Identification of winter moth (Operophtera brumata) refugia in North Africa and the Italian Peninsula during the last glacial maximum

Jeremy C. Andersen1 | Nathan P. Havill2 | Yaussra Mannai3 | Olfa Ezzine4 | Samir Dhahi3 | Mohamed Lahbib Ben Jamâa3 | Adalgisa Caccone5 | Joseph S. Elkinton1

Abstract

Numerous studies have shown that the genetic diversity of species inhabiting temperate regions has been shaped by changes in their distributions during the Quaternary climatic oscillations. For some species, the genetic distinctness of isolated populations is maintained during secondary contact, while for others, admixture is frequently observed. For the winter moth (Operophtera brumata), an important defoliator of oak forests across Europe and northern Africa, we previously determined that contemporary populations correspond to genetic diversity obtained during the last glacial maximum (LGM) through the use of refugia in the Iberian and Aegean peninsulas, and to a lesser extent the Caucasus region. Missing from this sampling were populations from the Italian peninsula and from North Africa, both regions known to have played important roles as glacial refugia for other species. Therefore, we genotyped field-collected winter moth individuals from southern Italy and northwestern Tunisia—the latter a region where severe oak forest defoliation by winter moth has recently been reported—using polymorphic microsatellite. We reconstructed the genetic relationships of these populations in comparison to moths previously sampled from the Iberian and Aegean peninsulas, the Caucasus region, and western Europe using genetic distance, Bayesian clustering, and approximate Bayesian computation (ABC) methods. Our results indicate that both the southern Italian and the Tunisian populations are genetically distinct from other sampled populations, and likely originated in their respective refugium during the LGM after diverging from a population that eventually settled in the Iberian refugium. These suggest that winter moth populations persisted in at least five Mediterranean LGM refugia. Finally, we comment that outbreaks by winter moth in northwestern Tunisia are not the result of a recent introduction of a nonnative species, but rather are most likely due to land use or environmental changes.
1 | INTRODUCTION

Natural movements of organisms following the expansions and contractions of glacial ice sheets during the Quaternary climatic oscillations have resulted in the geographic and genetic isolation of species/populations across temperate regions (Hewitt, 1996, 2000; Schmitt, Rober, & Seitz, 2005; Schmitt & Seitz, 2001). For example, in Europe, many studies have shown how the persistence of populations in separate glacial refugia (e.g., the Iberian, Italian, and Balkan peninsulas, the Caucasus region, North Africa, and microrefugia in the Alps) during the Quaternary climatic oscillations promoted genetic diversity in contemporary populations (Andersen, Havill, Cacconne, & Elkinton, 2017; Habel, Lens, Rodder, & Schmitt, 2011; Huck, Buedel, & Schmitt, 2012; Niedziakowska et al., 2016; von Reumont, Struwe, Schwarzer, & Misof, 2012; Rofes et al., 2018; Schmitt, 2007; Schmitt & Müller, 2007; Sim, Hall, Jex, Hegel, & Colman, 2016; Stolting et al., 2015; Torroni et al., 2001). For outbreaking species, reconstructing the biogeographical history of the species can help to determine the factors that might be promoting their outbreak (e.g., Moyal et al., 2011; Song et al., 2018; Zhu et al., 2016).

One species that frequently reaches outbreak densities across much of its native range of Europe and western Asia (Elkinton, Boettener, Liebhold, & Gwiazdowski, 2015) is the winter moth, Operophtera brumata (Lepidoptera: Geometridae). Throughout its distribution, winter moth is a serious pest of many broadleaf tree and shrub species, including oak (Quercus spp.), maple (Acer spp.), and birch (Betula spp.) trees (Wint, 1983). Winter moth is also present in North Africa, where it was first reported in the 1970s (Ferguson, 1978; Hausmann & Viidalepp, 2012). More recently it has been observed at outbreak densities in oak forests (Quercus spp.) in northwestern Tunisia beginning in 2009 (Mannai, Ezzine, Nouira, & Ben Jamâa, 2015), though the reasons for these outbreaks are currently unknown. In this region, mature oak forests extend mainly to the regions of Kroumirie and Mogods, from the Algerian–Tunisian border in the west, to Lake Ichkeul in the east (Debazac, 1959; DGF, 2005), and are home to five oak species: Quercus suber L., Quercus canariensis Willd, Quercus ilex L., Quercus coccifera L., as well as a Tunisian–Algerian endemic species Quercus afares Pomel (Boudy, 1950). In Tunisia, winter moth has mainly been observed on Q. canariensis in both mixed oak (Ain Zena, 36.435°N, 8.515°E) and pure Q. canariensis oak (Mzara Forest, 36.769°N, 8.72°E) forests, with lower densities being found on Q. afares and Q. suber compared with Q. canariensis in Ain Zena. These differences may be the result of differences in budburst timing of different oak species (Mannai, Ezzine, Hausmann, Nouira, & Ben Jamâa, 2017).

As a result of the movement of nursery stocks, invasive populations of winter moth have also been introduced to North America (Ferguson, 1978), with populations established in Nova Scotia in the 1920s (MacPhee, 1967), Oregon in the 1950s (Kimberling, Miller, & Penrose, 1986), British Columbia in the 1970s (Gillespie, Finlayson, Tonks, & Ross, 1978), and in the northeastern United States in the 1990s (Elkinton et al., 2015). Therefore, we were curious as to whether or not the recent reports of outbreaks in northwestern Tunisian oak forests were the result of a recent introduction (as has been the case in North America), or whether a native population has achieved outbreak levels for some other reason. Previously, Mannai et al. (2017) explored the native/nonnative status of outbreaking Tunisian winter moth populations by sequencing a fragment of the mitochondrial (mtDNA) locus cytochrome oxidase I (COI) from field-collected larvae in Tunisian oak forests, and compared these sequences to those published in NCBI GenBank and the Barcode of Life Data Systems (BOLD) database. This work confirmed the field identifications of winter moth in northwestern Tunisia. Unfortunately, due to the overall lack of genetic diversity of this COI fragment, the native/nonnative status of these populations could not be determined. This low diversity in COI was similarly observed by Gwiazdowski, Elkinton, deWaard, and Sremac (2013) where winter moth samples collected from across Europe were predominantly assigned to two haplotypes that differed by only two substitutions (~0.5% divergence), thereby hampering the ability of COI data alone to determine relationships among populations.

Previously, we examined the use of glacial refugia during the last glacial maximum (LGM) by winter moth (Andersen et al., 2017). In that work, we genotyped field-collected individuals from across Europe with 24 highly polymorphic microsatellite loci. In contrast to mtDNA sequence data that showed limited biogeographic structure between winter moth populations, we found evidence that individuals could be assigned to either an eastern or a western European genetic lineage, with a clear hybrid zone in central Europe (Andersen et al., 2017). Using approximate Bayesian computation (ABC) methods, we also hypothesized that the contemporary genetic diversity of European populations of this species was largely shaped by its retreat to refugia in the Iberian, Balkan, and Caucasus regions, and by hybridization between the Iberian and Balkan lineages postrecolonization.

Absent from the sampling in Andersen et al. (2017), however, were moths from the Italian peninsula and from North Africa, both regions known to have acted as glacial refugia for other species (see Hewitt, 2000; Schmitt, 2007). Therefore, in this study we obtain new winter moth samples from Tunisia, Italy, and additional specimens from Spain, to further explore the biogeographic history of winter moth and its use of potential glacial refugia during the LGM. Specifically, our research objectives were as follows: (a) to genotype winter moth individuals collected in potential glacial refugia not sampled in Andersen et al. (2017; i.e., Southern Italy and North Africa), and (b) to determine whether outbreaking populations of winter moth.
moth in Tunisia represent a native population dating back to the LGM or whether they were recently introduced to the region, and if so from where.

2 | METHODS

2.1 | Data collection

Adult male winter moths were collected using pheromone-baited traps (Elkinton et al., 2010; Elkinton, Lance, Boettner, Khrimian, & Leva, 2011) deployed in Barcelona, Catalonia, Spain, and in Ortì, Calabria, Italy. After moths were collected from the traps, they were placed in glassine envelopes (Uline Corporation, USA), and stored at −80°C. In addition, we sampled larval winter moth collected from trees in Mzara Forest, Tunisia (pure forest of Q. canariensis, alt 653 m.). Once at the laboratory, larvae were placed in 96% ethanol, and then stored at −80°C prior to molecular analyses. See Table 1 for complete locality information. To this dataset, we added genotype scores from individuals presented in Andersen et al. (2017), including all 28 individuals from Tbilisi, Republic of Georgia, 30 randomly selected individuals from Germany, 30 randomly selected individuals from Serbia, and 20 randomly selected individuals from Spain (bringing the total for Spanish samples to 30).

2.2 | DNA extraction and microsatellite amplification

Whole genomic DNA was extracted using the Omega Bio-tek E.Z.N.A.® Tissue DNA Kit (Omega Bio-tek) following the manufacturer’s instructions. For adult males, prior to DNA extraction, the uncus and wings were removed and stored as morphological vouchers. Vouchers were deposited at the Yale Peabody Museum of Natural History, New Haven, Connecticut, USA. Specimen was then homogenized using 3/16” stainless steel beads (GlenMills Inc.) using a FastPrep-24 Sample Homogenizer (MP Biomedicals). Twenty-four polymorphic microsatellite loci were amplified following protocols presented in Havill et al. (2017), and genotyping was performed at the DNA Analysis Facility on Science Hill at Yale University in comparison to the GeneScan 500 LIZ size standard (Thermo Fisher Scientific) using a 3730xl DNA Analyzer (Thermo Fisher Scientific). Fragment lengths were scored using the microsatellite plugin in the software program Geneious v. R11 (https://www.geneious.com). To address potential variation in allele fragment lengths generated during different genotyping runs (see Hoffman & Amos, 2005), the same laboratory methods, personnel, sequencing instrument, and allele bins in Geneious were used for all runs, including those performed by Andersen et al. (2017). Individuals for which ≥20 microsatellite loci were successfully amplified were included in subsequent analyses. This subset included up to 30 randomly selected individuals from

| Country | Location | N | Collector | Date     | Latitude | Longitude |
|---------|----------|---|----------|----------|----------|-----------|
| Georgia | Tbilisi  | 28| G. Japoshvili | 2xii2015 | 41.8068  | 44.7268   |
| Germany | Bühren   | 6 | F. Krueger | 18xii2006 | 51.4658  | 9.6778    |
| Germany | Göttingen| 8 | F. Krueger | 7xii2007  | 51.5460  | 9.9111    |
| Germany | Reinhardshagen | 8 | F. Krueger | 22xii2006 | 51.4763  | 9.5154    |
| Germany | Schlüchtern| 8 | F. Krueger | 20xii2006 | 50.2425  | 9.4538    |
| Italy   | Ortì     | 26| E. Castiglione and F. Manti | 11iii2018 | 38.1475  | 15.7196   |
| Serbia  | Belgrade | 9 | M. Glavendekić | 27xii2006 | 44.7643  | 20.4364   |
| Serbia  | Pančevo | 11| M. Glavendekić | 4xii2006  | 44.8524  | 20.6532   |
| Serbia  | Pančevo | 10| M. Glavendekić | 28xii2006 | 44.8524  | 20.6532   |
| Spain   | Barcelona| 10| J. Bau     | 12ix2018  | 41.9657  | 2.1907    |
| Spain   | La Langa | 2 | A. Pepi   | 6xii2015  | 40.0895  | 2.6607    |
| Spain   | Uriz     | 1 | M. Lombardero | xii2008  | 43.1184  | 7.6517    |
| Spain   | Uriz     | 1 | M. Lombardero | xii2008  | 43.1184  | 7.6517    |
| Spain   | Lugo     | 1 | M. Lombardero | xii2008  | 42.9922  | 7.5451    |
| Spain   | Uriz     | 1 | M. Lombardero | xii2008  | 43.1190  | 7.6497    |
| Spain   | Uriz     | 1 | M. Lombardero | xii2008  | 43.1190  | 7.6497    |
| Spain   | Uriz     | 1 | M. Lombardero | xii2008  | 43.1190  | 7.6497    |
| Spain   | Uriz     | 1 | M. Lombardero | xii2008  | 43.1190  | 7.6497    |
| Spain   | Lugo     | 1 | M. Lombardero | xii2008  | 42.9922  | 7.5451    |
| Spain   | Lugo     | 8 | M. Lombardero | 9i2014    | 42.9926  | 7.5441    |
| Tunisia | Mzara Forest | 17| O. Ezzine | 27iv2018 | 36.7690  | 8.7200    |
each of the following locations (a) Spain, (b) Serbia, (c) Republic of Georgia, and (d) Germany. Locality collection information for each individual in the analyses is provided in Table 1, and microsatellite genotype information is as a tab-delimited supplemental file titled “WinterMothTunisiaStructure.txt.”

### 2.3 Bayesian genetic clustering

The assignment probabilities (Q) of each individual to distinct genetic clusters (K) were estimated using the Bayesian genetic clustering software STRUCTURE v.2.3.2 (Falush, Stephens, & Pritchard, 2003; Pritchard, Stephens, & Donnelly, 2000) for values ranging from K = 1 to K = 8. For each value of K, ten independent analysis were run, each with random starting values, a burn-in period of 100,000 generations, and a total runtime of 1,000,000 generations. For each run, we utilized the admixture model, with correlated allele frequencies using default settings for both. The optimal number of clusters was then estimated using the Delta-K method of Evanno, Regnaut, and Goudet (2005) as implemented in the software package STRUCTURE HARVEST (Earl & vonHoldt, 2012), and values for Q were averaged across runs using CLUMPAK (Kopelman, Mayzel, Jakobsson, Rosenberg, & Mayrose, 2015).

### 2.4 Genetic diversity and differentiation

For individuals from each country, we calculated several population genetic statistics, including the number of individuals genotyped (n), the average number of alleles per locus (Na), the effective number of alleles (Eff_Na), the average observed heterozygosity across loci (H_o), the average heterozygosity within each population (H_s), the total heterozygosity (H_t), the average inbreeding coefficient (G_{is}), and the presence of deviation from Hardy–Weinberg Equilibrium using GENODIVE v.2.0b27 (Meirmans & van Tienderen, 2004). The proportion of null alleles present in the dataset and “excluding null alleles” (ENA)-corrected measures of genetic differentiation (F_{ST}) among populations from each country was estimated were

---

**Figure 1** Proportional assignment (Q) of individuals to genetic clusters (K) based on summary across independent STRUCTURE runs using CLUMPAK.
estimated using FReeNA (Chapuis & Estoup, 2007), the subsequent pairwise corrected $F_{ST}$ distance matrix was then used to calculate a "NeighborNet" network with the program SPliTTree v.4.14.2 (Huson & Bryant, 2006). The probability of population differentiation was then estimated using GENEPOP (Raymond & Rousset, 1995; Rousset, 2008).
TABLE 3  Demographic parameter priors for DnABC simulations

| Parameters                        | Prior               |
|----------------------------------|---------------------|
|                                  | Min     | Max     |
| Population sizes (effective pop. size) |         |         |
| Tbilisi, Georgia                 | 10,000   | 500,000 |
| Germany                          | 10,000   | 1,000,000 |
| Ortì, Italy                      | 10,000   | 500,000 |
| Serbia                           | 10,000   | 1,000,000 |
| Spain                            | 10,000   | 500,000 |
| Mzara Forest, Tunisia            | 10,000   | 500,000 |
| Ancestor 1 (NE3a)                | 10,000   | 100,000 |
| Ancestor 2 (NE3b)                | 10,000   | 100,000 |
| Ancestor 3 (NE1)                 | 10,000   | 100,000 |
| Divergence/merger time (years)   |         |         |
| tBottleneck                      | 10       | 200     |
| tRecentIntro                     | 1,000    | 10,000  |
| t1                               | 1,000    | 15,000  |
| tTunisia                         | 5,000    | 20,000  |
| t2                               | 10,000   | 30,000  |
| t3                               | 20,000   | 40,000  |
| t4                               | 20,000   | 50,000  |
| Admixture proportion (from the Spanish Lineage) |         |         |
| ra                               | 0.001    | 0.999   |
| Mutation rates (per generation)  |         |         |
| Mean µmic_1                      | $1 \times 10^{-5}$| $1 \times 10^{-4}$ |
| Mean pmic_1                      | 0.1      | 1.0     |
| Mean snimic_1                    | $1 \times 10^{-6}$| $1 \times 10^{-5}$ |

2.5  | Historical demography

To estimate the divergence time between the Tunisian population and Eurasian populations of winter moth using ABC as implemented in the software DnABC v.2.1.0 (Cornuet et al., 2014, 2008), we used a two-step approach to limit the number of potential scenarios (see Stone et al., 2017 for an example). In the first step, we first estimated the relationship of the southern Italian population to other European populations (i.e., Georgian, German, Serbian, and Spanish). We used the results from Andersen et al. (2017) to establish the relationships among the other European populations. Andersen et al. (2017) found strong support for the German lineage to be the result of admixture between the Spanish and Serbian lineages after the LGM. The authors were unable, however, to conclude whether the Serbian and Georgian lineages diverged before or after the divergence of the Spanish lineage, as these two scenarios received equivocal results. To reduce complexity in scenario design, here we use a tree with the Spanish lineage diverging prior to the divergence of the Serbian and Georgian lineages [Scenario 7 from Andersen et al. (2017)] as our root tree. Three scenarios (Figure S1) were compared to determine whether the Italian population was most closely related to the Spanish, Serbian, or Georgian populations by generating a reference table based on three million simulated datasets (one million per scenario).

The branching pattern among European populations from the scenario receiving the highest support from the first step was then used to explore the origins of the Tunisian population. Nine scenarios were explored (Figure S2); four scenarios for the Tunisian population diverging from the Spanish, southern Italian, Serbian, and Georgian populations with branching times corresponding with the LGM, and five scenarios for the Tunisian population diverging from the Spanish, southern Italian, Serbian, and Georgian as well as from the admixed German population with a branching time corresponding to contemporary establishment (i.e., within the last ~1,000 years). The contemporary scenarios also included an additional population size parameter to allow for changes in population size following the divergence from potential source populations (i.e., to allow the possibility of a genetic bottleneck). Initially, three additional scenarios were explored (not shown) where the Tunisian population acted as a source for the Spanish and Italian populations; however, these scenarios received posterior probabilities of 0.0000 in comparison to a European origin of the Tunisian population (posterior probability of 1.0000), and such “out-of-Africa” scenarios were not pursued further.

All scenarios for both steps included the following: (a) multiple population size parameters to allow for changes in population sizes following splitting/merging events; (b) default mutation model parameters, except that the minimum mean mutation rate to $1 \times 10^{-5}$, and maximum values for the mean and individual locus coefficient $P$s were both set to 1.0; and (c) four loci with especially large allelic ranges (2,339, 925, 2,191, and 12,042) were removed to improve the shape of the cloud of simulated datasets. For both sets of simulations, the following summary statistics were used: one sample; mean number of alleles, mean genic diversity, mean size variance: two sample; $F_{ST}$, classification index, and $(d\mu)^2$ distance. Distributions for parameter priors are presented in Table 3. The scenarios were then compared using and the “Logistic Regression” tests as part of the “Compute posterior probabilities of scenarios” analysis in DnABC based on comparisons of 1% of simulated datasets closest to the observed data (Cornuet, Ravigné, & Estoup, 2010; Cornuet et al., 2008). To determine the goodness of fit of simulated scenarios, we first conducted a principle components analysis (PCA) that examined the posterior distributions of summary statistics in comparison to the observed dataset, as implemented in the “Perform model checking” analysis in DnABC. We then estimated error rates by implementing the “Evaluate confidence in scenario choice” analysis in DnABC.

3  | RESULTS

3.1  | Data collection, DNA extraction, and microsatellite amplification

After filtering individuals from which less than 20 of 24 polymorphic microsatellites were amplified, ten winter moth individuals
collected in Barcelona, Spain, 26 individuals from Ortì, Italy, and 17 individuals from Mzara Forest, Tunisia were included in subsequent analyses.

3.2 | Bayesian genetic clustering

The probability of assignment (Q) of individuals to major genetic clusters (K) for K = 1 through K = 8 after averaging across independent Structure runs is presented in Figure 1. Assignments for major and minor clusters, as well as the number of runs supporting each cluster, after summary in Clumpak are presented in Figure S3. Results from StructureHarvester indicated that the optimal number of genetic clusters (K) present in the dataset was K = 6 (Figure S4). At K = 6, all ten independent Structure runs were assigned to one major cluster using Clumpak, and these genetic clusters predominantly corresponded to the geographical origin of samples (Figures 1 and 2).

3.3 | Genetic diversity and differentiation

Population genetic statistics for each population are presented in Table 2. The German population displayed the highest number of effective alleles (Eff_Na = 4.987) as well as the highest level of observed heterozygosity (H_o = 0.670) while the Mzara Forest, Tunisia population had the lowest number of effective alleles (Eff_Na = 2.817) and the lowest level of observed heterozygosity (H_o = 0.450). As per Andersen et al. (2017), all sampled localities displayed highly significant deviations from Hardy–Weinberg Equilibrium (p = .001 for all). The average value ± standard error (SE) for the null allele frequency was 0.053 ± 0.024. The average ENA-corrected measure of

![Figure 3: Schematic representation of FST distances between countries visualized as a NeighborNet in SplitsTree](image)

![Figure 4: Evolutionary scenario of European and North African populations of winter moth based on simulations conducted in DIYABC. Lines widths are drawn relatively proportional to the mean estimated population size. At each node, the mean divergence/merger time is shown along the y-axis (1 ka = 1,000 years). Ninety-five percent confidence intervals for population sizes, divergence/merger times, and for the amount of admixture (ra) are presented in Table 3](image)
ANDERSEN ET AL.

The NeighborNet analyses indicated that all sampled populations showed a high degree of genetic distinction, with all but the German population being placed at the end of a long branch (Figure 3).

3.4 | Historical demography

Results from simulations conducted during the first step in our DnABC analyses indicated that the Ortì, Italy population was most closely related to the Spanish population (Figure S5 and Table S2). Model-checking analyses for the posterior distributions for summary statistics based on this dataset formed a distinct cloud around the observed dataset (Figure S8). Evaluations of the confidence in the scenario choice indicated a posterior predictive error rate of 0.287. Based on this scenario (Figure 4), the Tunisian population likely diverged from the Spanish population ~16,200 years ago (95% CI = 10,100–19,800 years ago; Table 3). All estimates for demographic parameters for the most likely scenarios were calculated based on the analysis of 1,000,000 simulated datasets and are presented in Figures S9 and S10, for the scenarios estimating the relationship of the southern Italian population and the Tunisian populations, respectively, and summarized for the Tunisian scenario in Table 4.

4 | DISCUSSION

We previously demonstrated that winter moth persisted in at least three glacial refugia (Iberian, Aegean, and Caucasus) during the LGM (Andersen et al., 2017). By analyzing newly sampled populations in North Africa and the Italian Peninsula, we provide evidence for two additional LGM refugia. Contemporary populations sampled in Tunisia and southern Italy differ from all other European populations (Figures 1 and 2) suggesting that the lineages that derived from these refugia did not recolonize Europe following the retreat of glaciers following the LGM.
ANDERSEN ET AL.

(Figure 4). For the southern Italian lineage, it is possible that the Alps presented a significant geographic barrier to their recolonization as has been seen in other systems (see Hewitt, 2000). However, as we do not have samples from central or northern Italy included in this analysis, we can only speculate in regards to barriers to recolonization for this lineage. In regard to the Tunisian lineage, it is very likely that after crossing the Strait of Gibraltar and colonizing northern Africa during the LGM that populations of winter moth became genetically isolated as deciduous forests became geographically isolated in this region with the Mediterranean Sea preventing gene flow with European populations. Again, we can only speculate as to the role of the Mediterranean Sea as a barrier without sampling across northern Africa, southern Spain, and from Mediterranean islands (such as Sicily and Sardinia). However, when our results (i.e., locally distinct populations based on nuclear loci) are combined with mitochondrial studies that have uncovered limited genetic diversity among winter moth populations (Gwiazdowski et al., 2013; Mannai et al., 2017), it suggests that winter moth may be a relatively new addition to the European ecological community, and either became established in Europe prior to the onset of the last glacial cycle, or speciated in Europe shortly before this event.

Our results provide evidence that the winter moth population sampled in northwestern Tunisia arrived in North Africa during the LGM. Therefore, factors other than a recent biological invasion are likely responsible for the current outbreak. These factors might include reduced interspecific competition, particularly from other oak defoliators such as *Lymantria dispar* L. (Lepidoptera: Erebidae). *Lymantria dispar* has been reported to outbreak densities in Tunisian oak forests (with four large outbreaks reported between 1925 and 1995; Ben Jemâa, Mnara, Villemant, & Khaldi, 2002); however, no outbreaks of this species have been observed in the region since 2000 (M’nara, 2005). It is therefore possible that the absence of *L. dispar* outbreaks has allowed other defoliators, such as *Tortrix viridana* L. (Lepidoptera: Tortricidae), *Eraniss defoliaria* (Clerck) (Lepidoptera: Geometridae), and possibly winter moth to grow to outbreak densities (Mannai, 2017). In addition, climate change in northern Africa might have influenced the population dynamics of winter moth, as already demonstrated in northern Scandinavia (Jepsen, Hagen, Ims, & Yoccoz, 2008). Indeed, northern Tunisia saw dramatic changes in mean annual temperature during the 20th century, with an increase of 2.5–3°C on average, and future climate models predicting an additional increase of 3–4°C coupled with decreased rainfall in the 21st century (Hulme, Doherty, Ngora, New, & Lister, 2001). Environmental factors can play an important role in shaping plant-herbivore interactions (Kuglerová, Skuhrovec, & Münzbergová, 2019). It is therefore possible that changes in temperature and/or precipitation might promote winter moth populations either directly by reducing developmental times and/or indirectly by limiting the amount of secondary compounds produced by stressed oak trees. It is also possible that human-mediated disturbances in the region might be limiting the impact of native natural enemies (such as pathogens, parasitoids, and predators). Further work in this region is needed to understand what factors are resulting in the outbreaks of winter moth in order to preserve these culturally and ecologically important native oak stands.

5 | CONCLUSIONS

Here, we find the use of two additional glacial refugia, North Africa and southern Italy, by winter moth during the LGM. Our results indicate that populations that utilized both refugia during the LGM diverged from a population that migrated into the Iberian Peninsula, and that subsequently each population became genetically distinct. Curiously, our findings continue to support the results from Andersen et al. (2017) that southern populations appear to be less genetically distinct than northern populations. As such, the biogeographical history of winter moth might represent an important case study for the role of secondary contact in promoting the genetic diversity of outbreaking pest species. Finally, we hope that this new genetic data, and the discovery of additional biogeographic structuring, can be used in conjunction with the recently published winter moth genome (Derks et al., 2015), to promote the use of winter moth as a model system for future research questions, including studies of local adaptation—a subject for which winter moth has previously been studied in the pregenomics era (Buse & Good, 1996; van Dornen, Backeljau, Matthysen, & Dhondt, 1997; Holliday, 1977; Visser & Holleman, 2001).

ACKNOWLEDGMENTS

This work could not have been accomplished without the tremendous efforts of numerous winter moth collectors. In particular, we would like to thank Josep Bau for collecting samples in Barcelona, Catalonia, Spain, and Carmelo Bonsignore for organizing the collection of samples in Ortì, Calabria, Italy. A complete list of all collectors is presented in Table 1, and we are extremely grateful for their contributions. We would also like to thank Brian Griffin and DeAdra Newman for technical assistance, and Jeff Boettner, Hannah Broadley, Rodger Gwiazdowski, and Michelle Labbé for their comments on earlier drafts of this manuscript. Funding was provided by the USDA Forest Service, Northern Research Station; USDA Forest Service (agreement number 17-CR-11242303-066 to AC); and USDA APHIS (grant number AP17PPQS&T00C068 awarded to JSE); and by INRGREF.

CONFLICT OF INTEREST

None declared.

AUTHOR CONTRIBUTIONS

All authors contributed equally to the study design and manuscript preparation. YM, OE, SD, and MLBJ oversaw sample collection. JCA would like to thank Josep Bau for collecting samples in Barcelona, Catalonia, Spain, and Carmelo Bonsignore for organizing the collection of samples in Ortì, Calabria, Italy. A complete list of all collectors is presented in Table 1, and we are extremely grateful for their contributions. We would also like to thank Brian Griffin and DeAdra Newman for technical assistance, and Jeff Boettner, Hannah Broadley, Rodger Gwiazdowski, and Michelle Labbé for their comments on earlier drafts of this manuscript. Funding was provided by the USDA Forest Service, Northern Research Station; USDA Forest Service (agreement number 17-CR-11242303-066 to AC); and USDA APHIS (grant number AP17PPQS&T00C068 awarded to JSE); and by INRGREF.

ORCID

Jeremy C. Andersen https://orcid.org/0000-0002-9273-6490
Nathan P. Havill https://orcid.org/0000-0002-4004-8266
Yaussa Mannai https://orcid.org/0000-0001-8972-7537
Olfa Ezzine https://orcid.org/0000-0003-0859-4299
Samir Dhahri https://orcid.org/0000-0002-4954-8670
DATA AVAILABILITY STATEMENT
Genotype data for this study is provided as a tab-delimited file titled "WinterMothTunisiaStructure.txt" uploaded as supplemental material.

REFERENCES
Andersen, J. C., Havill, N. P., Caccone, A., & Elkinton, J. S. (2017). Postglacial recolonization shaped the genetic diversity of the winter moth (Operophtera brumata) in Europe. Ecology and Evolution, 7, 3312–3322.

Ben Jemâa, M. L., Mnara, S., Villelmain, C., & Khalidi, A. (2002). Lymantria dispar L. (Lepidoptera, Lymantriidae) en Tunisie: état actuel des connaissances et perspectives de recherche. Bulletin De Protection Intégrée Des Forêts À Quercus Ssp. 25, 101–108.

Boudy, P. (1950). Economie Forestière Nord Africaine. Tome II. Monographie et Traitement des Essences Forestières. Paris, France: Éditions Larose.

Buse, A., & Good, J. E. G. (1996). Synchronization of larval emergence in winter moth (Operophtera brumata L.) and budburst in pedunculate oak (Quercus robur L.) under simulated climate change. Ecological Entomology, 21, 335–343.

Chapuis, M.-P., & Estoup, A. (2007). Microsatellite null alleles and estimation of population differentiation. Molecular Biology and Evolution, 24, 621–631. https://doi.org/10.1093/molbev/msi191

Cornuet, J.-M., Pudlo, P., Veyssier, J., Deheu-Garcia, A., Gautier, M., Leblois, R., ... Estoup, A. (2014). DIYABC v2.0: A software to make approximate Bayesian computation inferences about population history using single nucleotide polymorphism, DNA sequence and microsatellite data. Bioinformatics, 30, 1187–1189.

Cornuet, J.-M., Ragnirev, V., & Estoup, A. (2010). Inference on population history and model checking using DNA sequence and microsatellite data with the software DIYABC (v1.0). BMC Bioinformatics, 11, 401.

Cornuet, J.-M., Santos, F., Beaumont, M. A., Robert, C. P., Marin, J.-M., Balding, D. J., ... Estoup, A. (2008). Inferring population history with DIY ABC: A user-friendly approach to approximate Bayesian computation. Bioinformatics, 24, 2713–2719.

Debazeau, E. F. (1959). Le végétation forestière de la Kroumirie. Nancy, France: Station de Recherches et Expériences Forestières.

Derks, M. F. L., Smit, S., Salis, L., Schijlen, E., Bossers, A., Mateman, C., ... Megens, H.-J. (2015). The genome of winter moth (Operophtera brumata) provides a genomic perspective on sexual dimorphism and phenology. Genome Biology and Evolution, 7, 2321–2332.

Direction Générale des Forêts (2005). Inventaire Forestier National en Tunisie. Tunis, Tunisia: Ministère de l’Agriculture.

Earl, D. A., & vonHoldt, B. M. (2012). STRUCTURE HARVESTER: A website and program for visualizing STRUCTURE output and implementing the Evanno method. Conservation Genetics Resources, 4, 359–361.

Elkinton, J., Boettner, G., Liebhold, A., & Gwiazdowski, R. (2015). Biology, spread, and biological control of winter moth in the eastern United States. Morgantown, WV: USDA Forest Service Publication.

Elkinton, J. S., Boettner, G. H., Sremac, M., Gwiazdowski, R., Hunkins, R. R., Callahan, J., ... Campbell, N. K. (2010). Survey for winter moth (Lepidoptera: Geometridae) in northeastern North America with pheromone-baited traps and hybridization with the native Bruce spanworm (Lepidoptera: Geometridae). Annals of the Entomological Society of America, 103, 135–145.

Elkinton, J. S., Lance, D., Boettner, G., Khrimian, A., & Leva, N. (2011). Evaluation of pheromone-baited traps for winter moth and Bruce spanworm (Lepidoptera: Geometridae). Journal of Economic Entomology, 104, 494–500.

Evanno, G., Regnaut, S., & Goudet, J. (2005). Detecting the number of clusters of individuals using the software STRUCTURE: A simulated study. Molecular Ecology, 14, 2611–2620.

Falush, D., Stephens, M., & Pritchard, J. K. (2003). Inference of population structure using multilocus genotype data: Linked loci and correlated allele frequencies. Genetics, 164, 1567–1587.

Ferguson, D. C. (1978). Pests not known to occur in the United States or of limited distribution. Winter moth Operopthera brumata (L.) Lepidoptera: Geometridae. Cooperative Plant Pest Report, 3, 687–694.

Gillespie, D. R., Finlayson, T., Tonks, N. V., & Ross, D. A. (1978). Occurrence of winter moth, Operopthera brumata (Lepidoptera, Geometridae), on southern Vancouver-Island, British-Columbia. Canadian Entomologist, 110, 223–224.

Gwiazdowski, R. A., Elkinton, J. S., deWaard, J. R., & Sremac, M. (2013). Phylogeographic diversity of the winter moths Operopthera brumata and O. bruceata (Lepidoptera: Geometridae) in Europe and North America. Annals of the Entomological Society of America, 106, 143–151.

Habel, J. C., Lens, L., Rodder, D., & Schmitt, T. (2011). From Africa to Europe and back: Refugia and range shifts cause high genetic differentiation in the Marbled White butterfly Melanargia galathea. BMC Evolutionary Biology, 11, 215.

Hausmann, A., & Viidalepp, J. (2012). The geometrid moths of Europe (Vol. 3). Vester Skerninge, Denmark: Apollo Books.

Havill, N. P., Elkinton, J. S., Andersen, J. C., Hagen, S. B., Broadley, H. J., Boettner, G. H., & Caccone, A. (2017). Asymmetric hybridization between nonnative winter moth, Operopthera brumata (Lepidoptera: Geometridae), and native Bruce spanworm, O. bruceata, in the northeastern United States, assessed with novel microsatellites and SNPs. Bulletin of Entomological Research, 107, 241–250.

Hewitt, G. M. (1996). Some genetic consequences of ice ages, and their role in divergence and speciation. Biological Journal of the Linnean Society, 58, 247–276.

Hewitt, G. (2000). The genetic legacy of the Quaternary ice ages. Nature, 405, 907–913.

Hoffman, J. I., & Amos, W. (2005). Microsatellite genotyping errors: Detection approaches, common sources and consequences for paternial exclusion. Molecular Ecology, 14, 599–612. https://doi.org/10.1111/j.1365-294X.2004.02419.x

Holliday, N. J. (1977). Population ecology of winter moth (Operopthera brumata) on apple in relation to larval dispersal and time of bud burst. Journal of Applied Ecology, 14, 803–813.

Huck, S., Buedel, B., & Schmitt, T. (2012). Ice-age isolation, postglacial hybridization and recent population bottlenecks shape the genetic structure of Meum athamanticum in Central Europe. Flora, 207, 399–407.

Hulme, M., Doherty, R., Ngara, T., New, M., & Lister, D. (2001). African climate change: 1900–2100. Climate Research, 17, 145–168.

Huson, D. H., & Bryant, D. (2006). Application of phylogenetic networks in evolutionary studies. Molecular Biology and Evolution, 23, 254–267.
Jepsen, J. U., Hagen, S. B., Ims, R. A., & Yoccoz, N. G. (2008). Climate change and outbreaks of the geometrids Operophtera brumata and Epiprita autumnata in subarctic birch forest: Evidence of a recent outbreak range expansion. Journal of Animal Ecology, 77, 257–264.

Kimberling, D. N., Miller, J. C., & Penrose, R. L. (1986). Distribution and parasitism of winter moth, Operophtera-brumata (Lepidoptera, Geometridae), in western Oregon. Environmental Entomology, 15, 1042–1046.

Kopelman, N. M., Mayzel, J., Jakobsson, M., Rosenberg, N. A., & Mayrose, I. (2015). CLUMPKA: A program for identifying clustering modes and packaging population structure inferences across K. Molecular Ecology Resources, 15, 1179–1191.

Kuglerová, M., Skuhrovec, J., & Münzbergová, Z. (2019). Relative importance of drought, soil quality, and plant species in determining the strength of plant-herbivore interactions. Environmental Entomology, 44, 665–677.

MacPhee, A. W. (1967). Winter moth Operophtera brumata (Lepidoptera - Geometridae) a new pest attacking apple orchards in Nova Scotia and its coldhardiness. Canadian Entomologist, 99, 829–834.

Mannai, Y. (2017). Etude des Lépidoptères défoliateurs des chênes au Nord-Ouest de la Tunisie: Biodiversité et interactions insectes/plantes-hôtes (p. 271). Tunis, Tunisia: Faculté des Sciences Mathématiques, Physiques et Naturelles de Tunis. Université El Manar.

Mannai, Y., Ezzine, O., Hausmann, A., Nouira, S., & Ben Jamâa, M. L. (2017). Budburst phenology and host use by Operophtera brumata (Linnaeus, 1758) (Lepidoptera: Geometridae) in three Mediterranean oak species. Annals of Forest Science, 74, 3. https://doi.org/10.1007/s13595-016-0600-3

Mannai, Y., Ezzine, O., Nouira, S., & Ben Jamâa, M. L. (2015). First report of the winter moth Operophtera brumata on Quercus canariensis and Q. afares in North West of Tunisia. Tunisian Journal of Plant Protection, 10, 69–73.

Meirmans, P. G., & van Tienderen, P. H. (2004). GENOTYPE and GENODIVE: Two programs for the analysis of genetic diversity of plant-herbivore interactions. Molecular Ecology Notes, 4, 792–794.

M’Mara, S., Ben Jamâa, M. L., & Nouira, S. (2005). Bilan des observations sur Lymantria dispar (L.) en phase de latence en Tunisie. Bulletin De Protection Intégrée Des Forêts À Quercus Ssp, 28, 147–154.

Moyal, P., Tokro, P., Bayram, A., Savopoulou-Soulmani, M., Conti, E., Eizaguirre, M., ... Andreidis, S. (2011). Origin and taxonomic status of the Palearctic population of the stem borer Synergus umbraculus (Lepidoptera, Geometridae). Journal of Zoological Systematics and Evolutionary Research, 49, 945–959.

Raymond, M., & Rousset, F. (1995). GENEPop (VERSION-1.2) - Population-genetics software for exact tests and ecumenism. Journal of Heredity, 86, 248–249.

Rofes, J., Cucchi, T., Hanot, P., Herman, J., Stephan, P., Cersoy, S., ... Tresset, A. (2018). Postglacial recolonization and Holocene diversification of Crocidura suaveolens (Mammalia, Soricidae) on the north-western fringe of the European continent. Quaternary Science Reviews, 190, 1–10.

Rousset, F. (2008). GENEPop007: A complete re-implementation of the GENEPop software for Windows and Linux. Molecular Ecology Resources, 8, 103–106.

Schmitt, T. (2007). Molecular biogeography of Europe: Pleistocene cycles and postglacial trends. Frontiers in Zoology, 4, 1–13.

Schmitt, T., & Müller, P. (2007). Limited hybridization along a large contact zone between two genetic lineages of the butterfly Erebia medusa (Satyrinae, Lepidoptera) in Central Europe. Journal of Zoological Systematics and Evolutionary Research, 45, 39–46.

Schmitt, T., Rober, S., & Seitz, A. (2005). Is the last glaciation the only relevant event for the present genetic population structure of the meadow brown butterfly Maniola jurtina (Lepidoptera: Nymphalidae)? Biological Journal of the Linnean Society, 85, 419–431. https://doi.org/10.1111/j.1095-8312.2005.00504.x

Schmitt, T., & Seitz, A. (2001). Allozyme variation in Polymatryrus coronid (Lepidoptera: Lycaenidae): Identification of ice-age refugia and reconstruction of post-glacial expansion. Journal of Biogeography, 28, 1129–1136.

Sim, Z., Hall, J. C., Jex, B., Hegel, T. M., & Coltman, D. W. (2016). Genome-wide set of SNPs reveals evidence for two glacial refugia and admixture from postglacial recolonization in an alpine ungluate. Molecular Ecology, 25, 3696–3705.

Song, W., Cao, L.-J., Li, B.-Y., Gong, Y.-J., Hoffman, A. A., & Wei, S.-J. (2018). Multiple refugia from penultimate glaciations in East Asia demonstrated by phylogeography and ecological modelling of an insect pest. BMC Evolutionary Biology, 18, 152.

Stolting, K. N., Paris, M., Meier, C., Heinze, B., Castiglione, S., Bartha, D., & Lexer, C. (2015). Genome-wide patterns of differentiation and spatially varying selection between postglacial recolonization lineages of Populus alba (Salicaceae), a widespread forest tree. New Phytologist, 207, 723–734.

Stone, G. N., White, S. C., Csóka, G., Meliga, K., Mutun, S., Pénzes, Z., ... Nicholls, J. A. (2017). Tournament ABC analysis of the western Palearctic population history of an oak gall wasp, Synergus umbraculus. Molecular Ecology, 26, 6685–6703.

Torroni, A., Bandelt, H. J., Macaulay, V., Richards, M., Cruciani, F., Rengo, C., ... Scozzari, R. (2001). A signal, from human mtDNA, of postglacial recolonization in Europe. American Journal of Human Genetics, 69, 844–852.

van Dongen, S., Backeljau, T., Matthysen, E., & Dhondt, A. A. (1997). Population-genetics software for exact tests and ecumenicism. Molecular Ecology, 6, 369–374.

von Reumont, B. M., Struwe, J.-F., Schwarzer, J., & Misof, B. (2012). Phylogeography of the burnet moth Operophtera brumata (Lepidoptera, Geometridae), in western Oregon. Environmental Entomology, 41, 163–173.

Wint, W. (1983). The role of alternative host-plant species in the life of a polyphagous moth, Operophtera-brumata (Lepidoptera, Geometridae). Journal of Animal Ecology, 52, 439–450. https://doi.org/10.2307/456479

Zhu, G.-P., Ye, Z., Du, J., Zhang, D.-L., Zhen, Y.-H., Zheng, C.-G., ... Bu, W.-J. (2016). Range wide molecular data and niche modeling revealed the Pleistocene history of a global invader (Halymorpha halys). Scientific Reports, 6, 23192.

SUPPORTING INFORMATION
Additional supporting information may be found online in the Supporting Information section.

How to cite this article: Andersen JC, Havill NP, Mannai Y, et al. Identification of winter moth (Operophtera brumata) refugia in North Africa and the Italian Peninsula during the last glacial maximum. Ecol Evol. 2019:9:13931–13941. https://doi.org/10.1002/ece3.5830