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Spatial Population Genetics: It’s About Time

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Abstract
Many important questions about the history and dynamics of organisms have a geographical component: How many are there, and where do they live? How do they move and interbreed across the landscape? How were they moving a thousand years ago, and where were the ancestors of a particular individual alive today? Answers to these questions can have profound consequences for our understanding of history, ecology, and the evolutionary process. In this review, we discuss how geographic aspects of the distribution, movement, and reproduction of organisms are reflected in their pedigree across space and time. Because the structure of the pedigree is what determines patterns of relatedness in modern genetic variation, our aim is to thus provide intuition for how these processes leave an imprint in genetic data. We also highlight some current methods and gaps in the statistical toolbox of spatial population genetics.
1. INTRODUCTION

The field of population genetics is shaped by a continuing conversation between theory, methods, and data. We design experiments and collect data with the methods we will use to analyze them in mind, and those methods are based on theory developed to explain observations from data. With some exceptions, especially the work of Sewall Wright (1940, 1943, 1946) and Gustave Malécot (1948), much of the early body of theory was focused on developing expectations for discrete populations. This focus was informed by early empirical data sets, most of which (again, with exceptions, like Dobzhansky & Wright 1943; Dobzhansky 1947) were well-described as discrete populations—like samples of flies in a vial (Lewontin 1974). In turn, the statistics used to analyze these data sets—e.g., empirical measurements of $F_{ST}$ (Wright 1951) or tests of Hardy-Weinberg equilibrium (Hardy 1908, Weinberg 1908)—were defined based on expectations in discrete, well-mixed populations. In part, this was because much more detailed mathematical predictions were possible from such simplified models. The focus of theory and methods on discrete populations, in turn, influenced future sampling designs.

To what degree does reality match this picture of clearly delimited yet randomly mating populations? Often, the answer is: not well. Organisms live, move, reproduce, and die, connected by a vast pedigree that is anchored in time and space. But it is difficult to collect data and develop theory and methods that capture this complexity. Sampling and genotyping efforts are limited by access, time, and money. Developing theory to describe evolutionary dynamics without the simplifying assumption of random mating is more challenging. For example, without discrete populations, many commonly estimated quantities in population genetics—migration rate ($m_{ij}$), effective population size ($N_e$), admixture proportion—are poorly defined. A long history of theoretical work describing geographical populations (e.g., Fisher 1937, Haldane 1948, Slatkin 1973, Nagylaki 1975, Sawyer 1976, Barton 1979) has not had that strong an effect on empirical work (but see Schoville et al. 2012). However, modern advances in acquiring both genomic and geospatial data are making it more possible—and necessary—to explicitly model geography. Today’s large genomic data sets are facilitating the estimation of the timing and extent of shared ancestry at a much finer geographic and temporal resolution than was previously possible (Li & Durbin 2011, Palamara et al. 2012, Harris & Nielsen 2013, Ralph & Coop 2013). And advances in theory in continuous space (Felsenstein 1975; Barton & Wilson 1995; Barton et al. 2002; Barton et al. 2010, 2013a,b; Hallatschek 2011), data sets of unprecedented geographical scale (e.g., Leslie et al. 2015, Aguillon et al. 2017, Shaffer et al. 2017), new computational tools for simulating spatial models (Haller & Messer 2018, Haller et al. 2019), and new statistical paradigms for modeling those data (Petkova et al. 2016; Ringbauer et al. 2017, 2018; Bradburd et al. 2018; Al-Asadi et al. 2019) are together bringing an understanding of the geographic distribution of genetic variation into reach.

The goals of this review are to frame some of the fundamental questions in spatial population genetics and to highlight ways that new data sets and statistical methods can provide novel insights into these questions. In the process, we hope to provide an introduction for empirical researchers to the field. We focus on a small number of foundational questions about the biology and history of the organisms we study: where they are, how they move, where their ancestors were, how those quantities have changed over time, and whether there are groups of them. We discuss each question as a spatial population genetic problem, explored using the spatial pedigree. It is our hope that by framing our discussion around the spatial pedigree, we can build intuition for—and spur new developments in—the study of spatial population genetics.

2. THE SPATIAL PEDIGREE

The spatial pedigree encodes parent-child relationships of all individuals in a population, both today and back through time, along with their geographic positions (Figure 1). If we had complete
Now

Parents

Grandparents

Great-grandparents

Longitude
Latitude

Figure 1
An example pedigree (a) of a focal individual sampled in the modern day, placed in its geographic context to make the spatial pedigree (b). Dashed lines denote matings, and solid lines denote parentage, with red hues for the maternal ancestors and blue hues for the paternal ancestors. In the spatial pedigree, each plane represents a sampled region in a discrete (nonoverlapping) generation, and each dot shows the birth location of an individual. The pedigree of the focal individual is highlighted back through time and across space.

knowledge of this vast pedigree, we would know many evolutionary quantities that we otherwise would have to estimate from data, such as the true relatedness between any pair of individuals, or each individual’s realized fitness. If we wanted to characterize the dispersal behavior of a species, we could simply count up the distances between offspring and their parents. Or, if we wanted to know whether a particular allele was selected for in a given environment, we could compare the mean fitness of all individuals in that environment with that allele to that of those without the allele. The location of an individual can be a slippery concept, depending on the biology of the species in question (as can the concept of individual). For instance, where in space does a barnacle hitchhiking on a whale live, or a huge, clonal stand of aspen? To reduce ambiguity, we will think of the location of an individual as the place it was born, hatched, or germinated.

In practice, we can never be certain of the exact spatial pedigree. Even if we could census every living individual in a species and establish the pedigree relationships between them all, we still would not know all relationships between—and locations of—individuals in previous, unsampled generations. And in a highly dispersive species, or a species with a highly dispersive gamete, there may be only a weak relationship between observed location and birth location. Despite these complexities, the idea of the spatial pedigree is useful as a conceptual framework for spatial population genetics. Although it may be difficult or impossible to directly infer the spatial pedigree, we can still catch glimpses of it through the window of today’s genomes.

2.1. Peeking at the Spatial Pedigree

Early empirical population genetic data sets were confined to investigations of a small number of individuals genotyped at a small number of loci in the genome. Many of the statistics measured from early data, such as \( \pi \), \( \theta \), or \( F_{ST} \), rely on averages of allele frequencies at a modest number of loci and thus reflect mostly deep structure at a few gene genealogies. These statistics are informative about the historical average of processes like gene flow acting over long timescales but are relatively uninformative about processes in the recent past or, more generally, about heterogeneity over time. However, the ability to genotype a significant proportion of a population and generate
whole genome sequence data for many individuals is opening the door to inference of the recent pedigree via the identification of close relatives in a sample. In turn, we can use this pedigree, along with geographic information from the sampled individuals, to learn about processes shaping the pedigree in the recent past.

2.2. Simulating Spatial Pedigrees

It has only recently become possible to simulate whole genomes of reasonably large populations evolving across continuous geographic landscapes. Below, we illustrate concepts with simulations produced using SLiM v3.1 (Haller & Messer 2018), which enables rapid simulation of complex geographic demographies. We recorded the genealogical history along entire chromosomes of entire simulated populations as a tree sequence (Haller et al. 2019). In most simulations, we retained all individuals alive at any time during the simulation in the tree sequence, allowing us to summarize not only genetic relatedness but also genealogical relatedness (e.g., to identify ancestors from whom an individual has not inherited any parts of their genome). We used tools from pyslim and tskit (Kelleher et al. 2018) to extract relatedness information and plotted the results in Python with matplotlib (Hunter 2007). Using SLiM’s spatial interaction capabilities, we maintained stable local population densities by increasing mortality rates of individuals in regions of high density, and we chose parameter values to generate relatively strong spatial structure (a dispersal distance of between 1 and 4 units of distance and a mean population density of 5 individuals per unit area). Our code is available at https://github.com/gbradburd/spgr. This new ability to not only simulate large spatial populations but also record entire chromosome-scale genealogies as well as spatial pedigrees has great promise for the field moving forward.

2.3. Estimating “Effective” Parameters from the Spatial Pedigree

On one hand, every single one of my ancestors going back billions of years has managed to figure it out. On the other hand, that’s the mother of all sampling biases.

—Randall Munroe, https://xkcd.com/674/

Inferences about the past based on those artifacts that have survived to the present are fraught with difficulties and caveats in any field, and population genetics is no exception. The ancestors of modern-day genomes are not an unbiased sample from the past population—they are biased toward individuals with more descendants today. It is obvious that we cannot learn anything from modern-day genomes about populations that died out entirely, but there are more subtle biases. For example, if we trace back the path along which a random individual today has inherited a random bit of genome, the chance that a particular individual living at some point in the past is an ancestor is proportional to the amount of genetic material that individual has contributed to the modern population. If we are looking one generation back, this is proportional to how many offspring they had, i.e., their fitness. If we are looking a long time in the past (in practice, more than 10 or 20 generations (Barton & Etheridge 2011]), this is their long-term fitness. Individuals from some point in the past who have many descendants today may or may not look like random samples from the population at the time. Different methods query different time periods of the pedigree and are thus subject to these effects to different degrees.

A familiar example of the limits of inference is the difference between census and effective population size (Wright 1931, Charlesworth 2009). More subtle is the difference between dispersal distance (σ) and effective dispersal distance (σ_e) (reviewed in Cayuela et al. 2018). Both are mean distances between parent and offspring: σ is the average observed from, say, the modern population, but σ_e is the mean distance between parent and offspring along a lineage back through the
spatial pedigree. The difference arises because lineages along which genetic material is passed do not give an unbiased sample of all dispersal events; they are weighted by their contribution to the population. For instance, on the one hand, if there is strong local intraspecific competition, individuals that dispersed farther from their parents (and their siblings) may have been more likely to reproduce and be ancestors of today’s samples, which would make $\sigma_e > \sigma$. On the other hand, if there is local adaptation, individuals may be dispersing to habitats dissimilar to that of their parents and are unlikely to transmit genetic material to the next generation because they are maladapted to the local environment, and so $\sigma_e < \sigma$ (for a review, see Wang & Bradburd 2014).

3. THINGS WE WANT TO KNOW

3.1. Where Are They?

Perhaps the first things we might use to describe a geographically distributed population are an estimate of the total population size and a map of population density across space. Quantifying abundance and how it varies can be crucial, e.g., for understanding what habitat to prioritize for conservation (Zipkin & Saunders 2018), which regions have higher population carrying capacities than others (Roughgarden 1974), or whether a particular habitat is a demographic source or sink (Pulliam 1988). The best measure of population density would be a count of the number of individuals occurring in various geographic regions, and the best measure of total population size would be the sum of the counts across all regions. How close can population genetics get us to estimating these quantities by indirect means? Some fundamental limitations are clear: for instance, if we only sample adults, we cannot hope to estimate the proportion of juveniles that die before adulthood. None of our sampled genetic material inherits from these unlucky individuals, so they are invisible as we move back through the pedigree. In addition, it is tricky to separate the inference of population density from that of dispersal, as the rate of dispersal to or from a region of interest governs whether it makes sense to consider the pedigree in that region on its own. Partly because of this issue, it is also difficult to estimate total size from a sample of a nonpanmictic population, and so it is more intuitive to discuss local density first, then scale up to the problem of total population size. The spatial pedigree contains a record of both total and regional population sizes, and we can use the genomes transmitted through it to learn about these quantities.

3.1.1. Population density. Genetic information about population sizes or densities most clearly comes from numbers and recency of shared ancestors, i.e., the rate at which coalescences between distinct lineages occur as one moves back through the pedigree. The relationship between population density and coalescence is, in principle, straightforward: With fewer possible ancestors, individuals’ common ancestors tend to be more recent.

Some calculations may be helpful to further build intuition (although the details of the calculations are not essential to what follows). Consider the problem of inferring the total size of the previous generation in a randomly mating population from the number of siblings found in a random sample of size $n$. There are $x(x-1)/2$ sibling pairs in a family with $x$ offspring, so the probability that, in a population of size $N$, a given pair of individuals are siblings in that family is $x(x-1)/N(N-1)$. Averaging across $N/2$ families (and assuming monogamy), the probability that a given pair of individuals are siblings (in any family) is therefore $E[X(X-1)]/2(N-1)$, where $X$ is the number of offspring in a randomly chosen family. This implies that in a sample of size $n$, $n(n-1)/4$ divided by the number of observed sibling pairs provides an estimate of effective population size, $N_e = N/E[X(X-1)]$. A similar argument is presented by Möhle & Sagitov (2003); also note this is inbreeding effective population size (see Wang et al. 2016 for a review of different types of effective population size). In other words, we should be able to use relatedness
to estimate population size up to the factor $E[X(X - 1)]$, which depends on demographic details. This same logic can be applied locally: The (inbreeding) effective population density near a location $x$, denoted $\rho_e(x)$, can be defined as the inverse of the rate of appearance of sibling pairs near $x$, per unit of geographic area and per generation (Barton et al. 2002).

We can extend this line of reasoning to see that the local density of close relatives should reflect local population size. The chance that two geographic neighbors actually are siblings decreases with the number of possible nearby parents: Wright (1946) quantified this with the neighborhood size, $N_{loc} = 4\pi \rho_e \sigma_e^2$, where $\pi$ is the mathematical constant, $\rho_e$ is the effective population density, and $\sigma_e$ is the effective dispersal distance. Each pair of siblings gives a small amount of information about population density within distance $\sigma_e$ of the pair. The more distant the coalescent event relating a pair of individuals, the vaguer the information they provide about population density, because there is greater uncertainty both in the inference of their precise degree of relatedness from genetic data, and in the location of their shared ancestors. Because genetic estimates of population size derive from rates of coalescence, they generally can only be used to estimate population sizes in the locations and times where those coalescences occur.

Relatives up to recent cousins can be identified in a large sample to varying degrees of certainty from genomic data (reviewed in Wang et al. 2016), using multilocus genotypes (Nomura 2008, Waples & Waples 2011, Wang 2014), shared rare alleles (Novembre & Slatkin 2009), or long shared haplotypes (e.g., Li et al. 2014).

An alternative approach to estimating local density is to measure the rate of genetic drift in an area. To do this, one compares allele frequencies in at least two samples collected in a region in different generations, either at different points in time or, in a species with overlapping generations, between contemporaneous individuals of different life stages, e.g., saplings and mature trees (Williamson & Slatkin 1999). The rate of drift is governed by, and therefore provides an estimate of, local population size; this is analogous to the variance effective size of a population (Ewens 2004, Charlesworth 2009, Wang et al. 2016). Because these approaches are pinned to the recent past, either by the generations sampled or by the degree of closeness of relatives, they should offer a more accurate estimate of recent population density than heterozygosity does, especially when density has changed over time.

3.1.2. Total population size. Estimates of the largest geographic scale—total population size—turn out to be sensitive to changes over the longest timescales. Total effective population size is often estimated using measures of genetic diversity such as $\pi$ (Nei & Li 1979, Tajima 1989), the mean density of nucleotide differences between two randomly sampled chromosomes. Because each such difference is the result of a mutation sometime since the chromosomes’ most recent common ancestor at that location on the genome, $\pi$ is equal to twice the mean time to most recent common ancestor within the sample, multiplied by the mutation rate. In other words, if the per-nucleotide mutation rate is $\mu$, and $2T$ is the mean length of a path through the pedigree along which two samples have inherited a random bit of genome from their most recent common ancestor, then $\pi \approx 2T\mu$. The quantity $T$ itself is an interesting summary of the pedigree and depends on geography, sampling, and dispersal in complex ways. What does it have to do with population size? In a randomly mating, diploid population of constant effective size $N_e$, the discussion above implies that $T = 2N_e$. In practice, populations are rarely constant, and so this is viewed as an average over the last few $N_e$ generations.

3.1.3. Heterozygosity and population density. We have seen that local population size is reflected in the pedigree through rates of shared ancestry: All else being equal, smaller populations
Figure 2

(a) Map of population density of the simulated population scenario we use for examples throughout. Populations have local density-dependent regulation within two valleys with high local carrying capacity (indicated by contours), separated and surrounded by areas with lower carrying capacity. The western valley has a carrying capacity about twice as large as that of the eastern valley. (b) Heterozygosity of a sample of 100 individuals across the range. Circle sizes are proportional to heterozygosity, and circle color indicates relative values (green, above the mean; white, mean; purple, below the mean). Low heterozygosities occur on the edges of the range, and because there is a strong barrier between the two valleys, mean heterozygosity is decreased by approximately 40% in the lower-density eastern valley.

should have more shared ancestry and therefore less overall genetic diversity. To gain intuition, we can look at how genetic diversity on the smallest scale—heterozygosity—behaves in a simulated example. Consider two adjacent valleys, one densely populated, the other sparsely populated, but both equal in geographic area and separated by a barrier to dispersal (Figure 2a). In general, we expect pairs of individuals in the low-density valley to have shared ancestors in the more recent past than pairs of individuals in the densely populated valley. This happens simply because each
group of siblings makes up a larger proportion of the more sparsely populated valley. Indeed, individuals in the sparsely populated valley have more shared ancestors in the more recent past, so the two chromosomes sampled in each individual from that valley are more closely related, i.e., have lower heterozygosity, as shown in Figure 2b.

Heterozygosity correlates with population density in this example with a strong barrier, but does it predict relative density more generally? Differences in heterozygosity can be shaped by other forces. For example, we might also expect a similar difference in heterozygosity if the two valleys had the same population density per unit area but one valley was larger than the other, because heterozygosity reflects population size. Or, if dispersal distances tend to be shorter in the more densely populated valley—e.g., because resources are more plentiful, so that individuals do not have to disperse as far to find food—there may be more inbreeding over short distances, despite higher genetic diversity within the valley as a whole. And, of course, modern population densities may not be well-predicted by historical densities.

Migration into and out of a region presents another complication. Roughly speaking, geographic patterns of relatedness are determined by a tension between population size and migration. As discussed above, genealogical relationships form on a timescale determined by population density, because in a geographic region containing N individuals, the chance that two individuals are siblings is of order 1/N. Shared ancestry—and thus genetic variation—is spatially autocorrelated to the extent that this process of coalescence brings lineages together more quickly than migration moves them apart as one traces individuals’ ancestry back through the pedigree. If the migration rate into a geographic region is much greater than 1/N—i.e., if more than a few individuals in each generation had parents living outside of the region—the pedigree in a region cannot be considered autonomously. In other words, regions cannot be analyzed independently unless a region of interest sees less than one migrant per generation [i.e., outside the parameter regime of the structured coalescent (Nagylaki 1998)].

3.2. How Do They Move?

Dispersal is fundamental to many questions in ecology and evolution, such as predicting the spread of pathogens (Biek & Real 2010), responses to climate change (Parmesan 2006), or population demography and dynamics (Schreiber 2010). We use the term “dispersal” to refer to the process that governs the displacement between geographic locations of parent and offspring. For some species, it may be feasible to directly measure dispersal using, e.g., telemetry or banding (Cayuela et al. 2018), but this is often prohibitively expensive or difficult. Although we usually do not know the spatial locations of ancestors in the pedigree, the spatial locations of modern individuals can give strong clues about this.

3.2.1. Dispersal (individual, diffusive movement; σ). We call the absolute value of the spatial displacement between parent and offspring locations the “dispersal distance” and refer to it as σ. The dispersal distance describes the quantity one could calculate by, e.g., collecting a large number of germinating seeds and taking the square root of the average squared distance from those seeds to their parents. It determines how quickly spatial populations mix: Larger dispersal distance spreads relatedness, and thus genetic variation, throughout the species’s range faster.

Dispersal distance could be trivially estimated by taking the average of these displacements across history if we knew the spatial pedigree through time without error. In practice, we often are only able to directly observe the location of individuals in the present day. However, spatial distances between individuals of different levels of relatedness give us information about dispersal distance, even without parental locations. For example, the distance between siblings is the result of
two dispersals, so an average over many pairs of siblings of this distance squared gives an estimate of \(2\sigma^2\). The mean squared displacement along a path of total length \(n\) through the pedigree is \(\sigma^2 n\): First cousins would give four times the squared dispersal distance, and so forth. This spatial clustering of close relatives is shown in Figure 3a. Unfortunately, basing an estimator of \(\sigma\) on this observation is usually infeasible in practice, unless a large, unbiased fraction of the population has been genotyped (e.g., Aguillon et al. 2017).

Empirical data sets do not usually contain many close relatives, but using phylogenetic methods on nonrecombining loci (e.g., the mitochondria), we can obtain good estimates of times since common ancestor between much more distantly related individuals. This general idea was put into practice by Neigel & Avise (1993), who used the spatial locations of samples and their mitochondrial tree to infer \(\sigma\) using phylogenetic comparative methods. However, the relationship between age of common ancestor and distance apart clearly can only apply if the geographic distance it predicts is much less than the width of the species range. In many species this is not true, because the typical time to common ancestor between two sampled individuals is long enough for their lineages to have crossed the species range several times (Barton & Wilson 1995).

Instead of identifying close relatives, we might use the decay of mean genetic relatedness with distance (shown in Figure 3b) to learn about dispersal. Empirical studies usually find that the mean time since most recent common ancestor of two individuals at a random point on the genome is larger for more distant individuals (Epperson 2003, Charlesworth et al. 2003, Sexton et al. 2013), and this relationship of isolation by distance clearly should contain some information about \(\sigma\). An approximate formula for this relationship in homogeneous space (the Wright-Malécot formula) can be derived in various ways (Malécot 1948, Sawyer 1976, Barton & Wilson 1995, Rouset 1997, Barton et al. 2002, Robledo-Arnuncio & Rouset 2010, Ringbauer et al. 2017, Al-Asadi et al. 2019), but is composed of two parts: (a) how long it has been since the two individuals last had ancestors that lived near to each other, and (b) how long before that they actually shared an ancestor. The second part is not well understood, but Malécot (1948) assumed that the second stage acts as in a randomly mating population with some local effective size, \(N_{loc}\). Rouset (1997) used this formula to show that the slope of \(F_{ST}\) against the logarithm of distance is, under these assumptions, an estimator for Wright’s neighborhood size \(4\pi\rho_\sigma^2\). Readers are directed to Barton & Wilson (1995) for more discussion of these methods.

### 3.2.2. Dispersal on heterogeneous landscapes

Of course, organisms do not live on featureless toroids, and individuals in one part of a species’s range may disperse more than those in another due to heterogeneity in their nearby landscape. In spatial population genetics, one common way to describe this heterogeneity is with a spatial map of resistance. The resistance value of a portion of the map is inversely proportional to the speed at which organisms tend to disperse, or (equivalently) the distance they tend to travel, when in that area. These models were introduced to spatial population genetics by McRae (McRae 2006, McRae & Beier 2007, McRae et al. 2008) and represented an important step forward in modeling dispersal (over, e.g., least-cost path analysis). However, resistance models do not allow spatial differences in dispersal variance or asymmetry in gene flow, as would be seen up a fecundity gradient or following a bias in offspring dispersal downwind. Note, furthermore, that we are distinguishing the resistance model of organism movement from the (current) methods used to fit such models; as implemented, resistance methods make use of an approximation that can result in estimates that diverge widely from the truth, especially if gene flow is asymmetric (Lundgren & Ralph 2019).

One of the most promising current spatial methods is MAPS (Migration And Population-size Surfaces) (Al-Asadi et al. 2019), which uses information about long shared haplotype tracts (recent ancestry) to build a map of both local population size and dispersal ability. MAPS builds on
(Caption appears on following page)
3.2.3. Dispersal flux. We would often like to measure the amount of gene flow across some barrier or between two geographic regions. To do this, we can ask for the number of dispersal events in a given year that cross a given line drawn on a map (e.g., the boundary between two adjacent regions)—that is, how many individuals are born each year whose parents were born on the other side of the line? This quantity, which we call the dispersal flux across the boundary, is useful for investigating connectivity between different areas and the role of the landscape in shaping patterns of dispersal. Dispersal flux is a geographical analog of the migration rate between discrete, isolated populations. Note that dispersal flux is not constrained to be equal in both directions across a boundary: It may be easier to disperse, e.g., down-current than up-current. For instance, Figure 3c shows the number of dispersal events crossing two lines on our simulated valley landscape over a few generations. The line on the left goes through the denser valley and so has a large flux in each direction. The line on the right goes along the edge of the sparser valley, and so has lower flux overall, and more east-to-west flux (more red than blue arrows crossing it) due to the higher population density to the east of the line.

As before, if we had complete knowledge of the spatial pedigree, we could calculate the flux across any line simply by counting the number of parent-offspring pairs that span it. However, it is rare to find any one individual that is a direct genetic ancestor of another in a data set (apart from perhaps some parent-child relationships). As a result, we almost never infer a direct chain of dispersal events, only dispersal since coalescence events. The likely locations of these coalescent events depends on the maps of population density and fecundity, thus entangling the problems of inferring dispersal flux and population density.

For instance, across a barrier with low flux, the average time to coalescence for pairs of individuals on opposite sides of the line is longer than that between individuals on the same side (Bradburd et al. 2013, Dufoiret-Frebourg & Blum 2014, Ringbauer et al. 2018). Flux across a line is also related to mean dispersal distance, $\sigma$, discussed above. The expected flux across a straight boundary in a homogeneous area, per unit time and per unit of boundary length, is equal to the birth rate per unit area multiplied by $2\sigma/\pi$ [where $\pi$ is the mathematical constant; this is “Buffon’s Needle” (Ramaley 1969)], so estimates of flux can be converted to estimates of dispersal distance.
3.3. Where Were Their Ancestors?

Above, we have discussed how population dynamics of movement and reproduction are reflected in the spatial pedigree. A complementary view is to start with individuals sampled in the present day, and then, looking backward in time, to ask where in space they have inherited their genomes from, i.e., where their ancestors lived. Because the signal in genetic data comes from shared ancestry and coalescence in the pedigree, thinking from this reverse-time perspective is useful in interpreting genetic data. However, summaries of how ancestry spreads across space through the pedigree can be important in their own right. For example, many humans are interested in the locations of their own ancestors back in time. More generally, we are often also interested in the partitioning of genetic diversity across space, e.g., quantifying the strength of inbreeding locally, or assessing the capacity of a population to become locally adapted. Spatial distributions of ancestry compared across sections of the genome could also be informative about other processes, such as the origins and mechanisms of selective sweeps.

3.3.1. Geographic distribution of ancestry. As one looks further back in time, the geographic distribution of an individual’s ancestors is determined by the history of population size changes and movements. At any point in the past, each portion of each individual’s genome can be associated with the ancestor from whom they inherited it. Their genetic ancestry can therefore be apportioned across space according to the locations of the ancestors from whom they inherited their genome. There are a number of natural questions that arise from a description of this process. How much of an average individual’s ancestry is still within a given region, at some point in the past? How long in the past was it since a typical individual in one location had an ancestor in another geographic region (say, the next valley over)? How far back in time do you have to go before the average proportion of ancestry that an individual inherits from the local region they live in drops below (say) 90%?

This spatial distribution of all genetic ancestors is depicted at several points back in time for four individuals in a simulation in Figure 4. As we look further back in time, we can see the cloud of ancestry spread out across space. The spread is diffusive, i.e., the mean displacement of a given nth-generation ancestor is of order $\sigma_n \sqrt{n}$, as long as this is small relative to the width of the region. Not shown on the map are the many genealogical ancestors from which the focal individuals have not inherited genome—these spread at constant speed rather than diffusively (Kelleher et al. 2016b), so that even by 15 generations in the past, each focal individual is genealogically descended from nearly everyone alive, even those living in the other valley (Chang 1999).

Although Figure 4 does not highlight ancestors shared by any pair of the focal individuals, it is clear that these occur in the points of overlap between their two clouds of ancestry. These shared ancestors may be informative about relative population densities in the two valleys.

3.3.2. Relative genetic differentiation. In the two-valley simulation of Figure 4, ancestors are shared within valleys over the time period shown, but not between valleys. The difference in the geographic distribution of shared ancestors results in higher relatedness between the pairs of individuals from the same valley, and it underlies all measurements of relative genetic differentiation. The most common measure of how much more closely related neighbors are to each other relative to the population as a whole is $F_{ST}$ (Wright 1951), which can be thought of as estimating one minus the ratio of mean coalescence time within subpopulations to that within the entire population (Slatkin 1991). The normalization by mean overall coalescence time makes this measure relative, and therefore comparable across populations of different sizes.

In stable populations, relative genetic differentiation is determined by the tension between geographic mixing (due to dispersal flux between areas) and local coalescence (determined by...
Figure 4
Spatial locations of the ancestors of four individuals at (a) 1, (b) 15, and (c) 75 generations in the past. All individuals from whom the four individuals have inherited genome are denoted by a circle, with one color for each individual, and with circle area proportional to the amount of genome inherited. The landscape is the same as in Figure 3. Since the simulation uses overlapping generations, ancestors of different degrees are present at the same time.
population density). In the more remote past, each individual’s ancestors are distributed across all of geography, so that the long-ago portion of the pedigree looks like that of a randomly mating population. Tracing the pedigree backward in time, remote ancestors have “forgotten” the geographic position of their modern-day descendants. The transition between these two phases—recent spatial autocorrelation (scattering) and distant random mating (collecting) (Wakeley 1999, Wilkins 2004)—happens on the timescale over which a lineage crosses the species range (Wakeley 1999). Since lineage motion is diffusive, in a continuous range of width \(L\), coalescences between lineages of distant individuals happen on a timescale of \(L^2\) generations. Relative genetic differentiation is determined by the likelihood that coalescence between two nearby individuals occurs during the scattering phase—i.e., by the length of time that two spatial clouds of ancestry of Figure 4 are correlated with each other, relative to the local effective population size. So, in a species with limited dispersal (i.e., small \(\sigma\)), \(F_{ST}\) may not be the best measure of this quantity—others are possible that depend less on the deep history of the population. For instance, one could ask for the proportion of the genomes of two neighbors inherited in common from ancestors in the last \(T\) generations relative to the same quantity for distant individuals, where \(T\) is chosen to reflect the timescale of interest. The appropriate measure will depend on the application at hand.

3.4. Clustering, Grouping, and Admixture

Many tools for modern analysis of genetic data partition existing genetic variation into clusters labeled as (discrete) populations (e.g., Pritchard et al. 2000, Alexander et al. 2009). The ubiquity of gene flow means that any conceptual model of discrete populations must also include admixture between them. Models of discrete population structure and admixture provide visualizations that can be helpful for describing and making sense of patterns of relatedness and genetic variation. Coupled with a quantification of relative genetic differentiation, they can be used in conservation to delineate discrete management units. However, in practice, these discrete groups may in fact be determined by clustering of sampling effort rather than any intrinsic divisions within the population. Explicitly incorporating the geographic proximity of samples can help alleviate these concerns (Bradburd et al. 2016, 2018), but it would still be desirable to delineate groups in a way explicitly linked to the actual history of relatedness, i.e., the spatial pedigree.

The real-world situation in which admixture between discrete populations is perhaps most clearly defined is when populations that have been completely separated for a long period of time come back into contact. In this case, the discrete populations are defined by partitioning ancestral individuals, and admixture of subsequent generations is determined by inheritance from them. For instance, suppose that the two valleys of our running example were isolated during a long period of glaciation, followed by glacial retreat and expansion into secondary contact. The valleys clearly act as discrete populations during glaciation, during which there are few if any pedigree connections between the valleys. Pedigrees only start to overlap more recently between individuals in each valley. To obtain an admixture proportion, we can fix some reference time in the past and label the genomes of each individual in subsequent generations according to which of the two valleys it inherited from at the reference time. This is shown in Figure 5a. Eight generations after contact, this quantity clearly reflects the history of isolation; individuals in or near each valley have inherited most of their genomes from ancestors in the closest valley. However, if we return 75 generations later, we see a more complicated story (Figure 5b). The continued spatial mixing has resulted in a significant portion of ancestry in the western valley inherited from glacial-era ancestors in the eastern valley, and vice versa. Further complicating the issue, the west-to-east flux across the ridge between the valleys is higher than that in the opposite direction because
Figure 5
Admixture proportions after a recent secondary contact. Each pie shows the proportion of a diploid individual's genomes that are inherited from the western (red) and eastern (blue) valleys, respectively. The plots show the geographic distribution of that ancestry (a) 8 generations, and (b) 75 generations after a barrier along the top of the ridge is removed.

the western valley has a higher population density. As a result, a much higher proportion of the genomes in the eastern valley are now inherited from individuals that, during glaciation, were found in the western valley.

This simulation highlights both the relevance of a model of discrete population structure and admixture to spatially continuous data, and also its limitations. Outside the time during and immediately following glaciation, describing genotyped individuals on this landscape using a model of discrete populations with admixture becomes less helpful. Without a concrete interpretation based in the population pedigree, partitioning genetic variation into discrete groups and labeling samples as admixed between those groups can encourage the incorrect interpretation that discrete populations are Platonic ideals, real and unchanging.
3.5. How Have Things Changed over Time?

As we have discussed above, different methods are sensitive to signal from different time slices of the pedigree. Of course, the geographic distributions and dispersal patterns of most species, not to mention the landscape itself, have changed over the timescale spanned by the pedigree of modern individuals. For example, high-latitude tree populations may only have been established following the retreat of the glaciers some tens of generations ago (Whitlock & McCauley 1999), and invasive species may have entered their current environment even more recently than that. In addition to organisms moving themselves across landscapes, landscapes are often changing out from under organisms’ feet. Anthropogenic land use and global climate change are radically altering both where organisms can live and where they disperse (Parmesan et al. 1999). The result is that many empirical systems are not in equilibrium, so that the dynamics we estimate from measurements of extant individuals may not be representative of those in the past. Exacerbating this problem, it can take a long time to reach equilibrium in areas of high population density or across regions of low flux (Crow & Aoki 1984, Whitlock 1992, Slatkin 1993, Whitlock & McCauley 1999). Theory and methods that can address change across both time and space are still lacking, and they present an exciting set of challenges for the field.

3.5.1. Difficult discontinuous demographies. The specter of spatial discontinuity in the pedigree—those situations in which none of the individuals living in a particular region at a particular point in time are descended from any of the individuals who had inhabited that region at some point in the past—haunts the inference of historical processes from modern genetic data. Spatial discontinuities in the pedigree occur when regions become unoccupied (e.g., due to ecological disturbance or agonistic interactions) and are recolonized by immigrants from elsewhere. Discontinuity is expected over long enough timescales that ranges have shifted, and vicariance–secondary-contact events have occurred. There is abundant evidence of large-scale population movements or changes in human data sets that include ancient or historical samples (Skoglund et al. 2014, Pickrell & Reich 2014, Lazaridis et al. 2014, van den Haak et al. 2015, Joseph & Pe’er 2018), as well as in other species over recent timescales using museum specimens (Bi et al. 2013, 2019). Spatial discontinuity can pose a problem if the models we use for inference are predicated on the assumption that the observed geographic distribution of genetic variation was generated in the sampled geographic context. In principle, we can include large-scale population movement in our inference methods, but only if we have some clue that it has occurred. Identifying lack of fit due to such historical events remains an open problem. As we develop more data sets that include genotype data from historical individuals, we will better be able to evaluate the prevalence of spatial discontinuity and its effects on our inferences across natural systems.

3.5.2. Genealogical strata. The field of population genetics has many heuristics for genetic data; different types of genetic information are informative about events that occurred at different times in a population’s history. For example, patterns of sharing of rare alleles can be informative about events in the recent past, and long shared haplotypes even more so. In contrast, pairwise statistics like $\pi$ or $F_{ST}$ mostly inform us about events long ago, on a scale of $N_e$ generations. Different statistics thus provide us with windows into different strata of the pedigree. Rates of sibling- and cousin-ship would shed light on the last few generations of history, but carry the (in many cases unreasonable) requirement of a sample that is a sizable fraction of the entire population. Methods for inferring the distant history of populations using a few samples are the most well-developed (e.g., Gutenkunst et al. 2009, Li & Durbin 2011, Kamm et al. 2017), but these can incorporate very
little in terms of geographic realism, and indeed much of the signal of geography may have washed out on this coalescent timescale (Wilkins 2004). The length spectrum of long, shared haplotypes—also known as identical-by-descent segments—can be used to peer into the strata of the pedigree between these two timescales. The lengths of segments of genome inherited in common by two individuals from \( t \) generations ago scales roughly with \( 1/t \), so these can carry substantial information about ancestors from only tens of generations ago. This fact has been used by Al-Asadi et al. (2019) to create estimated maps of population density and migration rate for different, recent periods of time. However, the different strata cannot be so cleanly disentangled—e.g., although 1-cM-long shared haplotypes are older on average than 2-cM-long ones, the distributions of ages of these shared haplotypes overlap considerably. Moreover, the correspondence between shared haplotype length and age of common ancestor depends strongly on population size history (Ralph & Coop 2013).

4. NEXT STEPS FOR SPATIAL POPULATION GENETICS

In this review we have focused on demographic inference—seeking to learn about past or present demographic processes from population genetic data. As demography—birth, reproduction, death—is the mechanism by which natural selection occurs, it is fundamental not only to history and ecology but also to evolutionary biology. Describing the geographic distribution of genetic variation—its patterns, the processes that have generated and maintained it, and its consequences—is a fundamental goal of the field. We have already sketched out plenty of work for the field—quantitative inference with continuous geography is for the most part an unsolved problem. Nonetheless, we describe below a few more areas in which we hope or expect further progress in the field to be made.

4.1. Selection

Some of the most important aspects of geographical population genetics we have not reviewed are related to natural selection. A significant avenue for future work is to develop methods that jointly estimate selection and demography for spatially continuous populations. We have mostly ignored this above, but clearly local adaptation and natural selection impact the structure of the pedigree, including its spatial patterns. For example, after a hard selective sweep, everyone in the population has inherited a single beneficial allele, and so the lineages of all extant individuals at that locus will pass through the lucky individual in whom that mutation occurred. There is a great deal known about the expected geographical action of selection, e.g., how beneficial alleles spread across space (Fisher 1937, Ralph & Coop 2010, Hallatschek & Fisher 2014), how locally adaptive alleles can be maintained in the face of gene flow (Slatkin 1973, Kruuk et al. 1999, Ralph & Coop 2015), or how hybrid zones are structured (Barton & Hewitt 1985, Sedghifar et al. 2015). In all these situations, selection can produce dramatically different patterns in the genealogies of different regions of the genome. Incorporating selection into spatially continuous models will facilitate a greater union of population genetic processes and ecological models of population dynamics, and it will help shed light on important evolutionary questions.

4.2. Simulation and Inference

Spatial population genetics suffers from a lack of concrete mathematical results that can be used for inference. Although we have a good understanding of the decay of expected pairwise relatedness with distance (e.g., the Wright–Malécot formula), a deeper understanding of the coalescent process is required to make quantitative predictions of more complex population genetic statistics.
How can we fill this gap? A recent advance in the field, which may help improve inference models with and without selection, has been the advent of powerful and flexible simulation methods, such as SLiM (Haller & Messer 2018, Haller et al. 2019) and msprime (Kelleher et al. 2016a). These methods allow researchers to produce genome-scale data sets from simulations of arbitrary complexity. This capability offers an excellent pedagogical tool and opportunity to build intuition for patterns of genetic variation in complex scenarios for which theory may be lacking. In addition, these simulation methods can be used in statistical inference. For example, simulated data sets can be used in the training of machine learning methods (e.g., Schrider & Kern 2018), or to compare between different generative models using approximate Bayesian computation (Marjoram & Tavaré 2006). Simulation-based inference presents an exciting way forward for problems that may be analytically intractable, although for large problems, the required computational effort is daunting. As an intermediate solution, a promising class of spatial coalescent models allows rapid simulation with geographically explicit models (Barton et al. 2010, Guindon et al. 2016), although the precise relation to forward-time demographic models is not well understood.

4.3. Identifiability and Ill-Posedness

Another important gap in current methods for spatial population genetics inference is a quantitative understanding of the limits of our ability to learn about quantities of interest. Demographic inference with population genetics data is an example of an inverse problem—the mapping from parameters to data is relatively well-understood, but the inverse map is not. Many inverse problems are ill posed, meaning that many distinct models give rise to statistically indistinguishable data sets, i.e., there is some degree of statistical nonidentifiability (Petrov & Sizikov 2005, Stuart 2010). For example, as with estimating effective population size using the site frequency spectrum (Myers et al. 2008), we may never be able to infer rapid changes in effective population density (although see also Bhaskar & Song 2014). Likewise, the timing of heterogeneity in flux across a particular part of the landscape may be difficult to disentangle from the magnitude of its change, as is the case with migration/isolation models of dynamics in discrete demes (Sousa et al. 2011). More generally, it may be difficult to learn about local biases in the direction of flux, or large-scale flow in the average direction of dispersal, without genetic data from historical individuals. Mathematical theory and simulation-based tests of inference methods should allow us to further determine the spatial and temporal resolutions with which we can view history.

4.4. The Past

The growing availability of genomic data from historical or ancient samples, collected from archaeological sites or museums, is one of the most exciting developments in population genetics and has the potential to revolutionize the field of spatial population genetics. These historical data can provide a kind of fossil calibration, helping to anchor spatial pedigrees in space and time. Although it may be unlikely that any particular genotyped historical individual is the direct genetic ancestor of any sampled extant individual, the geographic position of the historical sample and its relatedness to modern individuals is nonetheless informative about the geography of any modern individual’s ancestors. These historical data can also illuminate temporal heterogeneity in some of the processes described in Section 3.4, which might otherwise be invisible in data sets composed of only modern individuals.

5. CONCLUSION

There is a long history of empirical research describing spatial patterns in genetic variation (Dobzhansky 1947, Mourant 1954, Schoville et al. 2012). Most population genetics inference
methods assume a small number of randomly mating populations, and it is still unknown what
effect realistic geographies have on these inferences. This is starting to change, however, as larger
data sets with vastly more genomics data become available, giving us power to bring more of
a population’s pedigrees into focus. In turn, the ability to study pedigrees and their geography over
the recent past is bringing about a union between the fields of population genetics and population
ecology. Measures of dispersal and density based on the pedigree can also inform the study of the
factors governing the distribution and abundance of organisms over short timescales that might
have previously been out of reach for population genetics. As data sets, theory, and methods in
spatial population genetics continue to progress, it will be exciting to explore further integration
of the two disciplines.

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