly with climatic events in the Quaternary: the start of
the Holocene around 11 kya and the end of the Eemian interglacial
around 106 kya (Figure 3). Previous studies on the parasitoid oak
gall wasp community based on the same calibration have inferred
an increase in gene flow between refugia during the same time peri-
dods (Bunnefeld et al., 2018; Lohse et al., 2010, 2012). For example,
the cluster of peak $N_e$ times coincides with a large community-wide
peak in the frequency of admixture between refugia inferred by
Bunnefeld et al. (2018). Similarly, the large increase in $N_e$ inferred for
T. auratus at the end of the Eemian interglacial period coincides
with the divergence of refugial populations inferred for this spe-
cies previously (Bunnefeld et al., 2018; Stone et al., 2012). It seems
plausible that both events are associated with the geographic ex-
pansion of suitable oak habitat across refugial barriers during these
times (Brewer et al., 2002; Petit et al., 2002), which may have trig-
gerated range expansions in both host gall wasps and their associated
parasitoids.

4.3 | Population structure within Iberia and gene
flow from Eastern refugia

Population structure within southern European refugia has previ-
ously been demonstrated for some species, and it has been sug-
gested that given its complex topography Iberia should be considered
a mosaic of multiple micro-refugia rather than a single entity (Feliner,
2011; Hearn et al., 2014). However, our PSMC results suggest a com-
plete lack of population structure in six out of seven species in the
parasitoid guild (Figure S2) and imply high gene flow within Iberia.
This is perhaps unsurprising given that gall wasp-associated para-
sitoid wasps (and other chalcids) are able to disperse long distances
even across patchy habitats and host distributions (Compton et al.,
2000; Hayward & Stone, 2006).

Our estimates of population split times based on comparisons
of within- and cross-population PSMC trajectories agree broadly
with those of Bunnefeld et al. (2018) for most species. M. dorsalis
and O. nitidulus, the two species that show the least agreement with
past estimates, both show decreases in cross-population $N_e$ after
divergence that are compatible with ongoing gene flow between re-
fugia. The model space considered by Bunnefeld et al. (2018) was
limited to histories involving a single discrete burst of instantaneous
admixture between refugial populations, with limited potential to
detect periods of continuous post-divergence gene flow.

4.4 | Eurytoma brunniventris is an outlier

Eurytoma brunniventris is an outlier in our results in several ways: it is
the species with the highest genetic diversity, shows signals of popu-
lation structure within Iberia and is the only species for which our two
inference approaches disagree. While the single-step-change analy-
ses give no support for a change in $N_e$, the PSMC trajectory shows a
steady increase. To explore the sensitivity of the single-step-change
analysis to detect gradual changes in population size, we simulated
100 replicate data sets (assuming the same size and block length as
the real data) under the gradual change in $N_e$ inferred via PSMC for
E. brunniventris. We find that in each case, a single-step-change his-
tory fits the data significantly better (using a parametric bootstrap
analogous to that performed on the real data, see Methods) than a
null model of constant $N_e$, indicating that the single-step-change
analysis is indeed sensitive to gradual increases in population size.
A possible explanation for the discrepancy between the PSMC and
single-step-change analyses for E. brunniventris is that its genome
assembly, the least contiguous among our set of taxa (Table S3), is
too fragmented for reliable PSMC inference. Interestingly, both a
history of constant $N_e$ and a single-step-change model give a poor
fit to the observed frequency of $bSFS$ configurations in E. brunnin-
ventris. In particular, E. brunniventris shows an excess of both monomor-
phic blocks and blocks with a large number of variants (Figure S3),
suggesting that its history is not well approximated by any model
that assumes a single panmictic population. The lack of divergence of
within- and between-population PSMC trajectories for E. brunnin-
ventris would be compatible with substantial gene flow between the
Iberian and Hungarian populations (Figure 3). Alternatively, E.
brunniventris—an extreme generalist attacking a wide range of oak
gall wasp hosts (Askew et al., 2013)—may harbour genetic structure
as a result of recent divergence into cryptic host races (Nicholls et al.,
2018). However, in the absence of a better reference genome and
larger sample sizes, it remains unclear to what extent the disagree-
ment between PSMC and single-step-change analyses for E. brunnin-
ventris is indicative of a more complex history.

4.5 | Outlook

A general question for geographically widespread communities is
the extent to which component species continue to interact dur-
ing the assembly process (Agosta & Klemens, 2008; Janzen, 1985;
Ricklefs, 2015). Where communities are characterized by strong de-
pendencies between species, such as specific trophic or symbiotic
interactions, we might expect co-dispersal and coupled population
dynamics to result in similar demographic histories (e.g. Gaume et al., 2000); this scenario is compatible with ongoing selective effects of species on each other and potential coevolution (Hall et al., 2020; Wade, 2007).

In contrast, we might expect much lower demographic concordance for members of communities characterized by weaker and less specific species interactions and only diffuse coevolution (Hall et al., 2020). This is the pattern emerging for the parasitoid assemblies attacking oak gall wasps. Though comprising a consistent set of interacting taxa spanning thousands of kilometres of longitude, the component species show highly diverse histories of range expansion (Bunnefeld et al., 2018) and changes in population size. This finding is consistent with the ability of most of the parasitoid species to attack multiple gall wasp hosts (Askew et al., 2013), weakening both direct interactions between the trophic levels and competitive interactions between parasitoid species. While there is strong evidence that host gall wasp traits structure-associated assemblages of parasitoid enemies (Bailey et al., 2009) and of parasitoid specializations for exploitation of particular hosts (Weinersmith et al., 2017), there is little evidence for specific coevolution. The population histories of very few other multitrophic communities have been explored, and the extent to which the demographic diversity within the gall wasp system is typical of parasitoid–host and other biological systems remains unknown. Work on the arthropod communities associated with the pitcher plant Sarracenia alata (Sattler & Carstens, 2017) also shows variation in component species histories; further, this study suggests that more ecologically dependent associates show closer demographic and phylogeographic concordance with the host plant than less ecologically dependent species. An obvious question is whether any of the variation in patterns of $N_e$ change in our sampled parasitoid species can be attributed to variation in ecological traits (Papadopoulou & Knowles, 2016). Intriguingly, the four parasitoid species with support for a step change (in either direction) have a narrower host range (Askew et al., 2013) and a lower ancestral $N_e$ than those for which a null model of constant $N_e$ could not be rejected (Table S1). Although the number of species studied here is clearly insufficient for any statistical test of an association between host range and demographic history, these trends are compatible with the idea that ecologically specialist species (whose biology is critically dependent on interactions with a small number of other taxa) experience greater and/or more frequent changes in $N_e$ than generalists (whose demography is less tightly coupled to abundance of any specific interaction) (Östergård & Ehrlén, 2005; Rand & Tschamkite, 2007). Larger samples of species, incorporating wide diversity in host number and other relevant traits (such as dispersal ability) are required to assess the extent to which life history traits and demographic histories are correlated. Given that the genomes of parasitoid wasp can be sampled in a haploid state (by targeting males), it will be fascinating to test how much more signal about the demographic past of ecological communities can be extracted using a new generation of inference approaches that reconstruct the sequence of correlated genealogies directly from the data (Kelleher et al., 2019; Speidel et al., 2019).

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AUTHOR CONTRIBUTIONS
KL and GS designed the project; WW analysed the sequence data with contributions from KL; all authors wrote the manuscript.

DATA AVAILABILITY STATEMENT
• Raw reads have been deposited in the European Nucleotide Archive (ENA) (ERP023079) and the SRA (PRJEB20883)
• Genome assemblies are deposited in the ENA (PRJEB27189 and ERP109243)
• Mathematica notebook and blockwise data are available as Supporting Information

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