- 1 2 DR. QIXIN HE (Orcid ID: 0000-0003-1696-8203) 3 4 5 Article type : Original Article 6 7 Title: Inferring the geographic origin of a range expansion: latitudinal and longitudinal 8 9 coordinates inferred from genomic data in an ABC framework with the program X-ORIGIN 10 Qixin He<sup>1,2</sup>, Joyce R. Prado<sup>3</sup>, and L. Lacey Knowles<sup>4</sup> 11 12 <sup>2</sup>Department of Ecology and Evolutionary Biology, University of Chicago, Chicago IL, USA 13 60637 14 <sup>3</sup>Departamento de Ciências Biológicas, Escola Superior de Agricultura 'Luiz de Queiroz', 15 Universidade de São Paulo, Piracicaba, Brazil 16 <sup>4</sup>Department of Ecology and Evolutionary Biology, University of Michigan, Ann Arbor MI, 17 USA 41809-1079 18 19 <sup>1</sup>Corresponding Author: Oixin He<sup>1</sup>, E-mail: hegixin@uchicago.edu 20 Orcid id: 0000-0003-1696-8203 21 22 RH: Inferring the origin of range expansion 23 **Contact Information:** 24 Qixin He, heqixin@uchicago.edu 25 26 University of Chicago
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This is the author manuscript accepted for publication and has undergone full peer review but has not been through the copyediting, typesetting, pagination and proofreading process, which may lead to differences between this version and the <u>Version of Record</u>. Please cite this article as <u>doi:</u> <u>10.1111/mec.14380</u>

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31 Word count: Main Text 6613

32 Number of Figures: 5 + 3 SI

33 Number of Tables: 1 + 2 SI

Data accessibility: vcf files for Collared pika and SNP used in the analysis were deposited on Dryad for data archive (DOI: 10.5061/dryad.4s1gg); X-Origin pipeline tutorial, scripts, example files and input files used in the study are uploaded on GitHub and released under the DOI: <u>https://zenodo.org/badge/latestdoi/100994225</u>.

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39 Abstract

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Climatic or environmental change is not only driving distributional shifts in species today, but it 41 has also caused distributions to expand and contract in the past. Inferences about the geographic 42 43 locations of past populations, especially regions that served as refugia (i.e., source populations) and migratory routes are a challenging endeavor. Refugial areas may be evidenced from fossil 44 45 records or regions of temporal stability inferred from ecological niche models. Genomic data offer an alternative and broadly applicable source of information about the locality of refugial 46 47 areas, especially relative to fossil data, which are either unavailable or incomplete for most species. Here we present a pipeline we developed (called X-ORIGIN) for statistically inferring the 48 49 geographic origin of range expansion using a spatially explicit coalescent model and an Approximate Bayesian Computation testing framework. In addition to assessing the probability 50 51 of specific latitudinal and longitudinal coordinates of refugial or source populations, such inferences can also be made accounting for the effects of temporal and spatial environmental 52 heterogeneity, which may impact migration routes. We demonstrate X-ORIGIN with an analysis of 53 genomic data collected in the Collared pika that underwent post-glacial expansion across Alaska, 54 as well as present an assessment of its accuracy under a known model of expansion to validate 55 56 the approach.

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Population expansions leave signatures in the distribution of population genetic variation 61 across a landscape. This pattern of genetic variation is commonly used for making inferences 62 about the underlying demographic processes. For example, the decreasing pattern of genetic 63 diversity along expansion routes has been used to infer the origin of human migrations 64 (DeGiorgio, Jakobsson, & Rosenberg, 2009; Ramachandran et al., 2005). Similarly, such genetic 65 signatures have been applied to study post-glacial expansions in other species, as well as their 66 corresponding geographic refugia during glacial periods of the Pleistocene (reviewed in Hewitt, 67 2000). 68

However, this approach comes with an inherent issue. Specifically, genetic diversity 69 70 patterns (e.g., heterozygosity,  $F_{ST}$ ) can reflect not only signatures from recent distributional shifts, but also local habitat suitability or long-term geographic isolation (Austerlitz, Jung-71 72 Muller, Godelle, & Gouyon, 1997; Ray, Currat, & Excoffier, 2003). Thus, while the isolation-73 by-distance model applies relatively well to species that have a broad habitat, such as human 74 beings, species with narrower niches tend to track their habitats, displaying a genetic diversity pattern of isolation-by-barriers or resistance (McRae & Beier, 2007). Therefore, sole reliance on 75 76 the gradients of population size/heterozygosity or the principal components without spatial models is inadequate for making accurate inferences about the ancestral source population or 77 78 directions of expansion (François et al., 2010). Due to the rich, yet confounding information 79 retained in the genetic diversity patterns, most phylogeographic studies infer the location of 80 hypothesized refugia from the data that are independent of the genomic information (reviewed in Knowles, 2009). Ecological niche models (ENMs), for instance, could be applied to infer areas 81 82 with temporal stability as suitable habitats. In addition, the associated genetic data could then be used to evaluate the hypothesis that such geographic regions would have served as refugial 83 source population (e.g., see Carnaval, Hickerson, Haddad, Rodrigues, & Moritz, 2009; Knowles, 84 Massatti, He, Olson, & Lanier, 2016). 85

Attempts to address the issue of complex historical processes shaping the current genetic patterns have witnessed the development of spatially-explicit demographic models as well as spatial genetic indices. Ray, Currat, Berthier, & Excoffier (2005) systematically tested the likelihood of different geographic locations as human origins by evaluating the goodness-of-fit 90 of  $R_{ST}$  values from different spatial simulations of expansions using the empirical values. Itan, 91 Powell, Beaumont, Burger, & Thomas (2009) estimated the origin of lactase persistent mutations 92 in Europe by fitting empirical frequencies of lactase persistent mutations to those from spatial 93 simulations of the gene expansion along with dairy groups. These pioneer studies demonstrate 94 the potential of using spatially-explicit models for estimating migration histories. However, these 95 models do not take temporal changes in habitat suitability into account, which limit their 96 applicability in flora and fauna that underwent expansions largely driven by climatic oscillations.

Spatial genetic indices, on the other hand, are designed to pick up "range expansion"-97 specific signatures - that is, the directions of gene flow. By analyzing the allele frequency clines 98 99 created by consecutive founder events during the expansion of a population across a landscape, as captured by a directionality index  $\Psi$ , Peter & Slatkin (2013) demonstrated how information on 100 101 the geographic origin and the direction of expansion could be extracted from genomic data through asymmetrical gene flow. That is, regression between pairwise differences of  $\Psi$  and 102 103 geographic distances between populations can be used to *directly* infer the geographic origin of expansion. However, several aspects of this approach limit its utility in practice. For example, 104 105 this method does not account for the heterogeneity in the underlying landscape during the inference procedure (i.e., assuming a strict isolation-by-distance model).  $\Psi$  may also be biased 106 toward non-zero values when local population sizes differ substantially (Peter & Slatkin, 2013). 107 Also, although it is possible to recover a signature of expansion from the magnitude of  $\Psi$ , 108 109 assessing the significance of  $\Psi$ -values, and hence, the confidence of the inferred origin, is not straightforward. 110

Here, we present a pipeline specifically developed for making statistical inferences about 111 the geographic origin of range expansion (called X-ORIGIN) that addresses these aforementioned 112 113 shortcomings. This pipeline builds upon earlier developments in spatial demographic models (e.g., Ray et al., 2010) and spatially explicit summary statistics (e.g., Peter & Slatkin, 2013). 114 Specifically, with the X-ORIGIN we couple the  $\Psi$ -index (Peter & Slatkin, 2013) with a spatially 115 explicit coalescent model for hypothesis testing in an Approximate Bayesian Computation 116 (ABC; Beaumont, Zhang, & Balding, 2002) framework. Information based on current and/or 117 118 historical habitat suitability can be estimated using ENMs and subsequently incorporated into the spatially explicit coalescent model (i.e., a modified application of SPLATCHE2; Ray, Currat, Foll, 119 & Excoffier, 2010). In addition, with the ABC framework, the estimation of the geographic 120

origin of range expansion will not be sensitive to the uncertainties in the underlying demographic parameters if a wide range of priors of demographic parameters is specified in spatial simulations. Hereafter, we refer to the geographic origin of range expansion as a parameter,  $\Omega$ . Together, the significance of expansion and the confidence of a particular geographic location for the ancestral source population are provided by the X-ORIGIN. As such, the pipeline couples information from a series of independent analyses (Fig. 1), making X-ORIGIN a useful tool for inferring the geographic origin of ancestral sources with confidence.

It should be noted that there are general procedural parallels with the integrative 128 distributional, demographic, and coalescent (iDDC) approach for model selection, which also 129 involves a series of independent analyses (i.e., estimates of habitat suitability, demographic 130 modeling, and spatially explicit coalescent; He, Edwards, & Knowles, 2013). However, the x-131 132 ORIGIN pipeline differs in that (i) it infers a novel model parameter of interest  $\Omega$  (i.e., the actual latitudinal and longitudinal coordinates), and (ii) it utilizes information from spatial summary 133 statistics, specifically, pairwise population measures of  $F_{ST}$  and the directionality index,  $\Psi$  (Peter 134 & Slatkin, 2013). As such X-ORIGIN is an approach that focuses on the estimation of a specific 135 parameter of interest –  $\Omega$ , whereas the iDDC is an approach for model selection among a set of 136 biologically informed demographic hypotheses, the foci of which vary significantly among 137 138 studies (e.g., Bemmels, Title, Ortego, & Knowles, 2016; Knowles & Massatti, 2017; Massatti & Knowles, 2016). 139

140 Here we describe the approach and test the accuracy of the X-ORIGIN pipeline in inferring  $\Omega$  under a known expansion history (i.e., simulated history; see Fig. 2). Specifically, we model a 141 142 history of expansion that involves temporal shifts in the habitat suitability of a landscape (i.e., we validate the approach by implementing a complex model which cannot be accommodated by any 143 144 other currently existing programs). We also demonstrate the utility of the X-ORIGIN with an analysis of empirical data. Specifically, we analyze the SNP dataset collected in the Collared 145 pika (Ochotona collaris) (i.e., data from Lanier, Massatti, He, Olson, & Knowles, 2015). The 146 impact of the glaciations is pronounced in small Alaskan mammals (Galbreath, Cook, 147 Eddingsaas, & DeChaine, 2011; Knowles et al., 2016; Lanier et al., 2015). While previous 148 149 analyses in the Collared pikas also suggested that contemporary environmental factors contribute less to genomic structure than a dynamic history involving the founding of current populations 150

by ancestral source populations (Lanier et al., 2015), the location of putative ancestral sourcepopulations remains unclear.

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154 Methods

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#### 156 Statistic Inferences using the X-ORIGIN pipeline

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158 The X-ORIGIN pipeline couples information from a series of independent analyses to make 159 inferences about  $\Omega$ , the geographic location of ancestral source populations, by estimating the 160 posterior probability of  $\Omega$  under an ABC framework (Fig. 1). Scripts are provided in the X-161 ORIGIN pipeline for all the steps involved and a detailed tutorial is provided on GitHub (see 162 https://github.com/KnowlesLab/X-ORGIN).

Briefly, the approach employs a spatially explicit coalescent to generate expected patterns 163 of genomic variation under a set of priors, including a prior on  $\Omega$  and priors on demographic 164 parameters of the expansion process (i.e., k and m, the local population sizes and migration rates, 165 and an ancestral population size,  $N_A$ ). That is, genomic simulations of range expansion are 166 initiated at different random locations within the geographic range specified by the prior on  $\Omega$ 167 and for different population size and migration rate values. If there is no prior knowledge on 168 possible geographic origins, all demes on the map used for demographic simulations will be 169 170 tested. Otherwise, a prior on  $\Omega$  can be based on the fossil record, or a general candidate region might be based on the regression between pairwise population differences of  $\Psi$  and geographic 171 distances (see Peters & Slatkin 2013). 172

To make inferences using X-ORIGIN that consider the effects of spatial and temporal 173 174 environmental heterogeneity on the expansion process, X-ORIGIN models the impact of this environmental heterogeneity on the expansion process. Specifically, heterogeneity in habitat 175 176 suitability might be derived from ecological niche models (ENMs) for the present or the past (Sindato et al., 2016; Waltari et al., 2007), or from information on known barriers (e.g., mountain 177 ranges, glaciers, and bodies of water; Boehm et al., 2013; Knowles & Massatti, 2017; Waltari & 178 179 Hickerson, 2013). These suitability maps are used to inform demographic dynamics associated with the expansion process by specifying different likely migration events as a function of spatial 180 and/or temporal environmental heterogeneity. Specifically, the habitat suitability scores for each 181

deme determine local population sizes, thereby influencing the actual number of migrants across demes per generation. If distributional shifts are induced by climatic changes, then temporal shifts in habitat suitability can be incorporated into the demographic modeling (i.e., applying different relative weighting of suitability information from past versus current ENMs to mirror trends of climatic change; see Brown & Knowles, 2012), given that shifts in connectivity over time can influence the expansion process, and consequently, the patterns of genetic variation across the landscape.

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## 190 Programs called up in the X-ORIGIN pipeline

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In the X-ORIGIN pipeline, demographic and spatially explicit coalescent simulations are 192 performed in SPLATCHE2 (Ray et al., 2010) in conjunction with a customized script in the x-193 ORIGIN pipeline to allow for temporally changing landscapes. Local demographic parameters 194 195 (i.e., k and m) are informed from habitat suitability by scaling these parameters proportionally to the habitat suitability values of local demes (Fig. 1), which might be temporally dynamic (i.e., 196 197 the habitat suitability for a particular location may change each generation based on shifting climatic conditions; see Brown & Knowles, 2012). Each generation, m proportion of the 198 population migrates out of the local deme; migration occurs to the adjacent four cells (north, 199 south, west, east). After the exchange of individuals, local demes grow logistically with a rate r, 200 201 and are regulated by the local carrying capacity (which are also rescaled as a function of the habitat suitability of a deme); r can be set to a specific value (e.g., He et al., 2013), and as we do 202 here (r = 1), or it can also be estimated as a parameter. For each time-forward simulation (i.e., a 203 spatially explicit map of per generation local population sizes and migration events), a series of 204 205 corresponding time-backward coalescent genetic simulation are run, with a separate coalescent simulation generated for each independent locus in the study. The ancestry of an allele will trace 206 207 back from the present into ancestral source populations, where the pattern of gene lineage coalescence across the landscape and the timing of coalescence is defined by the time-forward 208 209 local demographic simulations (i.e., the per generation k and m parameter values). SNP mutation models are then used to simulate patterns of genomic variation in SPLATCHE2, where the state of 210 each SNP is generated across the independent coalescent simulations. 211

212 To generate patterns of genomic variation to compare with the empirical data, the simulated datasets are constructed by sampling the same populations (in geographic space), the 213 214 same number of individuals, and the same number of SNPs as the empirical scenario. Summary statistics are calculated for both the empirical and simulated datasets. These include the spatial 215 summary  $\Psi$  statistics calculated within, between, across all populations, as well as pairwise 216 population  $F_{ST}$ -values; ARLEQUIN 3.5 (Excoffier & Lischer, 2010) is used to calculate  $F_{ST}$ . Note 217 that other non-spatial statistics often used in ABC analyses were also considered (e.g., K, the 218 number of haplotypes, and H, observed heterozygosity). These additional summary statistics are 219 not used in the analyses presented here because of the lack information they contained under the 220 221 expansion scenarios (see Supplementary Fig. S1); however, a user could employ them in x-ORIGIN if they determine they are relevant to the expansion history under study. 222

223 The empirical summary statistics are compared to those from the simulated data using approximate Bayesian computation (ABC), as implemented with ABCESTIMATOR in 224 ABCTOOLBOX (Wegmann, Leuenberger, Neuenschwander, & Excoffier, 2010). Rather than 225 conducting ABC analyses directly on the summary statistics, principal components (PCs) are 226 227 extracted from all predictor variables to remove the effects of interactions between summary statistics, as well as to reduce "the curse of dimensionality" (i.e., when too many statistics are 228 included, the distance between the simulated and empirical values systematically increases, 229 reducing the accuracy of parameter estimates and making it more difficult to distinguish among 230 231 models) (Wegmann & Excoffier, 2010; Wegmann, Leuenberger, & Excoffier, 2009).

Five thousand simulations (0.5%) whose transformed summary statistics are closest to 232 those calculated from the empirical genomic data are retained for estimating the model 233 parameters (i.e.,  $\Omega$ , the geographic locations of the ancestral source populations, and the 234 235 demographic parameters k, m, and  $N_A$ ). In order to jointly estimate the likelihood of a specific deme as the origin  $\Omega$  (i.e., a specific longitude and latitude), the kernel densities of  $\Omega$  across the 236 retained simulations was estimated and used as the likelihood. This provides a non-parametric 237 way of smoothing and estimating the likelihood of the origin based on the limited retained 238 simulations (i.e., from the 0.5%, or five thousand retained simulations). 239

To check if the inferred model is capable of generating the observed data, the likelihood of the empirical data given the model is compared with the likelihoods of the retained simulations. The fraction of simulations that have a smaller likelihood than the empirical data is expressed as a *P*-value, with small *P*-values indicating that a model is highly unlikely (Wegmann
et al., 2010). Likewise, we conduct standard evaluations of the quality of the inferences from
ABC (e.g., bias in parameter estimates; described below).

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247 *Performance of the X-ORIGIN pipeline* 

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We tested the pipeline on a simulated scenario (Fig. 2A) to evaluate the performance of the 249 approach for inferring the geographic location of the source population,  $\Omega$ , under a temporally 250 changing landscape. Specifically, simulations were conducted on a  $50 \times 50$  deme landscape with 251 252 a centrally located geographic barrier that was present in the past but not the present and expansion proceeding from the upper left deme (Fig. 2A). Simulations were run for 500 253 254 generations, in which the barrier persisted for 250 generations. At the end of the simulations, 10 diploid individuals were sampled from 10 demes from across the distributional map. A range of 255 migration rate, ancestral population size, and carrying capacity values per deme were simulated 256 257 to check if the inferred origin is sensitive to particular details of the demographic expansion process (Table 1). 258

The accuracy of X-ORIGIN was evaluated by measuring the geographic distance between 259 260 the actual and inferred geographic location of the source population (i.e., differences in the actual and inferred latitudinal and longitudinal coordinates). In addition to evaluating the accuracy of 261 262 the estimated  $\Omega$  under the model in which expansion proceeded from the upper left deme (Fig. 2), we also tested whether the accuracy of  $\Omega$  varied depending upon the geographic origin of the 263 264 expansion. Specifically, we investigated the performance of the model by inspecting the average error of the inferred  $\Omega$  of 10 pseudo-observed datasets (i.e., PODs from the simulations) in which 265 266 the geographic origin of the expansion differed. Specifically,  $\Omega$  was systematically varied so that each deme across the entire map served as the source of expansion. 267

In addition, the accuracy of X-ORIGIN pipeline is compared with Peter & Slatkin's (2013) original "time difference of arrival location estimation" (TDOA) approach as well as a modified TDOA approach, which incorporates spatial heterogeneity in migration patterns (Olave, He, & Knowles, in prep). Specifically, we calculated the distance between the actual geographic origin with the one estimated from the TDOA approaches. The TDOA approach identifies the origin of the expansion by identifying the deme that explains the highest proportion of variation in the correlation of pairwise  $\Psi$  differences and the pairwise differences of geographic distances of the populations to the potential origin. The modified TDOA approach correlates pairwise  $\Psi$ differences with pairwise resistance differences (McRae & Nürnberger, 2006) in which heterogeneous landscape is considered (Olave et al., in prep), whereas the original TDOA (Peter & Slatkin, 2013), assumes migration occurs on a homogeneous landscape (i.e., according to a random diffusion model). We conducted a cursory examination of the robustness of X-ORIGIN to model mis-specification as well.

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282 Demonstration of X-ORIGIN with application to Alaskan Collared pika

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In addition to details about the ABC analyses (see below), here we briefly describe the empirical 284 285 genomic data we analyzed with X-ORIGIN, given that all data used here are from previous publications and are referenced below. Specifically, we analyze a genomic dataset collected in 286 287 the Alaskan Collared pika (for details on library construction and rigorous quality filtering see Lanier et al. 2015). Maps of environmental heterogeneity used in the X-ORIGIN analyses to infer 288 289  $\Omega$ , the geographic location of the ancestral source population for the Collared pika, were generated from ENMs for the present and the last glacial maximum, LGM (see details in 290 Knowles et al. 2016). 291

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293 Genomic dataset. We analyzed RADseq data for 8 populations; note we excluded the Pika Camp (Wrangell-St. Elias Mtns; GIS coordinates 61.2170, -138.2670) from our analyses because 294 295 previous analyses indicate that it was founded from a separate ancestral refugial source (Lanier et al. 2015). Of the 23,493 RADseq loci with at least one biallelic SNP across populations, we 296 297 analyzed 6816 loci with one SNP retained per RADseq loci in 50 individuals (i.e., 6-8 individuals per population, with the exception of Jawbone Lake, where n = 2; loci in less than 298 299 50% of the samples or were not present in more than one individual per population were excluded. Note that this is an expanded dataset relative to those previously published (i.e., Lanier 300 301 et al. 2015; Knowles et al. 2016) because we recovered more genetic information using ddRAD 302 aligned to a reference genome for *Ochotona princeps* (American pika; ID: 771).

The directionality index  $\Psi$  requires information on the ancestral versus derived states of SNPs because the statistic is calculated by counting the difference in derived allelic frequencies

between pairs of populations (see Eq. 1 in Peter & Slatkin 2013). Ancestral states of independent
biallelic SNPs were determined by aligning the sequences with *Ochotona princeps* (American
pika; ID: 771; <u>https://www.ncbi.nlm.nih.gov/genome</u>).

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Prior on  $\Omega$ , the geographic locations of the origin of expansion. The TDOA approach was 309 conducted to select candidate regions of origin to inform the prior on  $\Omega$  (as opposed to 310 considering the entire state of Alaska). Specifically, for each potential geographic location as the 311 site of the ancestral source population (i.e., each deme from the distributional map), linear 312 regression was performed between pairwise  $\Psi$  differences and the pairwise differences of 313 geographic distances of the populations to the potential origin. The linear regression was 314 repeated for each of the different potential geographic origins and the geographic locations with 315  $R^2$ -values larger than 0.5 were used to specify the prior on the geographic location of the 316 ancestral source population (regression analyses were conducted using modified scripts from 317 Peter & Slatkin, 2013, which we provide on KnowlesLab/Github). This generated a target area of 318 approximately 442,300  $km^2$  (i.e., 1302 demes, with a size of  $18.4 \times 18.4 km^2$  for each deme: 319 320 Table 1) to analyze in detail regarding the posterior probability of  $\Omega$ , the geographic location of the ancestral source population for the set of 8 Collared pika populations collected across its 321 range (see Lanier et al. 2015 for details). 322

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324 Estimates of habitat heterogeneity across space and time. Maps of environmental heterogeneity for the Collared pika were generated from ENMs (see details in Knowles et al. 2016). Briefly, 325 326 inferences about differences in habitat suitability across space were made for the present and the LGM from ENMs based on bioclimatic data for the present and paleoclimatic data from 21 kya. 327 328 The models were tested over combinations of regularization parameters from 0.25 to 3 in intervals of 0.25 and the Linear, Quadratic, Hinge, Product and Threshold features using 329 330 SDMTOOLBOX (Brown, 2014). Each model parameter class was replicated 25 times using crossvalidation. 331

In addition, temporal shifts in habitat suitability were represented using differences in the relative weighting of habitat suitabilities estimated for the present and LGM across time to reflect climatic trends in the region over the past 21,000 years (Brown & Knowles 2012). Specifically, the current ENM suitability map was used to represent the present to 5,000 years ago, an intermediate suitability map (i.e., an average suitability between the current and LGM
ENMs) for the time period 5,000 - 11,000 years ago, and the LGM ENM suitability map for
11,000 - 21,000 years ago.

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ABC analyses. Datasets were simulated for 2100 generations (based on a scaling factor of 10 to reduce the computational requirements; see He et al. 2013) to represent the range expansion from last glacial maximum. Priors for the local carrying capacity (k), ancestral population size ( $N_{ans}$ ), and migration rates (m) are set according to Lanier *et al.* (2015) (see Table 1). Note that a geographic grid of  $18.4 \times 18.4 \text{ } km^2$  corresponded to a single deme and expansion was modeled across the Alaskan landscape (i.e., over approximately 2,197,850  $km^2$ ).

As with tests of the general performance of X-ORIGIN, we compared the estimates of  $\Omega$ , 346 the geographic location of the ancestral source of expansion, with results from: 1) the TDOA 347 method, where heterogeneity in the present landscape is not incorporated (i.e., the geographic 348 349 distances separating populations were represented as pairwise Euclidean distances), 2) the modified TDOA method, where resistance distances based on heterogeneity in the current habitat 350 351 suitability is used, and 3) X-ORIGIN, where temporal shifts in the heterogeneity of the landscape over time are accounted for. To evaluate the accuracy of estimates of  $\Omega$ , five thousand pseudo-352 353 observations were generated from the prior distributions of the parameters. If the estimated parameters are unbiased, posterior quantiles of the parameters from the pseudo datasets should 354 355 be uniformly distributed (Cook, Gelman, & Rubin, 2006; Wegmann et al., 2010). The posterior quantiles of true parameters for each pseudo run were calculated based on the posterior 356 357 distribution of the regression adjusted 5000 simulations closest to the pseudo-observed datasets.

- 358
- 359 Results
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- 361 *Performance of the X-ORIGIN pipeline*
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For the example history considered here, which involved a central barrier that was present in the past, but not the present (i.e., there is both spatial and temporal heterogeneity in habitat suitabilities) X-ORIGIN gives more accurate inferences of  $\Omega$ , the geographic location of the source population of the expansion, than the TDOA approach. In fact, the performance of x-

ORIGIN was quite good, estimating the most likely origin within 1- 4 demes of the actual origin (mean *P*-value = 0.67) from different starting positions across the map (and hence, differences in when and where the expansion process interacted with the geographic barrier), except for the lower left grid of the geographic area (Fig. 3A, C; see Supplementary Fig. S2 for detailed examples of variation in inferences across PODs for different locations of origins).

372 In contrast, the majority of analyses with the TDOA approach give inferred locations that differ markedly from the actual area where the expansion originated, irrespective of where on the 373 map the expansion originates (Fig. 3B). The performance of the TDOA approach was especially 374 poor (i.e., large discrepancies between the inferred and actual geographic origin of expansion) 375 376 when the ancestral source area was near the barrier (Fig. 3D). This variation in accuracy highlights the importance of explicitly modeling the temporal heterogeneity of landscapes (also 377 see Wegmann, Currat, & Excoffier, 2006), as it strongly distorts the  $\Psi$  signatures, especially if 378 the heterogeneity is present in the early stage of the expansion. 379

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#### 381 Inferred geographic origin of expansion in the Alaskan Collared Pika

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For the set of Collared pika populations studied here, the highest likelihood (marginal density:  $1.82 \times 10^{-8}$ ; *P*-value: 0.996) for the location of the expansion origin,  $\Omega$ , is the Mackenzie Mountains in Yukon Territory, Canada (Fig. 4). This inference is based on the retained 5000 simulations whose summary statistics were to those of empirical data. The geographic origin of expansion (i.e., the latitudinal and longitudinal coordinates) was estimated using a twodimensional kernel density of the retained simulations, implemented using the kde2d function in the MASS package of R (Venables & Ripley, 2002).

The geographic origin of expansion inferred using X-ORIGIN differed from the TDOA results (Fig. 4). Moreover, neither the inferred area based on the pairwise  $\Psi$  matrix on a homogeneous landscape (TDOA-diffusion) nor the one based on a resistance map of the current landscape suitabilities (TDOA-resistance), are in areas with high likelihoods. That is, simulated genetic data sets where expansion proceeded from the inferred areas under the TDOA approaches do not correspond to the observed genetic data (i.e., there is a mismatch between the empirical summary statistic and those calculated from the simulations). 397 Based on the distances between actual versus inferred origin for each of the different method, X-ORIGIN outperformed TDOA, although the accuracy of inferred  $\Omega$ -values varied 398 399 depending upon the geographic origin of the expansion (Fig. 5). We also note that the accuracy was generally lower for the heterogeneous landscape inferred for pikas relative to the landscape 400 used to validate the X-ORIGIN package (Fig. 5 versus Fig. 3). In particular, populations that 401 originated from the southeast region exhibited the lowest accuracy (i.e., the greatest difference 402 between the inferred and actual value of  $\Omega$ ). This is most likely due to the lack of samples from 403 that area, and consequently little information of the direction of asymmetrical gene flow 404 expected under an expansion model (see Peter & Slatkin, 2013). Nevertheless, comparison of the 405 accuracy of inferences between X-ORIGIN and TDOA approaches, indicate those from X-ORIGIN 406 are more accurate for an expansion originating from the Mackenzie Mountain range. 407 408 Specifically, analyzing simulated data of expansions from the Mackenzie Mountain range (i.e., the PODs from the ABC simulations), the TDOA approaches give estimates that are generally 409 410 displaced by 15 to 30 demes from the actual origin of expansion (i.e., a discrepancy of 750 to 1500 km), and curiously these were more inaccurate than inferences with a south-west 411 geographic origin of expansion (Fig. 5), despite sampling of populations in that region (see 412 discussion below). 413

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415 DISCUSSION

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Patterns of genetic variation in individuals sampled in the present harbor rich information about 417 past movements of species. In contrast to those from non-spatial models of population 418 demography (e.g., changes in population size or admixture proportions; see Hey, 2005; Theis, 419 420 Ronco, Indermaur, Salzburger, & Egger, 2014), recent developments have focused on inferences from spatially explicit approaches. Specifically, departure from equilibrium status of population 421 422 movements under a diffuse model, 'isolation-by-distance', caused either by range expansion/contraction history, long distance admixture or habitat heterogeneity is tested through 423 different approaches. One general approach is to quantify discrepancies between spatial genetic 424 425 patterns and the expectations from geographic distances. For example, discrepancies between population's positions on a genetic PCA map can be visualized against a map of their 426 geographical distribution using Procrustes analyses to examine where on a landscape patterns of 427

428 genetic variation depart from isolation by distance (Knowles et al., 2016; Wang, Zöllner, & 429 Rosenberg, 2012), or a "geogenetic map" can be used to infer potential long-range admixture 430 among populations (Bradburd, Ralph, & Coop, 2016). Similarly, disruptions to past movement 431 might be inferred by relating the effective migration rates to expected genetic dissimilarities for 432 an interpolated geographical map of barriers or corridors among populations (see Petkova, 433 Novembre, & Stephens, 2016).

Instead of quantifying discrepancies from isolation-by-distance, our approach directly 434 435 models expected patterns of genetic variation using spatial genetic indices and makes inferences about historical movements – specifically, the geographic origin of expansion,  $\Omega$  – under an 436 ABC framework, while incorporating temporal shifts in habitat suitability over time. This is not 437 the first approach for directly evaluating genetic variation under models of historical movement. 438 439 For example, the spatial genetic indices applied here were developed to directly infer historical movements based on shifts in the genetic summary statistics across a landscape (Peter & Slatkin, 440 2013), and spatial-autocorrelation of genetic covariance information has been applied to 441 distinguish among spatially-explicit demographic scenarios (Alvarado-Serrano & Hickerson, 442 443 2016; Bertorelle & Barbujani, 1995; Coop, Witonsky, Rienzo, & Pritchard, 2010). However, our approach infers and evaluates the parameter  $\Omega$  – the actual latitudinal and longitudinal 444 coordinates for the origin of an expansion- that is not based on the assumption of a diffusion 445 model that provides statistical rigorousness and flexible applications for inferences about 446 447 historical expansion scenarios. First, we can evaluate the likelihood of different geographic locations for the origin of a population expansion, accounting for both spatial and temporal 448 449 heterogeneity in habitat suitability of the landscape. Second, with the freely available X-ORIGIN pipeline we developed, users can validate any inference by determining whether the inferred 450 model is capable of generating data that generally corresponds to the empirical data, which is 451 equally important as estimating the most likely model for the origin of expansion (i.e., the most 452 453 likely location for the origin of expansion may nonetheless be a poor fit to the observed data). Such attributes are not currently implemented in other methods for inference about expansion 454 455 histories (e.g., compare with Ray et al., 2005).

Below we discuss how these attributes make X-ORIGIN not only a practical tool, but as our analyses demonstrate, also one whose performance is better than not accommodating such dynamic histories. Likewise, we highlight how this pipeline can easily be adapted for a more 459 general inference approach beyond inferring the origin of expansions, especially with the development of new spatial indices. However, we also note the difference in performance of X-460 ORIGIN between a simple demographic history (i.e., the one used to validate the approach) and 461 the one with more extreme habitat heterogeneity, and caution users of the importance for 462 validating the accuracy of the inference, which can be implemented in the X-ORIGIN pipeline. We 463 apply this practice when interpreting the results from the X-ORIGIN analysis of the Collared pikas, 464 as well as discuss aspects of the data that might contribute to uncertainty in the inferred origin of 465 expansion, and the importance of corroborative evidence not based on the genetic data itself. 466

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# 468 *Factors impacting the accuracy of inferences about the geographic origin of expansion*

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The  $\Psi$  index directly captures the overall trend of differences in frequencies of derived 470 polymorphic alleles in populations based on the fact that expanding front of populations are 471 experiencing serial bottlenecks. Therefore,  $\Psi$  indices are informative as long as current 472 473 populations have not yet reached equilibrium. If the majority of the pairwise  $\Psi$  indices are close 474 to zero in the system (which is not the case for pikas; Supplementary Table 1), the lack of spatial gradient in the  $\Psi$  indices indicate that either there was not an expansion or a sufficient amount of 475 time since the expansion has passed such that its genetic signature can no longer be detected by 476 the  $\Psi$  indices (see also Peter and Slatkin 2013). We tested a scenario in the Pika dataset where 477 478 there is no expansion origin to examine the performance of X-ORIGIN. Specifically, we simulated 479 1000 replicate data sets in which all populations started from their sampling areas to reach 480 equilibrium states. For these datasets, although  $\Psi$  indices deviate strongly from zero, no origin can be estimated from TDOA as no positive relationship between pairwise differences of  $\Psi$  and 481 482 geographic distances among populations can be established (Supplementary Table 2). Likewise, with X-ORIGIN, marginal densities of the expansion model are extremely low (on the order of 483  $10^{-200}$  to  $10^{-12}$  as compared to  $10^{-8}$  for PODs that experienced expansion from a single origin) and 484 P-values are zero (Supplementary Table 2). Therefore, X-ORIGIN, like TDOA, will not give 485 486 misleading results about the potential origin for expansion when no such expansion occurred.

Any inference that extracts information on the geographic distribution of genetic variation requires adequate sampling of populations as well as number of independent SNPs (i.e., at least more than 1,000 independent SNPs; Peter & Slatkin, 2013; Bradburd et al., 2016). Our results 490 clearly show that inferences become less accurate when sampled populations are located further from the location where an expansion originated (e.g., see higher error rate at south-east corner 491 492 of Fig. 5A). Therefore, researchers should carefully consider the sampling design. In particular, 493 our results (see also Peter & Slatkin, 2013; Bradburd et al., 2016) suggest that obtaining accurate inferences that utilize spatial information about the distribution of genetic variation may be 494 495 dependent upon which populations are sampled, rather than whether there is sufficient power for such inferences related to the number of loci analyzed. Although it's beyond the scope of this 496 study, this general question is something that could be explored using the X-ORIGIN pipeline. 497

Another factor impacting the accuracy of inference relates to model mis-specification. 498 Specifically, complicated demographic scenarios such as those involving two or more 499 geographic origins of expansion will give misleading results if not accommodated (see also Peter 500 & Slatkin 2013). There are a number of ways to accommodate and/or test whether an assumed 501 expansion from a single source might be violated. For example, clustering algorithms can be run 502 503 to delineate populations into different groups with potentially different expansion origins and validated by a minimum spanning tree built from a matrix of  $\Psi$ -values (Peter & Slatkin, 2013), 504 505 followed by separate inferences of  $\Omega$  for each subgroup of populations. Alternatively, competing explanatory models with multiple origins versus one expansion origin can be analyzed in X-506 ORIGIN and compared in a model selection framework. Our results also suggest that any model, 507 even those that might be more probable than others, should be interpreted with caution if  $\Omega$  is 508 509 located in areas with low confidence (based on reference to simulated datasets), or if the most likely model nevertheless has a low probability of generating data that resembles the empirical 510 511 data (i.e., low *P-value*; Wegmann et al. 2010; see He et al. 2013 for details of model validation).

Despite positive aspects of X-ORIGIN related to estimating the likelihood of the expansion 512 513 origin, and consequently, uncertainty surrounding this inference (e.g., the geographic area spanned by the 90% highest posterior density of  $\Omega$ ), as well as validating the inference using 514 515 PODs (see Fig. 5), one unexplored issue is how errors early in the pipeline might get amplified and generate misleading results. We did a cursory examination of how such errors might impact 516 an inferred expansion origin. Specifically, we examined how robust the inferred origin might be 517 518 to uncertainties regarding the temporal changes in habitats - in this case, the duration of a barrier, as in the scenario we used to validate X-ORIGIN (see Fig. 2). When we varied the true 519 duration of the barrier to simulate data (i.e., simulate data with a barrier that persisted for 200 to 520

300 generations, rather than 250 out of the 500 generations), we observed no difference in the accuracy of the  $\Omega$  estimation (Supplementary Fig. S3). This shows that the pipeline can be robust to misspecification of temporal dynamics of a historical scenario (at least for the parameter space examined here). This clearly should not be interpreted as general evidence of robustness to model mis-specification. Rather we present it here to show that X-ORIGIN exhibits some robustness, but also to emphasize that all users can conduct their own investigation to robustness tailored to the specifics of their application.

There are of course other paths for errors that could impact the accuracy of inferences 528 about  $\Omega$ . For example, we use ENMs to estimate potential suitable areas to inform demographic 529 models (see Fig. 1). As a consequence, the results from X-ORIGIN could be impacted by poor 530 ENMs (i.e., validation and best practices of ENMs should be followed). In addition, applying 531 532 different transformation of habitat suitabilities into local carrying capacities can affect patterns of genetic variation (see Brown & Knowles 2012). There are different strategies one might take to 533 534 avoid biases that could result from unrealistic assumptions or errors in the upstream steps of the pipeline (Figure 1). For example, instead of using a fixed suitability score from an ENM model 535 536 for each deme, suitability scores between maximum and minimum range inferred for each deme might be randomly sampled during the simulation process to generate expected patterns of 537 genetic variation that incorporate some uncertainties in the ENM modeling. This might increase 538 the number of simulations required for inferring  $\Omega$  to get an unbiased and precise estimate under 539 540 an ABC framework, given that accommodating such uncertainties may increase the variance in observed patterns of genetic variation in simulated datasets. Likewise, different transformations 541 542 of habitat suitabilities into local carrying capacities (scaling habitat suitability linearly with local carrying capacity versus a step function; Brown & Knowles, 2012) could be incorporated as 543 544 alternative models to be tested (i.e., treated in a model selection framework, even when the primary interest is on inferring the origin of expansion,  $\Omega$ ). 545

Although such flexibility in accounting for uncertainty or potential errors in upstream steps (Fig. 1) is a strength of the X-ORIGIN package we developed, the application of X-ORIGIN (especially compared with TDOA; Peter & Slatkin, 2013) comes with much more computational expense. For example, a typical spatially explicit simulation of 2000 generations on a 150 x 150 grid layer and the generation of 1000 SNPs takes more than 7 minutes. Users are advised to calculate required computational resources before experimenting with the pipeline. This includes reducing the size of the  $\Omega$  prior (e.g., by applying TDOA as a preliminary step for data inspection, as applied in the Collared pika example).

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#### The Mackenzie Mountain region as the most likely origin of expansion in Collared pika

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As an alpine small mammal, suitable habitats for Collared pika are spatially highly 557 heterogeneous, but also temporally heterogeneous given that Alaska was directly impacted by 558 the glacial cycles (Fig. 2B). Previous analyses have suggested a potentially complex 559 biogeographic history involving expansion from multiple ancestral sources (Knowles et al., 560 2016; Lanier et al., 2015; Lanier & Olson, 2009). Limited sampling of populations inhibits 561 analysis of data subsets to explore such models with X-ORIGIN (i.e., multiple populations are 562 required to estimate potential sources of expansion), and therefore is beyond the scope of this 563 manuscript. Nevertheless, it is informative to consider how our inference compares to previous 564 565 characterizations for the populations analyzed here.

Previous studies that made inferences about the biogeographic and demographic history 566 of the Collared pika applied analyses that assumed equilibrium status (e.g.,  $F_{ST}$ , STRUCTURE 567 analyses, estimates of phylogenetic relationships among populations). For example, in an 568 analysis of the relationship between  $F_{ST}$ -values among populations and the geographic distance 569 570 separating them (Lanier et al., 2015), the most north-eastern sampled population Jawbone Lake (Fig. 4) appeared to be an outlier under the expectation of isolation-by-distance. Based on this 571 572 result, and the relative genetic distinctiveness of the Jawbone Lake population and the other two 573 north-central populations from the Yukon-Tanana Uplands (specifically, the Eagle Summit and Cresent Creek populations), these populations were analyzed separately and a distinct pattern of 574 isolation by distance at the regional level was interpreted as possible evidence of different 575 576 ancestral source populations (Lanier et al., 2015). However, our analyses here provide a 577 compelling argument for an alternative explanation. Specifically, the genetic similarities between Jawbone Lake and the Eagle Summit and Crescent Creek populations (See Fig. 5 in Lanier et al., 578 579 2015) may not reflect a refugial source that was differed from the refugial source of other sampled populations. Instead, it may reflect their proximity to the geographic origin of expansion 580 581 in an ancestral species,  $\Omega$  in the Mackenzie mountains (see Fig. 4), and more specifically, the similar geographic distance of the populations from the source of expansion. Even though our 582

583 validation tests indeed show that the degree of reliability about expansion can be considerable (e.g., differing by as much as 1500 km from the actual expansion origin depending upon where 584 on the landscape the expansion proceeded from; Fig. 5), the mean error surrounding estimates of 585  $\Omega$  as a function of the distance from the actual origin is quite low (i.e., less than 5 demes away, 586 or 250 km) for the geographic region with the highest likelihood of  $\Omega$  (Fig. 4). Interestingly, 587 Procrustes analyses in the Collared pikas, as well as other co-distributed alpine mammals, 588 suggest a stronger deviation along the longitudinal axis between genetic variation and geography 589 (i.e., genetic similarities more centrally located than the geographic space occupied by the 590 populations; Knowles et al., 2016). Our analysis supported this deviation as a result of an 591 592 expansion history along this axis, offering an alternative interpretation to the hypothesis of a centrally located refugium. 593

594 Lastly, ENMs for the LGM are not inconsistent with our estimate (Fig. 2B). However, if we consider information from the ENMs by themselves, the region of high habitat suitability 595 596 encompasses a broad area that does not offer much detail about the potential location of ancestral 597 populations. This even includes a potential north-western source population (Fig. 2B), even 598 though former genetic (Knowles et al., 2016; Lanier et al., 2015) and fossil studies (Gunderson, Jacobsen, & Olson, 2009; Lanier & Olson, 2013) suggest the lack of support for such a putative 599 600 ancestral source (e.g., in the Brooks Range). Both X-ORIGIN and TDOA analyses reinforce that despite projections from the ENM for the LGM, this region does not appear to be a likely 601 candidate as an ancestral source of expansion. 602

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#### 605 CONCLUSIONS

606 Our results show that failing to consider the impact of spatial and temporal heterogeneity on the expansion process can lead to much less accurate inferences (Fig. 3A compared with 3B, 607 608 and Fig. 5A compared with 5B). Furthermore, there are also ways to minimize potential errors when inferring the origin of expansion. For example, in our simulations, we place a broad prior 609 on parameters that are not targets of interest, but may influence estimates of  $\Omega$  (e.g., ancestral 610 population, carrying capacity; see Table 1), thereby accounting for uncertainty about the 611 612 demography of the expansion process. Moreover, the summary statistics used in the inference procedure (i.e.,  $\Psi$  and  $F_{ST}$ -values) are not sensitive to the absolute effective population sizes, but 613

614 rather the ratio of size differences between population pairs. Lastly, despite the lower accuracy of inferences for complicated scenarios, as with the analysis of the Collared pika, relative to simple 615 expansion scenarios (Fig. 3, 5), accounting for the effects of spatial and temporal heterogeneity 616 is generally more accurate than applying an oversimplified model if the goal is to infer the 617 geographic location of an expansions origin (Fig. 3). Therefore, we argue that the caveats and 618 concerns associated with inferring the origin of expansion do not nullify the utility of spatially 619 and temporally explicit models, such as those applied here in the new X-ORIGIN pipeline. In 620 particular, we show that it is incorrect to assume that environmental heterogeneity (whether 621 temporal or spatial) will not impact inferred origins of expansion, and that despite the caveats we 622 highlight with X-ORIGIN, they are less problematic than many implicit assumptions made in 623 approaches that ignore geographic and temporal constraints on population movements or 624 625 population size fluctuations (see Knowles & Alvarado-Serrano 2010). Moreover, the reliability of any inference about the origin of expansion under the more complex models implemented in 626 the X-ORIGIN pipeline can be (and should be) rigorously explored using validation procedures. 627

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#### 631 ACKNOWLEDGEMENTS

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Specimens and genomic data analyzed here were collected as part of past collaborative work with Hayley Lanier and Link Olson, for which we are grateful for their helpful discussions on small Alaskan mammals, and Collared pikas in particular. We appreciate the comments from two anonymous reviewers and Benjamin Peter that helped improve earlier versions of the manuscript. The authors declare no conflict of interest.

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785	TABLE
786	Table 1. Prior ranges for demographic and genetic parameters used in the demographic

787 simulations.

Paramete	ers Description	Prior Ranges	Distribution
m	migration rate between demes	$(10^{-3.6}, 10^{-2})$	log-uniform
Nans	ancestral population size before expansion	(36,880, 508,318)	uniform
K	carry capacity per deme	$(10^{3.3}, 10^{4.6})$	log-uniform
Lat	latitude range of origin	(1,073,893, 1,850,478)	uniform
Long	longitude range of origin	(616,487, 899,496)	uniform

- 788
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- 790 FIGURE LEGENDS

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Figure 1. The required data inputs (shown in boxes) and workflow of the X-ORIGIN pipeline are 792 793 highlighted in the schematic. Specifically, to infer the geographic location from which an expansion originates,  $\Omega$  (i.e., the actual latitudinal and longitudinal coordinates of the ancestral 794 source population), a habitat suitability map, candidate regions of  $\Omega$ , and priors for demographic 795 parameters are required. To consider how habitat heterogeneity might impact the range 796 expansion process, the habitat suitability map can be informed by spatial (as well as temporal) 797 variation in suitability (e.g., from ENMs based on contemporary bioclimatic variables, or 798 paleoclimatic variables; see He et al. 2013). Otherwise, the expansion process can be modeled as 799 800 a diffusion process (i.e., equal habitat suitability across space and time). Likewise, users have the option of either entering candidate regions of  $\Omega$  (e.g., a region identified by the regression 801 approach of Peter and Slatkin 2013; as discussed in the text), or the entire map area can be 802 evaluated during the inference procedure. The pipeline calls up different software packages for 803 804 downstream generation of simulations and estimation of the expansion origin, candidate regions of  $\Omega$ . Specifically, spatially explicit coalescent simulations are used to generate expected patterns 805 806 of genetic variation under a demographic model the expansion process (either informed or not by spatial and temporal heterogeneity of the landscape) using a modified version of the program 807 808 SPLATCHE2 (Ray et al. 2010). Summary statistics are calculated from each simulated data set using R script that are incorporated in the pipeline, which are compared with those calculated for 809 810 empirical data to inform the posterior distribution of  $\Omega$  using ABC. Note that all steps can be performed seamlessly in X-ORIGIN, which has a wrapper for connecting all the steps in R or 811 812 python scripts. Scripts for the pipeline are shown in grey shaded boxes, while external programs called in the pipeline are shown without boxes. 813

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Figure 2. Simulated scenario used to evaluate the performance of the X-ORIGIN pipeline for inferring the geographic origin of a range expansion. In the simulated scenario, A) expansion proceeded from the lower-left corner of the map (shown as the red dotted area) across a homogeneous landscape with a centrally located geographic barrier during the first 250 generations, but not the last 250 generations (i.e., there is spatial and temporal habitat heterogeneity, where the area of the barrier has zero suitability). Due to the symmetry of the landscape, we varied the origin of expansion in the simulations within the red dotted area instead

822 of the whole map. Circles mark populations that are sampled and for which summary statistics are calculated from multiple individuals. B) An empirical application of X-ORIGIN in the 823 824 Collared pika in which habitat suitability varied spatially and temporally across the Alaskan 825 landscape. Ecological niche models were used to estimate habitat suitabilities for the present and past (i.e., the LGM) using climatic data (see Lanier et al. 2015 for details about ENMs). 826 827 Specifically, the demographic expansion process proceeded across a temporally and spatially heterogeneous landscape, in which the habitat suitabilities from an ENM estimated for the LGM 828 was used to inform the first 5,000 years of the simulated demographic expansion, followed by 829 6,000 simulated years of expansion across an intermediate surface (i.e., a map with average 830 habitat suitability scores between those from the ENM for the present and LGM), and then 831 10,000 years of expansion with the habitat suitabilities from an ENM based on current climatic 832 conditions. 833

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Figure 3. Distribution of the mean errors in the estimated  $\Omega$  across the map (i.e., for different 835 geographic locations for the origin of expansion) under the simulated scenario (see Fig. 2A, the 836 837 red dotted area) using (A) X-ORIGIN versus (B) the TDOA approach. Color of each deme shows the accuracy of origin estimation if the expansion starts from that particular deme, which is 838 measured by the distance between its inferred origin,  $\Omega$ , and the actual origin, averaged across 10 839 simulations. Also shown are the histograms of accuracy across all 5000 instances (C) from x-840 841 ORIGIN versus the TDOA approach. Distances are in the units of the number of demes from the actual origin. 842

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Figure 4. Estimates of the origin of expansion,  $\Omega$ , inferred in the Collared pika using X-ORIGIN 844 845 compared with the TDOA approach. The deme with the highest likelihood inferred from x-ORIGIN is marked with a black "X", whereas the location of origins estimated using TDOA 846 methods are marked with crosses. The heat map shows differences in the probability density 847 estimates of different demes across the map being the origin of expansion, as estimated in x-848 ORIGIN, with the greener shades representing higher probabilities; the shaded square area 849 represents the prior area for  $\Omega$ , a region identified by the regression approach of Peter and 850 Slatkin (2013). Each deme in the map has equal relative size (i.e., the map is projected using the 851 852 North American Datum – NAD83 – readjustment of the global positioning system that accounts

for the earth's curvature) and population localities of sequenced individuals are marked by greycircles.

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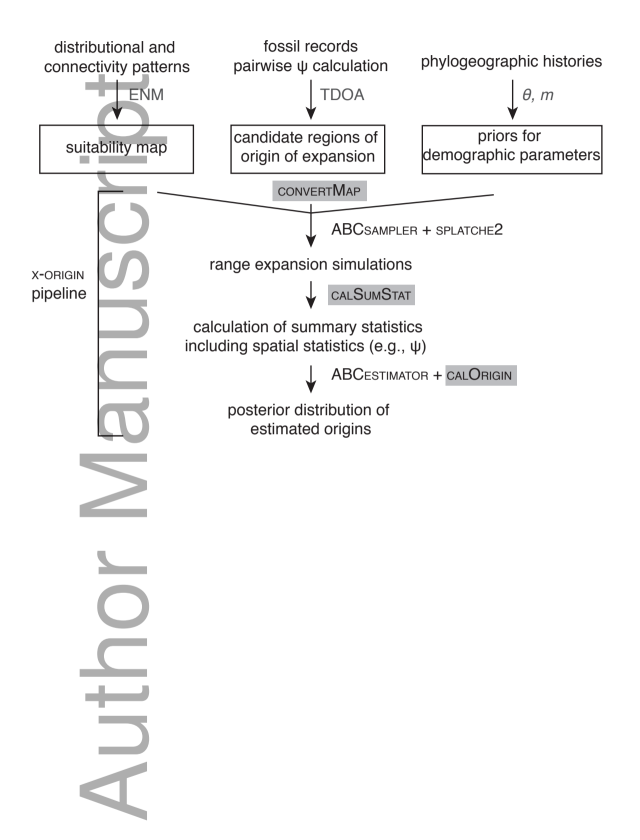
Figure 5. Distribution of mean errors in the estimated  $\Omega$  across the map (i.e., for different 856 geographic locations for the origin of expansion) for pseudo-observations in the Pika simulations 857 (see Fig. 2B) using (A) X-ORIGIN versus (B) the TDOA approach. 5000 pseudo-observations are 858 generated and color of each deme shows the accuracy of origin estimation if the expansion starts 859 from the particular deme, which is measured by the average distance between its inferred origins 860 and the actual origin. White area on the map contains demes where not all populations can be 861 colonized if the expansion starts from there. Also shown are the histograms of accuracy across 862 all 5000 instances (C) from X-ORIGIN versus the TDOA approach. Distances are in the units of 863 864 the number of demes from the actual origin and each deme is 18.4km in length.

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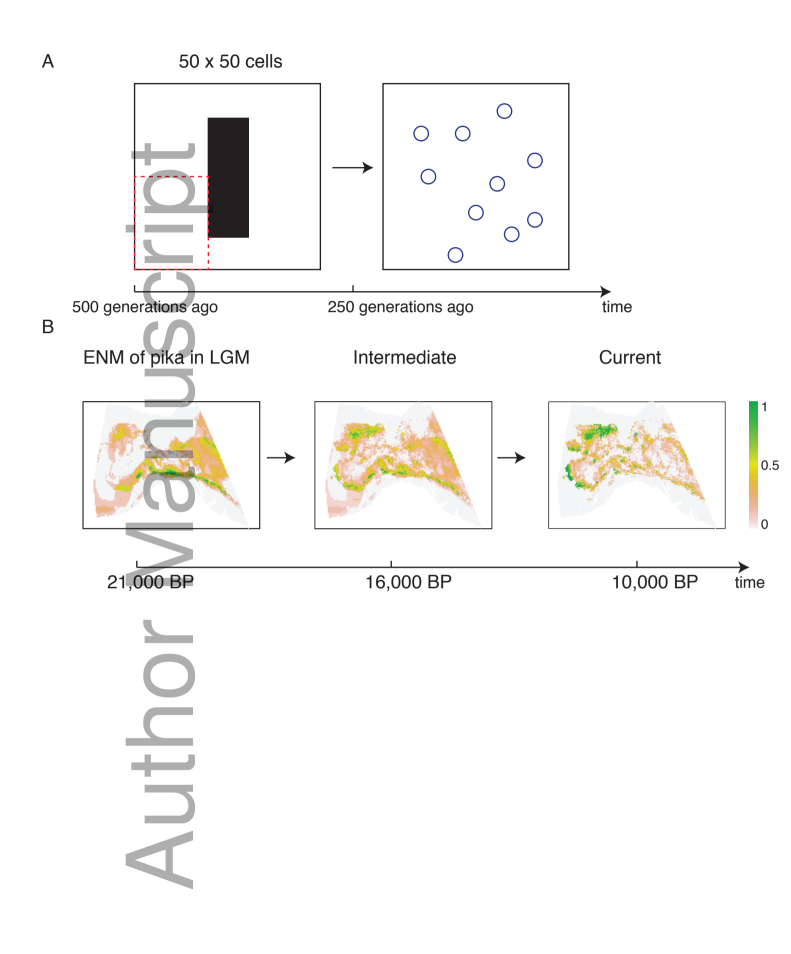
#### TABLE

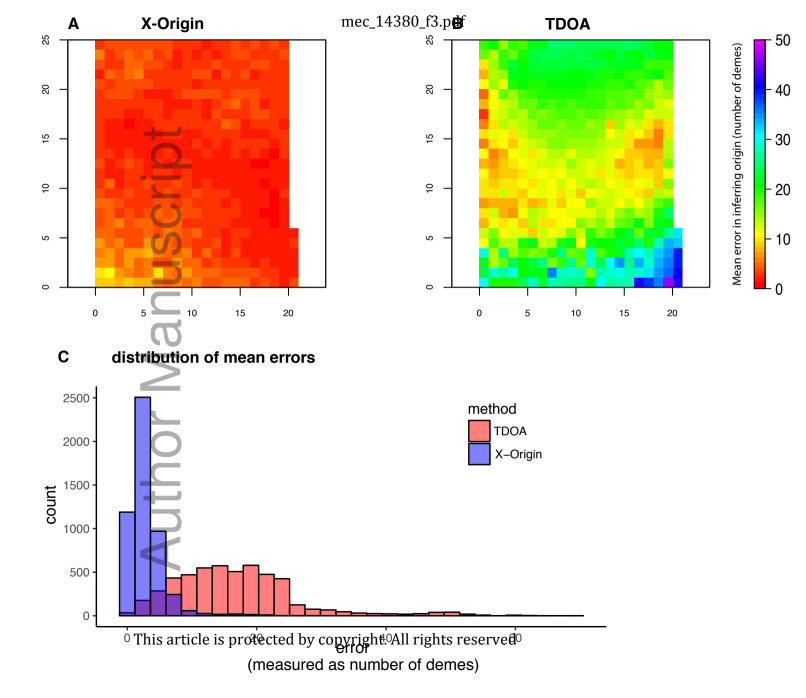
Table 1. Prior ranges for demographic and genetic parameters used in the demographic simulations.

Parameters Description		Prior Ranges	Distribution
m	migration rate between demes	$(10^{-3.6}, 10^{-2})$	log-uniform
N <sub>ans</sub>	ancestral population size before expansion	(36,880, 508,318)	uniform
К	carry capacity per deme	$(10^{3.3}, 10^{4.6})$	log-uniform
Lat	latitude range of origin	(1,073,893, 1,850,478)	uniform
Long	Plongitude range of origin	(616,487, 899,496)	uniform
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Comparisons of the origin inferences

