The Evolution of Comparative Phylogeography: Putting the Geography (and More) into Comparative Population Genomics

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Abstract

Comparative population genomics is an ascendant field using genomic comparisons between species to draw inferences about forces regulating genetic variation. Comparative phylogeography, by contrast, focuses on the shared lineage histories of species codistributed geographically and is decidedly organismal in perspective. Comparative phylogeography is approximately 35 years old, and, by some metrics, is showing signs of reduced growth. Here, we contrast the goals and methods of comparative population genomics and comparative phylogeography and argue that comparative phylogeography offers an important perspective on evolutionary history that succeeds in integrating genomics with landscape evolution in ways that complement the suprageographic perspective of comparative population genomics. Focusing primarily on terrestrial vertebrates, we review the history of comparative phylogeography, its milestones and ongoing conceptual innovations, its increasingly global focus, and its status as a bridge between landscape genomics and the process of speciation. We also argue that, as a science with a strong “sense of place,” comparative phylogeography offers abundant “place-based” educational opportunities with its focus on geography, prominent landscape features, and Earth history. We conclude that it nonetheless plays an important role in grounding our interpretation of genetic variation in the fundamentals of geography and Earth history.

Key words: Gott Earth projection, whole-genome sequencing, landscape genomics, place-based education, indigenous knowledge.

Significance

Comparative phylogeography shares many concepts and techniques with comparative population genomics, yet differs in its emphasis on geography, codistribution of species, and genetic structure within species. We review the history of comparative phylogeography and its global expansion in the last 12 years and suggest that it complements comparative population genomics in its ability to engage students, indigenous groups and the public via its focus on geography, prominent landscape features, and Earth history.

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**Introduction**

Comparative population genomics is an emerging field that leverages variation in genomes of multiple species to understand the origins of genomic diversity and natural selection and to make inferences about patterns of divergence between closely related species. By comparing genomic variation across multiple species, comparative population genomics aims to discover common forces acting on homologous or divergent regions of the genome. Comparative population genomics approaches have been used with great success since the 1970s, but with the rise of whole-genome sequencing (WGS) in recent years, the approach has gained renewed vigor and detail. In its most recent incarnation, the emphasis in comparative population genomics is less on the details of geography of each species, and more on sampling each species sufficiently so as to capture major patterns of variation and identify the demographic and genetic forces that have shaped them. Appropriate for its goals, the focus of modern comparative genomics is often on genetics, rather than geography.

Here we argue that comparative phylogeography—in many ways a more mature discipline than comparative population genomics, but one arguably with less ongoing conceptual innovation—stands as an important counterpoint to and partner of comparative population genomics. In so far as comparative population genomics suffers from a lack of a rigorous geographic perspective, and often seeks to marginalize over geography (see Comparative Phylogeography as Place-Based Evolutionary Biology section), comparative phylogeography brings an explicit geographic perspective that has not only revealed details of species history, but can complement the goals of comparative population genomics. Comparative phylogeography seeks to understand biotic history by finding landscape features—mountains, rivers, transition zones—that create vicariant breaks in genetic variation across a suite of species. A major emphasis in comparative phylogeography is to find drivers of genomic splits shared across suites of codistributed species, and thereby find landscape features that ultimately explain patterns of species occurrence, community composition, and genetic diversity (table 1). A related discipline, landscape genomics, usually focuses on a smaller spatial scale and emphasizes genomic adaptation to landscape and climate features (Manel et al. 2003; Manel and Holderegger 2013; Rissler 2016). The focus of comparative phylogeography is usually on organismal histories, rather than on the diversity of evolutionary forces shaping the genome (table 1). To understand the relationships between comparative phylogeography and comparative population genomics, we first ask whether comparative phylogeography is a viable field of inquiry and on scientific growth trajectory. We briefly review the historical origins and recent milestones of comparative phylogeography from our perspective as empiricists working on terrestrial vertebrates. We then update our understanding of the global distribution of phylogeographic studies and argue that, as a “place-based” discipline, comparative phylogeography has great appeal for local and regional scientific communities and public education around the world in ways that are challenging for comparative population genomics. We conclude that, although it has yet to fully embrace the genomics revolution, comparative phylogeography is still a vibrant discipline that stands to make important contributions to evolutionary biology and in particular can leverage its “sense of place” to help include diverse communities in the research enterprise around the globe.

**Is Comparative Phylogeography Dead?**

A Brief History of Comparative Phylogeography

Early in its evolution, phylogeography adopted a comparative perspective, seeking to find common patterns across codistributed species. Although the first phylogeographic studies were conducted on single species (Avise et al. 1979), very quickly the field adopted a comparative approach, motivated...
to identify patterns of intraspecific genetic variation shared across species and so point to common historical processes (Bermingham and Avise 1986; Avise et al. 1987). At the same time, using morphology and other cladistic characters, the field of vicariance biogeography had begun to decipher shared geographic patterns of speciation among closely related and codistributed taxa with phylogenetic trees (Cracraft 1982, 1986). Although the problems addressed in early comparative phylogeographic studies by definition adopted molecular tools, they were often inspired by morphological studies of vicariance biogeography and cladistic analyses, often on the same organismal lineages (e.g., Cracraft 1986; Edwards 1993; Dolman and Joseph 2012; Cracraft 1991; Nyari and Joseph 2013). Indeed, phylogeography was envisioned then as the bridge between population genetics and vicariance-centered phylogenetic systematics, a perspective that remains true today but which has also been expanded in modern parlance into a bridge between the divergent scales of landscape genetics and processes of speciation.

A literal focus on gene trees in space defined phylogeography from the outset and distinguished it from prior analyses of allele frequency variation, as revealed by blood types Edwards and Cavalli-Sforza 1963; Cavalli-Sforza and Edwards 1967; Bodmer 2015) or multilocus allozymes (Allendorf 2017). Using allozyme electrophoresis, researchers had accumulated hundreds of studies documenting the geographic pattern of genetic variation within species. Allozyme electrophoresis proved to be a useful tool to quantify levels of variation in protein-coding regions whose alleles could be differentiated by electric charge. Studies of allozyme variation were relatively inexpensive and were usually couched in terms of quantifying geographic variation or investigating spatial variation in selection and clines, and eventually developed into quantitative tests of modes of speciation. Many studies employing allozymes, as well as more recent studies employing microsatellites or single-nucleotide polymorphisms (SNPs), focused on the geographic distribution of alleles and models of isolation by distance that provided a sensitive framework for interpreting patterns of geographic variation (Slatkin 1985; Novembre and Slatkin 2009; François et al. 2010). There was also considerable interest in the ecological correlates and life history predictors of genetic variation—how the ecology and behavior of organisms modulated levels of genetic variation. Although phylogenetic trees were routinely made from allozyme data, the emphasis was on variation in allele frequencies: For example, the commonly used statistic \( F_{st} \) was largely used to estimate levels of gene flow among populations; only occasionally was \( F_{st} \) used to attempt to detect natural selection, primarily because the number of loci routinely surveyed in allozyme studies (anywhere from \( \sim 20 \) to \( \sim 60 \) loci) was too small to detect \( F_{st} \) outliers.

Restriction enzyme analysis was introduced to population genetics in the mid-1970s and fundamentally changed the way empiricists viewed genetic variation; whereas in the allozyme era, alleles were distinguished categorically, as similar or different to one another. By contrast, restriction fragment length polymorphisms (RFLPs) allowed researchers to quantify the degree of difference between alleles. Perhaps more fundamentally, RFLPs permitted the phylogenetic analysis of alleles, first deployed on a large scale with animal mitochondrial DNA (mtDNA). The first intraspecific “gene tree” was published by Avise et al. (1979). This rather understated publication ushered in a dramatically new view of genetic variation, one in which genealogies of alleles, within and between species, rather than allele frequencies, were the mode of description for genetic variation. Because of its genetic simplicity, rapid evolution, and abundance in animal cells, mtDNA provided easy access to genetic variation and geographic patterns in animals. In time, massive gene trees consisting of hundreds of alleles, allowed fine dissection of geographic patterns within species, such as the human species (Cann et al. 1987). A plethora of statistics of genetic variation, such as nucleotide diversity (\( \pi \)) and \( F_{st} \) incorporating divergence among haplotypes, arose in the context of RFLP variation (Nei and Li 1979; Allendorf 2017); many are still in use today.

**Milestones and Recent Innovations in Comparative Phylogeography**

The introduction of the polymerase chain reaction to phylogeography in the early 1980s, to both nuclear and mitochondrial genes, finally allowed direct visualisation at the nucleotide level to the DNA changes underlying RFLPs; although phylogeographers still struggled then with the apparently low levels of variation detectable at nuclear genes and the challenge of recombination within loci, we suggest that the modern form of phylogeography, in which sequence changes at multiple loci are analyzed in allele-frequency and phylogenetic frameworks, was established by the early 1990s. With the establishment of phylogeography as a discipline, it was only a matter of time before researchers endeavored to compare phylogeographic histories of multiple species. Often phylogeographic comparisons between species were conducted in the context of codistributed species that may have responded similarly to the same physical or environmental barriers on the landscape. Indeed, codistribution of species under study is a central tenet of comparative phylogeography and is often included in its definition (Gutiérrez-García and Vázquez-Domínguez 2011). In many ways, comparative phylogeography refocused attention to the geographic context of genetic variation in ways that the earlier focus on single species did not, or could only do so haphazardly. By analyzing the genetic diversity of codistributed species at key transition zones around the globe, comparative phylogeography allowed the identification of zones of phylogeographic divergence and promoted the idea that landscape features, such as mountains, rivers, and marine archipelagoes, could serve as
powerful generators of biodiversity and genetic divergence across multiple lineages. A special issue of *Molecular Ecology* in 1998 focused on comparative phylogeography. Bermingham and Moritz (1998) celebrated 10 years of achievement since Avise's seminal 1987 review, but they bemoaned the overreliance on the single locus provided by animal mtDNA.

The 2000s brought continued expansion of frameworks and methods in comparative phylogeography. Beheregaray (2008) noted spectacular growth of phylogeography and comparative phylogeography in the ensuing decade, with concomitant diversification in molecular markers used, by then routinely including nuclear SNPs and microsatellites (Garrick et al. 2015). However, Beheregaray also noted a gross paucity of studies from the Southern Hemisphere and warned of making general conclusions of process without more equal representation of the hemispheres. Hickerson et al. (2010) documented further expansion of the molecular marker toolkit and pointed toward the integration of ecological niche models, community assembly perspectives, and model-based inference of demographic histories into comparative phylogeography. The incorporation of techniques such as approximate Bayesian computation (ABC) and coalescent simulations flowed naturally from the increased distinction by practitioners between gene histories and population histories, and the focus, appropriately, on the population history as the end-goal of phylogeography. Hierarchical approximate Bayesian computation represents an important statistical approach specifically tailored to comparative phylogeography and the detection of concerted demographic responses to environmental events. Increasingly, estimates of demographic history are integrated with paleospecies distribution models (Carstens and Richards 2007; Alvarado-Serrano and Knowles 2014). Additional recent innovations in comparative phylogeography include the use of machine learning to identify comparative phylogeographic trends across clades (Espíndola et al. 2016; Carstens et al. 2018; Barrow et al. 2020; Fonseca et al. 2021). Finally, the use of phenotypic traits to guide phylogeographic sampling and erect hypotheses of intraspecific differentiation and interspecific codiversification have been proposed (Papadopoulou and Knowles 2016; Sullivan et al. 2019). Many of these innovations were celebrated in a 2016 symposium on comparative phylogeography organized by John Avise, Brian Bowen, and Francisco Ayala for the National Academy of Sciences (Avise et al. 2016). These tools have yet to be widely applied but suggest multiple promising avenues for future research and help promote a predictive, rather than descriptive, framework for comparative phylogeography.

As the number and scale of comparative phylogeographic studies increased, the complexity and heterogeneity of species histories across individual barriers became increasingly evident. Indeed, most studies of multiple taxa across individual barriers find heterogeneity in the timing of vicariant splits and population dynamics across the barrier (for a few of many examples, see Schneider et al. 1998; Barber and Klicka 2010; Naka and Brumfield 2018; Thom et al. 2020; Provost et al. 2021). Another facet of comparative phylogeography that has expanded in recent years is the ubiquity of dispersal as a mechanism for obscuring common vicariant patterns. Indeed, dispersal and population expansion from Pleistocene refugia have been common characteristics across many multispecies communities investigated, particularly in temperate regions and the marine realm (Hewitt 1996, 1999; Bernatchez and Wilson 1998; Floeter et al. 2007; Burney and Brumfield 2009; Lohman et al. 2011; Bagley and Johnson 2014; Smith et al. 2014). The extent to which lineages depart from a strict vicariant patterns is in many cases linked to morphological and life history traits related to dispersal and vagility (Paz et al. 2015; Papadopoulou and Knowles 2016; Puntz et al. 2017; Schiebelhut and Dawson 2018; Sullivan et al. 2019; Ribeiro et al. 2020). Along with historical niche modeling, the integration of trait-based ecology into comparative phylogeography has had a profound influence on the trajectory of the field.

**Coalescent Theory and the Death of Classical Phylogeography**

Coalescent theory, focusing on statistical properties of gene trees and introduced to a wide audience by the mid-1980s, was an important addition to the phylogeographers’ toolkit. Strikingly for empiricists, an authoritative history of the origin of coalescent theory does not mention Avise, Allan Wilson, or any of the empiricists working in phylogeography during the transition to coalescent theory, which was driven by more theoretical considerations (Kingman 2000). Theoreticians quickly adopting coalescent theory, such as Tajima (1983), provided a much more direct link with empirical data sets, and the production and visualization of gene trees, particularly large ones of the human and other iconic species (Cann et al. 1987), was an important driver of coalescent theory’s early development. The early connection between phylogeography and coalescent theory—for example, to estimate population size trajectories (Rienzo and Wilson 1991; Slatkin and Hudson 1991) or population migration rates (Slatkin and Maddison 1989)—was an exciting time, even for single-locus mtDNA or cpDNA studies, which generated new insights, many of which have proved remarkably resilient. The development of spatially realistic coalescent theory remains a vibrant area of population genetics today (Bradburd and Ralph 2019).

As coalescent theory developed to address the burgeoning multicloud data from next-generation sequencing approaches, it is reasonable to ask whether comparative phylogeography—in the sense of interpreting gene trees in space—has had its day. As in early empirical surveys, the first applications of coalescent theory in phylogeography viewed...
gene trees as fixed entities, requiring researchers to estimate them first and infer parameters second (Slatkin and Maddison 1989). However, a key transition point in the field was the shift away from “gene tree” thinking and toward population and lineage thinking: the realization that gene trees themselves are nuisance parameters from the perspective of estimating population genetic parameters (Edwards and Beerli 2000; Hey and Machado 2003; Rosenberg and Nordborg 2002), just as they are considered now for estimation of phylogenetic histories of species (Edwards 2009). New software for simulating coalescent histories within diverging lineages helped reinforce this view among empiricists (Beerli and Felsenstein 1999; Excoffier et al. 2000). Around the time of Hickerson et al.’s review, disagreements arose as to how phylogeography should incorporate the variation and stochasticity found in gene trees as the number of markers expanded. Lacey Knowles coined the term “statistical phylogeography” to acknowledge the inherent noisiness of gene trees and to encourage the use of summary statistics and ensemble analyses going forward (Knowles 2009). Others, like Alan Templeton, had promoted methods, such as nested clad analysis (NCA), that focused on more literal interpretation of branching patterns in gene trees as signals of population inferences (Templeton 1998). Coalescent simulations suggested that NCA overinterpreted signals in gene trees, resulting in many false positives (Panchal and Beaumont 2007), and arguments ensued as to whether the signals utilized by NCA were valuable, or whether likelihood or approximate Bayesian methods that factored in uncertainty in the gene trees were the way forward (Beaumont et al. 2010; Templeton 2010). Likelihood and simulation methods appear to have won that argument (Petit 2008)—while at the same time ushering in a suite of new analytical approaches that moved the field away from the gene trees that marked its founding.

The solution arising over the last two decades is to estimate population divergence histories from extensive multilocus data and coalescent models, often using simulations and ABC methods to compare different hypotheses (Knowles 2009), themselves generated from spatial modeling of species or habitats through past climates (Hugall et al. 2002; Carnaval et al. 2009). Many such methods avoid gene tree estimation entirely, for example, by using SNP data and the multidimensional site frequency spectrum (mdSFS) directly (Xue and Hickerson 2015). In turn, hierarchical methods using the mdSFS across species were developed to test for common histories of vicariance or population expansion without interrogating gene trees directly (Overcast et al. 2017; Xue and Hickerson 2017, 2020). Indeed, the software msBayes and its updates and relatives (Oaks 2014, 2019; Robinson et al. 2014; Bunnefeld et al. 2018) are arguably the only software focused specifically on comparative phylogeography, as opposed to single-species phylogeography.

Against this backdrop of innovation, however, comparative phylogeography as a field seems no longer to be growing, as judged by numbers of publications (fig. 1). The number of publications related to comparative phylogeography seems to have plateaued since about 2010, no longer growing linearly or even exponentially as many new fields do (supplementary tables S1 and S2, Supplementary Material online). The transformation of the tools of comparative phylogeography—from the data-analysis perspective of empiricists—may help explain these trends. For example, some might argue that the eclipse of gene trees from phylogeography, as well as the leveling-off of publications per year (fig. 1), is sufficient grounds to proclaim the field dead, or at least superseded by modern, gene-tree-free approaches. So, is comparative phylogeography dead, or transformed? We assert the latter. Although the focus on gene trees in space, across multiple codistributed taxa will likely be replaced by SNP- or haplotype-based approaches, the core goals of comparative phylogeography remain in the population genomics era. The major change in perspective, we suggest, is the increased focus on the expectations of coalescence histories across space and taxa, rather than the estimated gene trees themselves. Empirical estimation of single-locus gene trees may now more profitably serve the valuable role of generating hypotheses and sampling designs for more intensive, multilocus, genome-wide SNP-based approaches. Although there are now many examples of discordance between organelle and nuclear gene histories, often due to hybridization, selection, or nonrandom lineage extinction, rapid phylogeographic surveys of a region via estimated mitochondrial or chloroplast gene trees still prove very useful and have particular value in relatively underexplored regions, including much of the biodiverse tropics.

At the same time, it is unclear whether comparative phylogeography will, or needs to, embrace WGS as a methodological standard. Even with dropping sequencing costs, WGS is unlikely to be adopted widely by diverse researchers around the globe. Moreover, WGS is likely overkill and is not required to answer the core questions of the field (table 1); genome subsampling approaches such as ddRadseq, or capture methods, will likely carry the field forward for the foreseeable future (Oswald et al. 2017; Alter et al. 2017; Thom et al. 2020; DiBattista et al. 2020; de Medeiros and Farrell 2020). Conceivably, the greater access to WGS methods will cause the conceptual boundaries of comparative phylogeography to expand, incorporating more questions traditionally in the domain of population genetics (Edwards et al. 2015). By contrast, landscape genomics may well embrace WGS (table 1), given its focus on adaptation of organisms to environmental gradients and effects of habitat heterogeneity on genetic diversity, as well as connections with life history, dispersal, and population connectivity (Gagnaire 2020). On the other hand, large meta-analyses of genetic diversity across many species across the globe often require restricting attention to commonly employed markers, such as mtDNA ("comparative mitogenomics"), with concomitant uncertainty about what genome-wide processes might look like (Miraldo et al. 2015).
The importance of museum specimens in phylogeographic work in general has been widely discussed, and the value of specimens linked to the genomic resources forming the basis of phylogeographic studies—as means of replication and verification, further research, or as direct sources of historical DNA—is increasingly appreciated (Besnard et al. 2016; Yao et al. 2017; Schmitt et al. 2018; Nakahama 2021). However, the relationship of the three fields outlined in Table 1 to the museum collections community is uneven. To the extent that comparative phylogeography emerged in part from the systems community—and conceptual antecedents like vicariance biogeography suggest that this is at least partly true—the field also emerged from the museum community and its values. But its prime exponent, John Avise, was not a museum scientist and, in truth, archiving of specimens for phylogeographic research is highly patchy and inconsistent. Poor and inconsistent archiving of museum specimens for genomics work is well documented (Buckner et al. 2021), and museum specimens in landscape genomics are unusual, especially given that field’s orientation toward ecology. Thus, whether comparative phylogeography in the future will continue to benefit from the extra value provided by museum specimens is unclear.

Fig. 1.—Number of publications per year with some variant of “comparative phylogeography” in the title, as referenced in the InCite database (accessed March 28, 2021). Dashed red line indicates a line of constant increase in numbers of publications from 1999 to 2008 (slope, increase of 6.42 publications per year). Dashed bar for 2021 indicates that a partial year is represented. Data are from the 1229 articles in supplementary tables S1 and S2, Supplementary Material online.

Comparative Phylogeography: A Global Perspective

Is Comparative Phylogeography Becoming More Global?

Figures 2 and 3 and supplementary table S4, Supplementary Material online, present summaries of some of the major sites of comparative phylogeographic investigation today in the New and Old Worlds, respectively, built on earlier work, identified, or restudied since the first investigations of the codistributed faunas around the Gulf of Mexico and Atlantic Ocean in the mid-1980s (Bermingham and Avise 1986). Building on the overview of phylogeographic breaks presented by Riddle (2016), and focusing primarily on terrestrial vertebrates, we document 93 major sites of divergence across the globe, many in the tropics. It seems likely that the number of geomorphological barriers facilitating divergence in the tropics is still underestimated, and we present only some of the known physical barriers in the tropics, especially around the barrier-rich regions of the northern Andes and Amazon basin in South America (Naka et al. 2012; Cuervo 2013; Naka and Brumfield 2018). Tropical Southeast Asia is likely another region whose barrier numbers have been underestimated, although the full extent of bias against the tropics versus temperate regions has not been studied.

For figures 2 and 3, we have used a recently proposed Earth projection that provides technical and ethical
improvements upon classical projections used in comparative phylogeography. The particular global projection used in figures 2 and 3 was proposed only in February 2021 and represents the best 2D projection yet as measured by six distortion metrics, including isotropy, area, flexion, skewness, distances, and boundary cuts (Gott et al. 2021). It is well known that many projections of the globe induce biases in how researchers and the public perceive the world; this awareness of implicit biases in data visualization is beginning to influence many areas of science, including modern genomics (Narayan et al. 2021). For example, the Mercator projection, which causes the impression that many high-latitude countries are larger in area than those at the equator, is known to cause biases in perception of the global order (Haemer 1949; Castex 1993). Riddle (2016) appears to have used the Gall–Peter projection, first presented in 1855 but promoted widely in the 1970s. Although the Gall–Peter projection captures a more “equal-area” perspective than the Mercator, it still grossly distorts the shapes of landmasses at the poles and the equator (Vujakovic 1989). The Gott et al. (2021) projection is a radical and elegant departure from previous projections and requires two discs, rather than a single flat image, to represent the entire globe. We suggest that the Gott et al. (2021) projection presents the most scientifically and socially equitable context yet proposed for depicting geography; we encourage its more widespread use, in comparative phylogeography and other fields.

In an important review, Beheregaray (2008) pointed out that comparative phylogeography was underrepresented by what today we call the Global South, and encouraged international collaborations as one way of empowering countries in the developing world. We update his investigation in figure 4, comparing geographical trends in publications focusing on comparative phylogeography before and since his review (supplementary table S3, Supplementary Material online). Assigning one paper contribution to each country that is represented in the author line, we find that, although there still appears to be a strong bias toward publications coming from the Global North, since 2008 there has been a noted increase in publications coming from the Global South. Although, on average since 2008, the proportional contribution of the Global South to the literature on comparative phylogeography has increased by 39.1%, the contribution of Global South countries has increased by 97.5% (supplementary fig. S1, Supplementary Material online). The difference in the distribution of contributions by country between the two time periods is significant (two-sided Kolmogorov–Smirnov test, $D = 0.59813$, $P < 2.2 \times 10^{-16}$). By tabulating country-of-origin of authors of papers on comparative phylogeography, our analysis focuses on where the work is getting done, and not directly on the geographic focus of the work. Still, it is likely that most authors of papers in comparative phylogeography are focusing on systems close to home, with the result that more regions of the globe are being studied as the geographical diversity of authorlines expands. For example, since 2008, and even before that date, there has been a steady accumulation of studies in the regions like the Atlantic Forest and Amazon of South America, many of them driven by researchers in Brazil, whose contribution to the literature in comparative phylogeography has grown since 2008 by 140% (Raposo do Amaral et al. 2013; Thom et al. 2020). The contribution of China, considered a member of the Global South, to the comparative phylogeography literature since 2008 has grown by 236%. Although only three African countries (Madagascar, Côte d’Ivoire, and South Africa) contributed to the pre-2008 literature on comparative phylogeography, 21 countries in Africa published at least one paper in comparative phylogeography after 2008, with an average of 4.85 publications per country in the latter period. There has been growth in Africa as a focus of comparative phylogeography in general (Hydeman et al. 2017; Beddek et al. 2018; van Asch et al. 2019; Helmstetter et al. 2020), although the extent to which in-country researchers participate in this increase is variable (Lorenzen et al. 2012; Freilich et al. 2016). Although we record only five comparative phylogeography publications from two countries (Indonesia and Malaysia) in SE Asia before 2008, in the later period ten SE Asian countries (not including Singapore, a Global North country) produced a total of 73 publications, with greater than 7 coming from authors in each of Vietnam, India, Thailand, Cambodia, Philippines, and Indonesia (fig. 4). Very few phylogeographic studies have focused on the rich biodiversity of India (Reddy 2014); the few studies since 2008 have revealed exciting phylogeographic patterns in birds (Robin et al. 2010, 2015), mammals (Karanth et al. 2010), amphibians (Meegaskumbura et al. 2002; Bossuyt et al. 2004; Vijayakumar et al. 2016), and reptiles (Agarwal and Ramakrishnan 2017), with several leading to descriptions of new taxa, particularly in the Western Ghats. We hope these trends continue and that new regulations promoted by the Convention on Biological Diversity, including the Nagoya Protocol, not only empower researchers from the Global South but also encourage international collaboration, data sharing and open access, especially for fields like comparative phylogeography, which are generally not driven with an eye for profit.

One reason for the plateau in publication number observed in figure 4 is that comparative phylogeographic studies, by necessity, are more challenging and more expensive to complete than single-species studies. Garrick et al. (2015) noted that the number of loci in phylogeographic studies increased over time only in single-species studies, which consistently employed larger numbers of loci than comparative phylogeographic studies. The embrace of genome-wide methods in comparative phylogeography will be most pronounced in wealthier countries (Edwards et al. 2016; Afonso Silva et al. 2017; Oswald et al. 2017), even as the Global South increases its participation in the field. It is unclear whether comparative
phylogeography will ever fully embrace WGS (except perhaps in meta-analyses of single-species studies), or whether it requires it to achieve its goals (table 1).

**Comparative Phylogeography: A Bridge between Landscape Genetics and Speciation**

The focus on geography and potential isolating barriers on the landscape puts comparative phylogeography in a privileged position to link the micro- and macroevolutionary ends of the speciation continuum (Harvey et al. 2019) (fig. 5). Speciation research is highly heterogeneous in its focus. At one extreme, there are the many empirical studies and mathematical theories focusing on the genetics of reproductive isolating mechanisms and the role of pre- and postzygotic isolating mechanisms. On the other extreme, there are numerous studies focusing on the geography and demography of the speciation process, with phylogeographic approaches, bottlenecks, and introgression having prominent roles. Moreover, a large body of literature focuses on ecological factors and organismal traits influencing the rate of speciation across higher level clades (Rosenblum et al. 2012). Comparative phylogeography stands to bridge these extremes by grounding empirical studies in the demography of speciation and quantifying the contribution of the landscape and environmental change to the likelihood of speciation (Singhal and Bi 2017; Potter et al. 2018; Edwards et al. 2019; Harvey et al. 2019). Recent genomic studies of speciation have repeatedly shown the prevalence of ongoing gene flow as species diverge, producing in many cases sister species that differ phenotypically but genomically only at a few places.
along the genome. Prominent examples of this outcome are found in the butterfly genus *Heliconius* (Martin et al. 2013) and many bird species (Toews et al. 2016). The prevalence of gene flow during the speciation process has highlighted the important role of incompatibilities and premating isolating mechanisms in keeping gene pools functionally distinct at few or many loci across the genome. Comparative phylogeography can aid such studies by estimating, for example, how long recent species have been in sympatry or what landscape features may have facilitated the achievement of sympatry. Although these new speciation models have made processes like sympatric speciation and ecological speciation due to local adaptation more plausible, they have not seriously undermined the long-appreciated role of geographic isolation in many speciation scenarios. Geographic isolation, in turn, is a major cause of phylogeographic concordance across lineages that is a core focus of comparative phylogeography.

Comparative phylogeographic studies, particularly when conducted with genome-wide methods, can also help determine the prevalence of genome-wide genetic differentiation of interacting and diverging populations, and thereby help us gauge how often speciation is accompanied by limited genomic divergence. Many studies in comparative phylogeography are conducted with a primary focus on demographic process and genetic diversity, rather than the genetics of speciation per se. Nonetheless, as we accumulate studies in comparative phylogeography, we obtain a valuable catalog—unbiased by a focus on unusual and genetically unique speciation scenarios—of the environmental, demographic, and genomic context in which speciation takes place (Moritz et al. 2009). Harvey et al. (2019) suggested that the metapopulation framework, with its emphasis on gene flow, population divergence, and local extinction, will be a useful addition to models and studies aiming to understand the speciation

**FIG. 3.—**Major phylogeographic breaks in the Old World, with projection details as in figure 2. Numbers correspond to listing of breaks in supplementary table S4, Supplementary Material online.
process. Comparative phylogeography, with its clear connections to the metapopulation concept, can help estimate the population sizes, rates of gene flow, and divergence times that accompany suites of speciation events. Such demographic factors have been argued as a key ingredient to ground modern speciation studies in the messy realities of real populations in nature (Harvey et al. 2019). Moreover, by accumulating repeated instances of divergence across specific geographic barriers, comparative phylogeography can help determine whether different lineages have responded to similar geographic contexts and landscape barriers uniformly or variably (Peñalba et al. 2019).

Finally, comparative phylogeography can help gauge the frequency of population extinction and “ephemeral speciation,” which have been suggested to play an important role in unifying the disparate estimates of speciation rates.

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**Fig. 4.**—Global trends in geographic locations of authors of publications in comparative phylogeography in two time periods, 1992–2008 and 2009–present. (A) Proportion of publications with authors from countries listed on the x-axis to total publication output for two time periods. (B) Shifts in number of publications and proportion of publication by authors in countries listed as Global North or Global South (country designations from https://meta.wikimedia.org/wiki/List_of_countries_by_regional_classification). Data on which these figures are based are in supplementary table S3, Supplementary Material online. We recognize that the terms “Global North” and “Global South” may offend some readers and for that we apologize.

**Fig. 5.**—Depiction of how different approaches in comparative genetics scale geographically. In this example, focused on low dispersal vertebrates northern Australia, population to landscape studies might be done in the north Kimberley at scales of 10–200 km, phylogeographic to speciation studies at the scale of 100s—1000 km (Kimberley to Top End), and speciation to macroevolution at continental scale. Images are from Google Earth; see Potter et al. (2018) as an example.
across studies (Rosenblum et al. 2012). Although population extinction and the presence of ghost populations is intrinsically challenging to detect (Slatkin 2005), a variety of phylogeographic tools now permit researchers to postulate and test for the existence of now extinct species and populations and their contribution to extant genetic diversity. Such approaches can leverage direct interrogation of the genetics of extinct populations via ancient DNA or can infer such populations by investigating anomalies in the genetic landscape of extant populations. Recent studies of extinct hominin species, such as Neanderthals, Denisovans, and other groups provide compelling examples of the former approach. Direct interrogation of ancient DNA and integration with extant samples is becoming more prevalent in nonhuman species, particularly in Africa and the Arctic (Shapiro et al. 2004; Soubrier et al. 2016; Palkopoulou et al. 2018; de Manuel et al. 2020). A few studies have leveraged enough samples from multiple ancient codistributed species to warrant status as a comparative phylogeographic study (Lorenzen et al. 2011). Notably, Lorenzen et al. (2011) found that responses of different large mammals to climate change in the Late Quarternary were idiosyncratic, reflecting the frequent finding from studies of extant populations that different species respond to similar barriers in different ways and at different times. Other lines of evidence, such as evidence for deep mtDNA lineages having been overridden by nuclear DNA gene flow from larger, expanding populations, as well as theory (Alcala and Vuilleumier 2014), have been combined with distributional modeling to infer extinction of populations by genetic swamping (Singhal and Moritz 2012).

Comparative Phylogeography as Place-Based Evolutionary Biology

Marginalizing over Geography in Comparative Population Genomics

Given its roots back to the 1980s and 1990s, it is reasonable to suggest that comparative phylogeography predates comparative population genomics, especially when we consider that population genomics as a field did not really emerge until genome-wide analyses even for single species became routine in the late 2000s. One can additionally argue that—perhaps because of its relative novelty—comparative population genomics is more trendy, and certainly more ascendant, than comparative phylogeography. Still, as reviewed earlier, there have been important conceptual and methodological innovations coming from comparative phylogeography that reinforce its relevance. However, comparative phylogeography continues to focus on the core questions of codiversification that have dominated it since its inception. Comparative population genomics, on the other hand, already seems to be asking questions that hitherto had not been asked before by single-species studies, as this special issue of GBE amply shows. Thus, it is helpful to clarify the relationships between comparative phylogeography and comparative population genetics as a means of understanding what each has to offer the other.

A major strand of comparative population genomics seeks to explain the levels of genetic diversity observed across species. We now know that multiple ecological, demographic, and phylogenetic factors can influence levels of genetic diversity, a deceptively simple summary of genetic variation that is influenced most directly by effective population size, local recombination rate, and mutation rate. This simplicity remains, but researchers now seek multiple upstream factors influencing these two primary determinants (Ellegren and Galtier 2016). The frequency with which the equilibrium level of diversity in a species is realized is now thought to be relatively rare, because pervasive linked selection tends to reduce the realized diversity compared with that expected without linked selection, especially in species with large effective population size (Corbett-Dettig et al. 2015). Although the field is far from settled, some of the important suggested factors include life history traits, such as where the species lands on the r to K spectrum, including its mating system and average offspring number (Romiguier et al. 2014). Recent population genetic surveys across multiple species in a clade—studies in butterflies (Mackintosh et al. 2019) and pinnipeds (Peart et al. 2020) are recent examples—while not downplaying the role of geographic context in modulating genetic diversity, tend to marginalize over it, striving instead for a generality that is suprageographic. Geographic and demographic history necessarily plays an important role in modulating genetic diversity; but one of the main goals of comparative population genetics is to develop predictions for genetic diversity that transcend geography.

“Space is the Place” in Comparative Phylogeography

In contrast to the goals for comparative population genetics, for comparative phylogeography, “space is the place” (Battey et al. 2020). In seeking to understand how environmental variation across space and time, for a given region, has shaped the evolution of genetic diversity and opportunity for speciation, comparative phylogeography is place-based evolutionary biology. The emphasis on multiple taxa across a common region contrasts with the comparative population genomics, where often geography is explicitly ignored or factored out as a nuisance parameter, in order to make generalizations that transcend geography (Leroy et al. 2021) (table 1).

Comparative phylogeography shares with comparative population genomics a desire for generality, but that generality often comes in the form of discovering common geographic boundaries influencing the location and timing of lineage splits. A given comparative phylogeographic study is unabashedly regional—it’s focus is on how the specific
Comparative Phylogeography, Place-Based Education, and Mutual Understanding

With its strong emphasis on geography and landscape features, and its focus on specific biomes and regions, comparative phylogeography is appealing as a means to facilitate place-based education. Conducted inclusively, comparative phylogeography can also serve as a forum for mutual understanding between scientists and local communities and indigenous groups around the world. Place-based education is education using examples from student’s local communities and environments and has emerged as an important vehicle for conveying the immediacy and relevance of scientific principles while at the same time inculcating principles of sustainability (Smith 2002, 2007). Although place-based education has usually been envisioned at the level of very local communities—much smaller in scale than the typical comparative phylogeographic study (fig. 5)—the importance of geography to place-based education has also been acknowledged, both as an object of study but also as a means of analysis and connecting different scales of inquiry (Shimeld 2012; Preston 2015). The study of phylogeography is intrinsically place-based and can help convey important scientific principles to students, particularly when paired with tangible experiences like use of museum specimens (Cook et al. 2014). By drawing attention to details of specific regions of the globe, comparative phylogeography can serve as a powerful place-based context in which students can learn geography and the landscape features that have influenced local plants and animals. Another strength of comparative phylogeography, and phylogeography in general, is its integration of organismal biology and geography. This integration and relevance to organisms found in local communities can be a powerful force in engaging students in evolutionary biology.

Comparative phylogeography can also help mediate dialogues with local and indigenous communities in ways that comparative population genomics alone would find challenging. Indigenous knowledge systems are recognized as powerful sources for place-based education and learning communities (Davidson-Hunt and O’Flaherty 2007), and we suggest comparative phylogeography has great potential for dialogue and study co-design with indigenous peoples. With its frequent references to prominent landscape features and pathways for movement of organisms, comparative phylogeography provides compelling examples not only of scientific principles with clear connections to indigenous knowledge but also of the potential for collaboration between the scientific community and indigenous communities (Moritz et al. 2013; Colwell 2016). Indeed, many comparative phylogeographic studies have been conducted in the spirit of collaboration with indigenous peoples and indigenous peoples. Modern genomics and museum science, fields allied with comparative phylogeography, have both had exploitative relationships with indigenous peoples in the past. However, the record of collaboration is improving, and there is an increasing legal and moral expectation of consultative processes and a desire for mutual learning and understanding (Colwell 2016; Card et al. 2021). Recent work, such as the sequencing of the genome of the Tuatara (Sphenodon punctatus) of New Zealand, offers compelling high-profile examples of this inclusive approach to evolutionary biology (Gemmell et al. 2020).

Conclusion

Comparative phylogeography, with roots going back to the 1980s, offers an instructive counterpoint to the relatively new field of comparative population genomics. Although comparative population genomics, in its most extreme forms, eschews the details of geography in an effort to focus on selective forces acting on the genome, comparative phylogeography embraces geography and strives to link landscape history with codiversification of organismal lineages. Although the output of papers in comparative phylogeography has plateaued in recent years, we suggest that comparative phylogeography as a discipline is still vibrant and innovative, having been transformed from its roots in gene tree analysis by fields like coalescent theory and tools like simulation, trait-based hypotheses, and machine learning. The geographic breadth of comparative phylogeographic studies has increased in the last decade, with a greater contribution to the literature by countries in the Global South than in the previous two decades. Comparative phylogeography offers a compelling bridge between landscape genomics at the microgeographic level and the speciation process. Finally, with its emphasis on geography and landscape features, comparative phylogeography—more so, we suggest,
than comparative population genomics—offers great opportunities for promoting place-based education and engaging local and indigenous communities. We hope this review will encourage continued dialogue and cross-fertilization between comparative population genomics and comparative phylogeography, because both fields stand to be enriched by the other.

**Supplementary Material**

Supplementary data are available at Genome Biology and Evolution online.

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