Linking evolutionary and ecological theory illuminates non-stationary biodiversity

A. J. Rominger^{1, 2}, I. Overcast³, H. Krehenwinkel¹, R. G. Gillespie¹, J. Harte^{1, 4}, and M. J. Hickerson³

¹Department of Environmental Science, Policy and Management, University of California, Berkeley

²Santa Fe Institute ³Biology Department, City College of New York ⁴Energy and Resource Group, University of California, Berkeley

Corresponding author: Rominger, A.J. (ajrominger@gmail.com).

Keywords: Equilibrium dynamics, ecology-evolution synthesis, neutral theory, maximum entropy, next generation sequencing

¹ Abstract

2 stub

3 1 Equilibrium in ecology and evolution

- 4 We propose that combining insights from ecological theory and inference of evolutionary and
- 5 demographic change from genetic data will allow us to understand and predict the conse-
- 6 quences of non-equilibrial processes in governing the current and future states of biodiversity.
- 7 The time is ripe to fully harness the vast amount of genetic and genomic data being gener-
- ated at unprecedented scales [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13] to address fundamental
- 9 questions in ecology and evolution.
- The idea of an ecological and evolutionary equilibrium has pervaded studies of biodiver-10 sity both on geologic and ecological time scales, and from global to local scales [14, 15, 16, 17, 11 18, 19, 20, 21, 22]. Biodiversity theories based on assumptions of equilibrium, both mecha-12 nistic [18, 20, 22] and statistical [see the Glossary; 19, 23, 24] have found success in predicting 13 ahistorical patterns of diversity such as the species abundance distribution [25, 18, 19] and 14 the species area relationship [18, 19]. However, investigation of the underlying dynamics 15 producing these patterns has revealed that the equilibrium assumed by the theories is not 16 realistic [26], and that many processes, equilibrial and otherwise, can generate the same 17 macroscopic, ahistorical predictions [27, 28]. 18
- The consequences of non-equilibrium dynamics for biodiversity, from diversification to macroecology to conservation, are not well understood. The need to understand non-equilibrial biodiversity processes comes at a critical time when anthropogenic pressures are forcing biodiversity systems into states of rapid transition [29]. The extent to which ecosystems are governed by non-equilibrial processes has profound implications for conservation, which are only just beginning to be explored [30]. For example whether conservation should

focus on conventional preservationist paradigms or adaptive management [30]. Whether biodiversity rapidly and consistently tends toward a steady state also determines how species and communities will respond to global environmental change [29].

Tests of equilibrial ecological theory alone will not allow us to identify systems out of 28 equilibrium, nor permit us to pinpoint the mechanistic causes of any observed patterns 29 indicating non-equilibrial processes. The two shortfalls of equilibrial theory are: 1) if the 30 theory fits observed ahistorical patterns but the implicit dynamical assumptions were wrong, 31 we would make the wrong conclusion about the equilibrium of the system; 2) if the equilibrial 32 theories do not fit the data we cannot know why unless we have a perspective on the temporal 33 dynamics underlying the generation of those data. Existing efforts to directly infer the evolutionary and demographic dynamics underlying community assembly in the context of ecological theory testing are limited by a lack of data and analytical framework (see section 4). The advent of next generation sequencing approaches to biodiversity (cite) have lifted the data barrier, but we need a tool set of bioinformatic methods and meaningful predictions grounded in theory to make use of those data; we call for and sketch that tool set here.

$_{ ext{\tiny 40}}$ 2 Ecological theories and equilibrium

The development of the equilibrium theory of island biogeography (ETIB; [31]) ushered in the advent of mechanistically elegant, predictive theories of general patterns in biodiversity. The theory of MacArthur and Wilson also set the precedent of focusing on equilibrial predictions for biodiversity, instead of transient states. From this starting point, three classes of ecological theory have emerged, mechanistically niche-based theory, mechanistically neutral theory, and mechanistically agnostic, statistical mechanical theory. We will focus on neutral and statistical equilibria here. In so doing, we treat niches as in effect being drivers of nonequilibrium: powerful niche dynamics prevent a system from attaining a neutral or statistical equilibrium. Modeling niche dynamics is difficult due to the inherent high dimensionality of the parameter space implied by verbal niche models [e.g., 32], thus showing a lack of neutrality or statistical equilibrium is easier than directly demonstrating niche factors. We further explore the consequences of this approach in section 8.

53 2.1 Mechanistically neutral theory

Neutral theory [18] assumes that one mechanism—demographic drift—drives community assembly. By presuming that populations do not differ in fitness nor in resource use, neutrality avoids the intractability of over parameterization and arrives as an equilibrial prediction when homogeneous stochastic processes of birth, death, speciation and immigration have reached stationarity. Thus neutrality in ecology is analogous to neutral drift in population genetics.

59 2.2 Statistical theory

Rather than assume that any one mechanism, be it niche-based or neutral, dominates the 60 assembly of populations into a community, theories based on statistical mechanics assume 61 that all mechanisms could be valid, but their unique influence has been lost to the enormity of the system and thus outcome of assembly is a community in statistical equilibrium. In one class of such theories, it is assumed that whatever mechanisms are at play, they are only relevant in determining the values of ecological state variables, and then if the system is allowed to come to equilibrium its properties will be predicted by maximizing information entropy relative to the constraints of the state variables. One example is the maximum entropy theory of ecology (METE), one model realization of which assumes that the area (A_0) of an ecosystem, the total number of species (S_0) in some taxonomic group, the total number of individuals in those species (N_0) , and the total metabolic rate of those individuals (E_0) , capture all necessary information to characterize a community because that community 71 has reached a statistical equilibrium in which the imprint of specific mechanistic forces has been lost. While this theory finds widespread success in predicting ahistorical patterns of species abundance, size, and spatial distribution [19, 25, 33, 34] at single snapshots in time, it fails to match observed patterns in disturbed and rapidly evolving communities [35, 19].

$_{76}$ 3 Inferring non-equilibrium dynamics

Unlocking insight into the dynamics underlying community assembly will help us overcome
the limitations of analyzing ahistorical patterns with equilibrial theory. We need explicit
information about the rates of change of populations and species by processes of demographic
fluctuations, immigration, speciation, and extinction.

While the fossil record can elucidate deep time patterns for select, well-fossilized groups [36], and in limited geographic areas and temporal extents yielding good preservation [37], we require an approach that is applicable across taxa, and scales of space and time. Bridging ecological theory with models from phylogenetics has long given us potential general-use tools to gain insight into the dynamics underlying contemporary biodiversity patterns [38, 39, 40], while links from population genetics have been more recently explored [38, 39, 40, 41, 42, 43, 44, 45, 46, 47].

While inference of community dynamics from phylogenetic data has its own challenges, 88 particularly in reliably reconstructing extinction rates [48] and species trees topologies in 89 the face of ancestral admixture [49, 50], its general applicability to all life and the advent of economical methods for producing massive amounts of genetic data, make it a promising 91 approach. Likewise, inference of aggregate population histories underlying regional ecologi-92 cal dynamics using assemblage-level genomic polymorphism data faces statistical challenges, 93 such as non-identifiability of demographic histories [51, 52, 53] and estimating levels of variability across histories [54]. Nevertheless, the promise of unifying processes underlying patterns of species diversities and abundances with distributions of historical population size trajectories, colonisation times, speciation times and regional patterns of genetic connectivity begs investigation. The timing is particularly apt because increasingly reliably and inexpensive methods permit collection of genomic data across a broad range of non-model species.

3.1 Population-level inference

Inference of community dynamics that accounts for non-equilibrium historical complexities 102 needs to expand empirical dimensions beyond species abundances and diversities to include 103 axes of information that are historically dynamic with respect to generative models that 104 link spatial-temporal processes and regional genetics patterns across and between species. 105 Coupling phylogenetic and ecological models with extant taxa have been useful [55, 56, 57], 106 yet these approaches lack a link between community-level processes and within-species vari-107 ation, which could reveal and exploit information about aggregate population histories that 108 underlie non-equilibrium ecological models. Population genetics and phylogeography have 109 always had enormous potential for the inference of community expansion and assembly in 110 the general context of post-LGM warming [58], as well as the community assembly of islands. 111 Yet these studies suffer from the high uncertainty surrounding limited numbers of genetic 112 markers [59], overly generic models [60], ignoring spatial processes [61], and confounding 113 effects of spatial patterns of adaptation [62]. 114

Genomic-scale data, however, is now cost efficient and feasible to collect across a wide 115 swath of non-model species and much progress has been made in multi-species historical 116 demographic models [63]. Although spatial methods are also advancing [64, 65, 66, 67], 117 as are ever more powerful historical inferential approaches via genome-scale data [68], test-118 ing community-scale hypotheses with multi-taxa data would be profoundly improved and 119 enriched if population genetic models were grounded in macroecological and biogeographic 120 theory. Conversely, it has been long recognized that models in community ecology have been 121 overly reliant on species abundance distributions which are by themselves often insufficient 122 for distinguishing competing models of assembly without adding other dimensions of data 123

124 [69, 27].

In contrast to the theoretical developments in ecology that have often focused on ge-125 ographical and environmental processes underlying the equilibrium dynamics of aggregate 126 species distributions and regional patterns of diversity, the field of phylogeography has taken 127 a historically-centered and species-specific approach by using geographically contextualized 128 population genetic data for testing hypotheses regarding the causal relationship among geo-129 graphic phenomena, species distributions, and the mechanisms driving speciation. While the 130 development of ecological theory has generally focuses on predictions of abundances and dis-131 tributions under stationarity, the dynamic nature of landscape and habitat change suggests 132 that ecological theory could be greatly enriched by building a joint modeling framework with 133 population genetic theory that explicitly accounts for historical changes in populations and 134 does not rely on stationarity for generative model predictions. 135

3.2 Coalescent-based inference

One of the fundamental tools allowing for complex historical inference with population genetic data is coalescent theory [70, 71, 72, 73, 74]. Now broadly applied, coalescent theory can generate the statistical properties of any sample of alleles across the genome by modeling gene genealogies backwards in time under virtually any complex demographic history thereby allowing model-based estimation of historical parameters such as historical population size fluctuations, divergence and/or colonization times, and migration rates [75].

Estimating isolation, divergence and/or speciation times has been a particularly important application of population genetic data, and use of coalescent theory is of notable
importance in this endeavor because it statistically captures the stochastic discord between
population divergence times and gene divergence times [76, 77]. However, the isolation of
ancestral lineages into sibling lineages is often only part of a more complex history, as migration and admixture at parts of the genome between diverged populations is a common

feature across the tree of life [78, 49, 53, 79], although the frequency and statistical identifiability of this general observation remains highly contentious [80, 81]. In the context of island biogeography and invasion ecology, coalescent-based estimates of isolation times is of particular importance for understanding the dynamics and timing of island colonization, intra-island speciation, as well as invasion times [82, 83, 84, 85].

The history of population size change is also of fundamental importance for understanding 154 the dynamics of community assembly across a variety of ecological settings, and coalescent 155 theory has likewise become the standard tool for estimating size change histories with pop-156 ulation genetic and phylogeographic data on hand [86, 87]. This application of coalescent 157 modeling has been deployed for large numbers of species from which only small numbers of 158 genetic loci are sampled from populations [88], whereas recent advances allowing genome-159 level data enable far more detailed reconstructions of population history [89, 90] that allow 160 accommodating histories of isolation prior to population size change [68]. However, like 161 any model-based approach, missing assumptions about the complexity of underlying demog-162 raphy can result in biased inference [91], while even using a population history model that 163 matches in reality can not overcome inherent statistical problems in model identifiability [51]. 164 Pivotal to the understanding of demographic and evolutionary histories, coalescent theory 165 has also allowed modeling complex patterns of historical population structure [92, 93], gene 166 flow [94, 95], and even incorporation of extinct "ghost populations" [96, 97] with or without 167 the use of ancient DNA samples [98, 99]. Taking all of these elements of demographic his-168 tory together (i.e. structure, divergence, expansion, size change and migration), researcher, 169 simulation-based coalescent approaches such as approximate Bayesian computation [100, 101] 170 have become of notable importance for making statistical inference under complex histories 171 when solving the likelihood function becomes intractable [102]. 172

As important as it is for the inference of complex demographic history, coalescent theory
has also become an important modeling tool for understanding how natural selection shapes
patterns of genetic polymorphism [103, 104, 105]. Indeed, one of the most commonly used

techniques for detecting positive selection relies on a summary statistic that can be easily simulated under the coalescent given alternative models with neutrality or selection [106]. However, similar to challenges in ecological theory in distinguishing between neutral and non-neutral models of community assembly with species abundance distributions, population 179 genetic models of positive and/or purifying selection also have very similar predictions to 180 those derived from neutral histories of population growth and/or expansion [107, 108, 109, 181 110, 111, as well as other more complex models of selection such as polygenic adaptation 182 and interference selection [112, 113]. Regardless of these difficulties, it is likely that various 183 forms of selection and demographic expansion are not mutually exclusive such that making 184 downstream inferences about overall regional biogeographic histories may be accommodated 185 by using coalescent theory to jointly model selection and neutral demographic history to 186 mitigate biased inference [104, 105, 114, 115]. 187

Ultimately, it is at the community level of inference that coalescent-based population 188 genetic methods could be most useful for investigating ecological models that deviate from stationarity. Indeed, it is the inherent historical approach enabled by coalescent methods that can potentially enrich the ecological theoretical approaches to community assembly and stationarity. 192

189

190

191

In fact, this potential was brought up in the early days of phylogeography [116, 117], as 193 it was well recognized that population genetic data from multiple codistributed taxa could 194 augment investigation of traditionally ecologically-centered questions about the geographic, 195 geological, and/or climatological phenomena that have generated the observed distribution 196 of biodiversity. This proposed "comparative phylogeographic" approach offers the oppor-197 tunity of a natural experiment where focal objects (codistributed taxa), have been inde-198 pendently submitted to the same "natural" evolutionary treatments (geologic and climate 199 change scenarios) [118]. Researchers have generally taken one of two approaches, either by 200 reconstructing taxon-specific histories independently for comparison [119, 120, 121] or us-201 ing hierarchical statistical models that accommodate aggregate genetic datasets for testing 202

alternative historical scenarios and/or hypotheses at the community level [122, 123, 59, 124].

Despite over 30 years of comparative phylogeographic studies, there has been almost 204 a wholesale neglect of the growing body of theory from community ecology that seeks to 205 accommodate the relative importance of deterministic (e.g., niche filtering, competition) 206 and stochastic (i.e., neutral) processes governing the assembly of communities. Conversely, 207 ecological models of community assembly tend to view communities as static pools with an 208 ahistorical focus on equilibrium expectations. Indeed, a fertile cross pollination of these two 200 bodies of theory could yield a joint inferential framework to bridge together ecological neutral 210 theory with coalescent-based comparative population genetic modeling to better generate 211 predictions of temporal changes in regional patterns of both richness and abundance as 212 well as community-level patterns of genetic diversity and divergence. This whole new type 213 of inference could potentially decouple expectations of abundance distributions from time 214 dependencies by parameterizing the population genetic component of demographic histories underlying temporal changes in abundances. 216

NEED TRANSITION

217

Current efforts to integrate evolution into ecological theory

Current efforts to synthesize theoretical perspectives from evolution and ecology have made substantial contributions toward understanding what drives biodiversity patterns. However, a more concerted integration is needed, and indeed was not even feasible until recent and ongoing genetic, bioinformatic and theoretical advances. Approaches to date have been hindered by one or more of several general issues: 1) lack of a solid theoretical foundation, 4) inability to distinguish multiple competing alternative hypotheses, 3) lack of comprehensive genetic data, 4) lack of bioinformatic approaches to resolve species and their abundances.

Here we quickly survey the ways these shortcomings have prevented further advances and then move on to the cutting edge of the field.

Community phylogenetics [38] attempted to understand the roles of competition and environmental filtering on community assembly by assuming key ecologically-relevant traits are conserved along phylogenies; without a solid theory of trait-mediated competition and recruitment, nor a solid theory of trait evolution, this program broke down [125]. Largely lost is the opportunity to use phylogenetic information to understand the historical contingencies at play in community assembly [126, 127], a task which phylogenies might be able to perform, while they are often poor proxies for traits [125].

Joint studies of genetic and species diversity [44, 128, 129, 46] are largely correlative, lacking a strong theoretical core that could be used to make testable predictions. These studies also miss the opportunity to explore more than just diversity metrics, but full models of community assembly, population demography and molecular evolution. These studies are also held back by limited access to genetic data, a hurdle we are actively overcoming (see Boxes 1 and 2).

Phylogeographic studies of past climate change have provided insights of how specific groups have responded in non-equilibrium ways to perturbations [118, 119, 121, 124], but such studies cannot make inference about entire community-level processes, nor were they designed to tie into ecological theory and have not been leveraged by theoreticians to gain more realistic insights on demographic and geographic change.

Ecological neutral theory applied to microbial communities [130] has demonstrated that
the same ecological processes that operate at macro-scales may also scale down to communities of microbes. However, such studies have not made use of the immense phylogenetic
and functional genomic resources available for microbes. Nor has the problem of inferring
abundance from metagenomic and metabarcoding data been fully resolved (see Box 2).

4.1 Emerging approaches

Several approaches have been taken that better ground synthesis of ecological and evolu-253 tionary dynamics in rigorous theory. Because the NTB is implicitly an evo-ecological theory 254 [18, 131], despite typically being treated as ahistorical, it is natural to include evolutionary 255 information into inference about the theory's parameters. Etienne cast the solution of the 256 NTB's species abundance distribution as a coalescent problem [132] while Jabot and Chave 257 [133] used approximate Bayesian computation to improve estimates of the NTB's funda-258 mental biodiversity number using phylogenetic information. Efforts have also been made 259 to validate the underlying assumption of ecological equivalence, a key assumption of the 260 NTB, from a phylogenetic perspective [134]. While these efforts improved inference of the 261 parameters involved in making ahistorical predictions of species abundance, they did not 262 aim to improve the underlying realism of the evolutionary dynamic presumed by the NTB. 263 For example, while the NTB accurately predicts phylogenetic tree shape (sensu [133]) it 264 does not accurately reflect tree tempo [135]. The time to equilibration in the NTB is also 265 unrealistically long [26]. While protracted speciation has been proposed to correct some of 266 these tempo problems in the NTB [136], it remains to be tested, by a framework such as the 267 one we propose, whether these theoretical advances can accurately predict joint patterns of 268 population genetics, phylogenies, and communities. 269

Another approach has tested the ahistorical predictions of equilibrial ecological theory 270 through evolutionary snapshots of community assembly and change. Several applications 271 of the NTB in the fossil record have been used to show changes over geologic time in com-272 munity assembly mechanisms [137, 138]. In a similar theme, Rominger et al. [35] used the 273 geologic chronosequence of the Hawaiian Islands in combination with METE to investigate 274 how evolutionary changes in community assembly drove non-equilibrial patterns in networks 275 of plants and herbivorous insects. While Rominger et al. used genetic information to un-276 derstand how evolutionary rates vary between different arthropod clades in response to the 277

geologic chronosequence, these evolutionary snapshot studies lack a quantitative reconciliation of mechanisms inferred by analyses of ahistorical theory with independently inferred dynamics, either from genetic data or stratigraphic time series.

²⁸¹ 5 What is needed now

298

299

300

301

A key limitation of using ahistorical theory to infer dynamic mechanisms is that multiple 282 mechanisms, from simple and equilibrial to complex, can map onto the same ahistorical 283 pattern, such as the species abundance distribution [139, 140, 141, 142, 143]. This means 284 that even when a theory describes the data well, we do not really know the dynamics that led 285 to that good fit—an interpretational pitfall common in many studies that claim mechanistic insight even in novel evolutionary study systems [18, 137, 138]. Studies that do not have 287 a strong theoretical foundation, and instead rely on qualitative predictions such as higher 288 or lower phylogenetic dispersion [38], further exacerbate the problem of many mechanisms 289 mapping onto single phenomenological predictions. 290

Quantitative theoretical foundations and direct information about dynamics can break
this many-to-one mapping of mechanism onto theoretical prediction. This nicely parallels calls to incorporate additional information into community ecology and macroecological
studies [144]. We propose here a needed framework for integrating the dynamics inferred
from population and phylogenetic approaches with with ahistorical, equilibrial ecological theory. There are two complementary options for incorporating the insights of both ahistorical
ecological theory and genetic inference methods:

• Option 1: using dynamics from genetic inference to predict and understand deviations from ahistorical theories. This amounts to separately fitting ahistorical theory to typical macroecological data (often species abundance, but potentially including body size and trophic network links) while also fitting population genetic and/or phylogenetic

models to genetic data simultaneously captured for the entire community. Doing so requires serious bioinformatic advances that would allow the joint capture of genetic or genomic data from entire community samples, while also estimating accurate abundances for each species in those same samples. This is discussed in Box 2.

• Option 2: building off existing theories, develop new joint models that simultaneously predict macroecological and population genetic patterns. This amounts to building hierarchical models that take genetic data as input and integrate over all possible community states given explicit models of community assembly and population coalescence. Such a model approach also represents a major bioinformatic challenge, which is also discussed in Box 2.

5.1 What we could gain from this framework

302

303

304

305

306

307

308

309

310

311

Given the insights that could be gained from either option 1 or 2 above, we could finally 313 understand why ahistorical theories fail when they do—is it because of rapid population 314 change, or evolution/long-distance dispersal of novel ecological strategies? We could predict 315 whether a system that obeys the ahistorical predictions of equilibrial ecological theory is in 316 fact undergoing major non-equilibrial evolution. We could better understand and forecast 317 how/if systems out of equilibrium are likely to relax back to equilibrial patterns. With such 318 a framework we could even flip the direction of causal inference and understand ecological 319 drivers of diversification dynamics. This last point bears directly on long-standing and open 320 debates about the importance of competitive limits on diversification. Competition and lim-321 iting similarity have a long history of study as drivers of diversification. This has culminated 322 in ideas of diversity-dependent diversification [145, 146, 147]. What has not been done is link 323 this back to ecological assembly mechanisms, but the opportunity seems ripe considering 324 the abundance of work on niche differences and fitness differences [20, 148, 149, 150]. There has even been work on this from a phylogenetic viewpoint [151, 152]. Conclusions about phylogenetic patterns (e.g. diversification slowdowns) would be more believable and robust if combined with population genetic inference (e.g. declining populations) and community patterns (e.g. deviation from equilibrium).

NEED TRANSITION

330

Evo-ecological predictions for systems out of equilibrium

The data needed to fully test a non-equilibrial theory of ecology and evolution, synthesizing 333 historical and contemporary biodiversity patterns, are unprecedented in scale and depth. 334 Put simply, we require knowing the species identities of each individual in a sample as well 335 as information on some portion of their genomes such that we can estimate historical demography and diversification. In Box 2 we discuss progress toward generating such data. 337 We highlight two promising routes: 1) estimating abundance from targeted capture high 338 throughput sequencing data (i.e. metabarcoding) to be used in ahistorical ecological the-339 ory testing, and then separately fitting models of demography and diversification; and 2) 340 jointly estimating the parameters of coupled models of community assembly and popula-341 tion demographics. Assuming these two approaches are within reach (as we demonstrate in 342 Bioinformatic advances), we now discuss hypotheses to be tested in our non-equilibrium 343 framework.

6.1 Cycles of non-equilibrium

Ecosystems are likely to experience consistence disturbances (cite) which can occur on ecological time-scales, such as primary success (cite), or evolutionary time scales, such as evolution of novel innovations that lead to new ecosystem processes (cite). We hypothesize that

- these consistence disturbances will lead to cycles of non-equilibrium in observed biodiversity patterns.
- Figure 1 shows how we hypothesize biodiversity to transition between different phases of equilibrium and non-equilibrium. The figure presents deviation between data and ahistorical ecological theory on the y-axis, versus deviations from demographic or evolutionary equilibrium on the x-axis. A clockwise cycle through this space would indicate:
- Panel I \rightarrow Panel II: following rapid ecological disturbance, ecological metrics diverge from equilibrium
- Panel II \rightarrow III: ecological non-equilibrium spurs evolutionary non-equilibrium leading to both ecological and evolutionary metrics diverging from equilibrium values
- Panel III \rightarrow IV: ecological relaxation to equilibrium after evolutionary innovations provide the means for populations to re-equilibrate to their environments
- Panel IV \rightarrow I: finally a potential return to equilibrium of both ecological and evolutionary metrics once evolutionary processes have also relaxed to their equilibrium
- Cycles could also be much shorter, with a system only transitioning back and forth
 between Panel I and Panel II. This scenario corresponds to the system being driven only
 by rapid ecological disturbance, and this disturbance itself following a stationary dynamic
 leading to no net evolutionary response.
- Cycles through this space could also occur in a counterclockwise direction, being initiated by an evolutionary innovation. Under such a scenario we hypothesize the cycle to proceed:
- Panel I \rightarrow IV: non-equilibrium evolution (including sweepstakes dispersal) leading to departure from evolutionary equilibrium before departure from ecological equilibrium
- Panel IV \rightarrow III: non-equilibrial ecological response to non-equilibrium evolutionary innovation

• Panel III \rightarrow I: ecological and evolutionary relaxation

373

We hypothesize that the final transition will be directly to a joint equilibrium in ecological and evolutionary metrics (Panel I) because a transition from panel III to panel II is unlikely, given that ecological rate are faster than evolutionary rates.

A complete cycle cannot be observed without a time machine, but by combining ahistor-377 ical ecological theory and population/phylogenetic inference methods with community-level 378 genetic data we can identify where on the cycle our focal systems are located. Such an 379 approach assumes that abundance data have been estimated from sequence data, ahistorical 380 ecological theories have been fit to those abundance data, and models of population demog-381 raphy and/or diversification have been separately fit to the underlying sequence data. To 382 better under how our focal systems have transitioned between different equilibrium and non-383 equilibrium phases, we must more deeply explore the joint inference of community assembly 384 and evolutionary processes. In the following sections we do just that for each transition 385 shown in Figure 1. We bring to bear other aspects of joint eco-evolutionary inference, in 386 particular the 1) relationship between lineage age (inferred from molecular data) and lineage 387 abundance; 2) the nature of deviation from ecological metrics, specifically the shape of the 388 species abundance distribution; and 3) the nature of deviation from evolutionary metrics, specifically inference of past population change and selection. 390

³⁹¹ 6.2 Systems undergoing rapid ecological change

For systems whose metrics conform to demographic predictions of equilibrium, but deviate from equilibrial ecological theory, we predict that rapid ecological change underlies their dynamics. However, more information is needed to confirm that the system is being driven primarily by rapid ecological change. The first line of evidence could come from a lack of correlation between lineage age and lineage abundance—this would indicate that slow eco-evolutionary drift is interrupted by frequent perturbations to populations, making their size

independent of age (Fig. 2). Actual abundance should similarly be uncorrelated with inference of effective population size from genetic data. Further support for the ecology-only hypothesis could come from a lack of directional selection detected in community-wide surveys of large genomic regions (see Boxes 1 and 2). Taken as a whole, systems in which ecological metrics deviate from equilibrial theory while demographic and macroevolutionary metrics conform to equilibrial theory presents an opportunity to understand and test hypotheses relating to disturbance, assembly, and the shape of the species abundance distribution [e.g.;

406 6.3 Non-equilibrium ecological communities fostering non407 equilibrium evolution

A lack of equilibrium in an ecological assemblage means that the system will experience 408 change on its trajectory toward a future possibility of equilibrium. If ecological relaxation 409 does not occur—by chance, or because no population present is equipped with the adap-410 tations to accommodate the new environmental that drove the system from equilibrium to begin with—then the system is open to evolutionary innovation. Such innovation could take the form of elevated speciation or long-distance immigration of populations with different ecological strategies than those present, relating to the idea that community assembly is a 414 race between processes with potentially different, but stochastic rates [44], thus any process 415 might by chance produce the observed outcome. In this way, speciation and sweepstakes 416 immigration (and its human-added counterpart, invasion) represent similar processes. Spe-417 ciation and sweepstakes/invasion will yield very different phylogenetic signals, however their 418 population genetic signals in a non-equilibrium community may be very similar (e.g. rapid 419 population expansion). Thus where non-equilibrium communities foster non-equilibrium di-420 versification (either through speciation or invasion) we expect to see a negative relationship 421 between lineage age and abundance (Fig. 2) and signs of rapid population expansion in the 422

newest arrivals to the system.

Local extinctions results from non-equilibrium community dynamics are unlikely to scale
up to non-equilibrial macroevolution, so long as the extinctions are not global. For such
ecosystem collapse to occur, continental ecosystems would have to be self-organized to the
point of criticality, a hypothesis which has largely been refuted [153].

428 6.4 Non-equilibrium evolution fostering non-equilibrium ecological 429 dynamics

If evolutionary processes, or their counterpart in the form of sweepstakes immigration/invasion, generate new ecological strategies in a community, this itself constitutes a form of disturbance pushing the system to reorganize, thus our predicted cycle in Figure 1 from Panel I to IV to III. Evolutionary change would have to be extremely rapid to force ecological metrics out of equilibrium, because evolutionary change could be accommodated by ecological communities if it occurs slowly enough, thus we would expect to see phylogenetic signals of adaptive radiation, and corresponding signals of strong selection in genomic-scale sequence data.

438 6.5 Ecological relaxation

Ecological metrics can return to equilibrium either by ecological means (local immigration from the source pool, local extinction, rapid population changes) or by evolutionary means (novel innovations arising through selection and adaptation or sweepstakes immigration).

In either case, communities are likely to return to equilibrium given enough time without disturbance. Because ecological rates are typically faster than evolutionary rates, this ecological relaxation is likely to happen more quickly than evolutionary relaxation, and thus genetic inference may reveal a time-averaged demographic signature of non-equilibrium.

446 6.6 Evolutionary relaxation

Given sufficient time in ecological equilibrium, the time averaged demographic record revealed by genetic inference will likely also re-equilibrate.

Harnessing evo-ecological measures of non-equilibrium for a changing world

451 Conclude on why this all matters.

453

⁴⁵² 8 More future directions

- 1. Integration with fossils beyond dating phylogenies
- 2. Functional genomics, already underway for microbes, but soon to be realistic for macroorganism communities

456 References

- [1] Yu, D.W. et al. (2012) Biodiversity soup: metabarcoding of arthropods for rapid biodiversity assessment and biomonitoring. Methods in Ecology and Evolution 3, 613–623
- [2] Pompanon, F. et al. (2012) Who is eating what: diet assessment using next generation sequencing. Molecular ecology 21, 1931–1950
- [3] Taberlet, P. et al. (2012) Towards next-generation biodiversity assessment using dna metabarcoding. Molecular Ecology 21, 2045–2050
- [4] Ji, Y. et al. (2013) Reliable, verifiable and efficient monitoring of biodiversity via metabarcoding. Ecology letters 16, 1245–1257
- ⁴⁶⁵ [5] Zhou, X. et al. (2013) Ultra-deep sequencing enables high-fidelity recovery of biodiver-⁴⁶⁶ sity for bulk arthropod samples without per amplification. Gigascience 2, 4
- [6] Tang, M. et al. (2014) Multiplex sequencing of pooled mitochondrial genomes—a crucial step toward biodiversity analysis using mito-metagenomics. Nucleic acids research
 469 42, e166–e166
- [7] Bohmann, K. et al. (2014) Environmental dna for wildlife biology and biodiversity monitoring. Trends in Ecology & Evolution 29, 358–367
- [8] Gibson, J. et al. (2014) Simultaneous assessment of the macrobiome and microbiome in a bulk sample of tropical arthropods through dna metasystematics. Proceedings of the National Academy of Sciences 111, 8007–8012
- [9] Shokralla, S. et al. (2015) Massively parallel multiplex dna sequencing for specimen identification using an illumina miseq platform. Scientific reports 5, 9687
- [10] Linard, B. et al. (2015) Metagenome skimming of insect specimen pools: potential for comparative genomics. Genome biology and evolution 7, 1474–1489

- [11] Leray, M. and Knowlton, N. (2015) Dna barcoding and metabarcoding of standardized samples reveal patterns of marine benthic diversity. *Proceedings of the National*Academy of Sciences 112, 2076–2081
- [12] Dodsworth, S. (2015) Genome skimming for next-generation biodiversity analysis.

 Trends in plant science 20, 525–527
- Liu, S. et al. (2016) Mitochondrial capture enriches mito-dna 100 fold, enabling pcr-free mitogenomics biodiversity analysis. Molecular ecology resources 16, 470–479
- ⁴⁸⁶ [14] Sepkoski, J.J. (1984) A kinetic model of phanerozoic taxonomic diversity. III. Post-⁴⁸⁷ Paleozoic families and mass extinctions. *Paleobiology* 10, 246–267
- 488 [15] Alroy, J. (2010) The shifting balance of diversity among major marine animal groups.

 489 Science 329, 1191–1194
- [16] Rabosky, D.L. and Lovette, I.J. (2008) Density-dependent diversification in north american wood warblers. *Proceedings of the Royal Society B: Biological Sciences* 275, 2363–2371
- [17] Rabosky, D.L. (2009) Ecological limits and diversification rate: alternative paradigms to explain the variation in species richness among clades and regions. *Ecol. Lett.* 12, 735–743
- [18] Hubbell, S.P. (2001) The unified neutral theory of biodiversity and biogeography, vol. 32.
 Princeton University Press
- ⁴⁹⁸ [19] Harte, J. (2011) The Maximum Entropy Theory of Ecology. Oxford University Press
- [20] Chesson, P. (2000) Mechanisms of maintenance of species diversity. *Annu. Rev. Ecol.*Syst. 31, 343–366
- [21] Adler, P.B. *et al.* (2010) Coexistence of perennial plants: an embarrassment of niches. *Ecol. Lett.*, no–no

- ⁵⁰³ [22] Tilman, D. (2004) Niche tradeoffs, neutrality, and community structure: a stochastic theory of resource competition, invasion, and community assembly. *Proc. Natl. Acad.* ⁵⁰⁴ *Sci. U. S. A.* 101, 10854–10861
- ⁵⁰⁶ [23] Pueyo, S. *et al.* (2007) The maximum entropy formalism and the idiosyncratic theory of biodiversity. *Ecol. Lett.* 10, 1017–1028
- 508 [24] Shipley, B. et al. (2006) From plant traits to plant communities: A statistical mechanistic approach to biodiversity. Science
- 510 [25] White, E.P. et al. (2012) Characterizing species abundance distributions across taxa 511 and ecosystems using a simple maximum entropy model. Ecology 93, 1772–1778
- [26] Ricklefs, R.E. (2006) The unified neutral theory of biodiversity: do the numbers add up? *Ecology* 87, 1424–1431
- [27] McGill, B.J. et al. (2007) Species abundance distributions: moving beyond single prediction theories to integration within an ecological framework. Ecol. Lett. 10, 995–1015
- [28] McGill, B.J. (2010) Towards a unification of unified theories of biodiversity. *Ecology*Letters 13, 627–642
- [29] Barnosky, A.D. et al. (2012) Approaching a state shift in earth/'s biosphere. Nature 486, 52–58
- [30] Wallington, T.J. et al. (2005) Implications of current ecological thinking for biodiversity conservation: A review of the salient issues. Ecol. Soc. 10
- [31] MacArthur, R.H. and Wilson, E.O. (1967) The theory of island biogeography. Princeton
 University Press
- [32] Hutchinson, G.E. (1959) Homage to santa rosalia or why are there so many kinds of animals? The American Naturalist 93, 145–159

- [33] Xiao, X. et al. (2015) A strong test of the maximum entropy theory of ecology. Am.

 Nat. 185, E70–E80
- [34] Harte, J. et al. (2009) Biodiversity scales from plots to biomes with a universal speciesarea curve. Ecol. Lett. 12, 789–797
- [35] Rominger, A.J. et al. (2015) Community assembly on isolated islands: macroecology meets evolution. Glob. Ecol. Biogeogr.
- [36] Alroy, J. et al. (2008) Phanerozoic trends in the global diversity of marine invertebrates.

 Science 321, 97–100
- [37] Harnik, P.G. (2011) Direct and indirect effects of biological factors on extinction risk
 in fossil bivalves. Proceedings of the National Academy of Sciences 108, 13594–13599
- [38] Webb, C.O. et al. (2002) Phylogenies and community ecology. Annu. Rev. Ecol. Syst.
 33, 475–505
- [39] Emerson, B.C. (2002) Evolution on oceanic islands: molecular phylogenetic approaches to understanding pattern and process. *Mol. Ecol.* 11, 951–966
- [40] Lavergne, S. et al. (2010) Biodiversity and climate change: Integrating evolutionary and ecological responses of species and communities. Annu. Rev. Ecol. Evol. Syst. 41, 321–350
- [41] Li, L. and Ma, Z.S. (2016) Testing the neutral theory of biodiversity with human microbiome datasets. *Sci. Rep.* 6, 31448
- [42] McGaughran, A. (2015) Integrating a population genomics focus into biogeographic
 and macroecological research. Front. Ecol. Evol. 3
- [43] Laroche, F. et al. (2015) A neutral theory for interpreting correlations between species and genetic diversity in communities. Am. Nat. 185, 59–69

- ⁵⁴⁹ [44] Vanoverbeke, J. et al. (2015) Community assembly is a race between immigration and adaptation: eco-evolutionary interactions across spatial scales. Ecography
- ⁵⁵¹ [45] Vellend, M. and Geber, M.A. (2005) Connections between species diversity and genetic diversity. *Ecol. Lett.* 8, 767–781
- ⁵⁵³ [46] Papadopoulou, A. et al. (2011) Testing the Species–Genetic diversity correlation in the aegean archipelago: Toward a Haplotype-Based macroecology? Am. Nat. 178, 241–255
- [47] Dexter, K.G. et al. (2012) Historical effects on beta diversity and community assembly
 in amazonian trees. Proc. Natl. Acad. Sci. U. S. A. 109, 7787–7792
- [48] Quental, T.B. and Marshall, C.R. (2009) EXTINCTION DURING EVOLUTION ARY RADIATIONS: RECONCILING THE FOSSIL RECORD WITH MOLECULAR
 PHYLOGENIES. Evolution 63, 3158–3167
- ⁵⁶⁰ [49] Mallet, J. et al. (2016) How reticulated are species? Bioessays 38, 140–149
- [50] Xu, B. and Yang, Z. (2016) Challenges in species tree estimation under the multispecies coalescent model. *Genetics* 204, 1353–1368
- [51] Terhorst, J. and Song, Y.S. (2015) Fundamental limits on the accuracy of demographic
 inference based on the sample frequency spectrum. Proc. Natl. Acad. Sci. U. S. A.
 112, 7677–7682
- [52] Robinson, J.D. et al. (2014) ABC inference of multi-population divergence with ad mixture from unphased population genomic data. Mol. Ecol. 23, 4458–4471
- 568 [53] Sousa, V. and Hey, J. (2013) Understanding the origin of species with genome-scale data: modelling gene flow. *Nat. Rev. Genet.* 14, 404–414
- 570 [54] Hickerson, M.J. et al. (2014) Recommendations for using msbayes to incorporate un-571 certainty in selecting an abc model prior: a response to oaks et al. *Evolution* 68, 572 284–294

- [55] Emerson, B.C. and Gillespie, R.G. (2008) Phylogenetic analysis of community assembly and structure over space and time. *Trends Ecol. Evol.* 23, 619–630
- ⁵⁷⁵ [56] Rosindell, J. et al. (2015) Unifying ecology and macroevolution with individual-based ⁵⁷⁶ theory. Ecol. Lett. 18, 472–482
- 577 [57] Burbrink, F.T. et al. (2015) Predicting community structure in snakes on eastern
 578 nearctic islands using ecological neutral theory and phylogenetic methods. Proc. Biol.
 579 Sci. 282
- [58] Hewitt, G.M. (2004) Genetic consequences of climatic oscillations in the quaternary.
 Philos. Trans. R. Soc. Lond. B Biol. Sci. 359, 183
- [59] Chan, Y.L. et al. (2014) Detecting concerted demographic response across community
 assemblages using hierarchical approximate bayesian computation. Mol. Biol. Evol.,
 msu187
- [60] Papadopoulou, A. and Knowles, L.L. (2016) Toward a paradigm shift in comparative
 phylogeography driven by trait-based hypotheses. Proc. Natl. Acad. Sci. U. S. A. 113,
 8018–8024
- [61] Meirmans, P.G. (2012) The trouble with isolation by distance. Mol. Ecol. 21, 2839–
 2846
- [62] Hoban, S. et al. (2016) Finding the genomic basis of local adaptation: Pitfalls, practical
 solutions, and future directions. Am. Nat. 188, 379–397
- [63] Xue, A.T. and Hickerson, M.J. (2015) The aggregate site frequency spectrum (aSFS) for comparative population genomic inference. *Mol. Ecol.*
- ⁵⁹⁴ [64] Petkova, D. *et al.* (2016) Visualizing spatial population structure with estimated effective migration surfaces. *Nat. Genet.* 48, 94–100

- ⁵⁹⁶ [65] Joseph, T.A. *et al.* (2016) Demographic inference under a spatially continuous coales-⁵⁹⁷ cent model. *Heredity* 117, 94–99
- [66] Prates, I. et al. (2016) Inferring responses to climate dynamics from historical demography in neotropical forest lizards. Proceedings of the National Academy of Sciences

 113, 7978–7985
- [67] Brown, J.L. *et al.* (2016) Predicting the genetic consequences of future climate change:

 The power of coupling spatial demography, the coalescent, and historical landscape

 changes. *Am. J. Bot.* 103, 153–163
- [68] Terhorst, J. et al. (2016) Robust and scalable inference of population history from hundreds of unphased whole genomes. Nat. Genet.
- [69] McGill, B. (2003) Strong and weak tests of macroecological theory. Oikos 102, 679–685
- [70] Hudson, R.R. (1983) Properties of a neutral model with intragenic recombination.

 Theor. Popul. Biol. 23, 183–201
- [71] Tajima, F. (1983) Evolutionary relationship of DNA sequences in finite populations.

 Genetics 105, 437–460
- [72] Kingman, J.F.C. (1982) The coalescent. Stochastic Process. Appl. 13, 235–248
- [73] Kingman, J.F.C. (1982) On the genealogy of large populations. J. Appl. Probab. 19a, 27–43
- [74] Rosenberg, N.A. and Nordborg, M. (2002) Genealogical trees, coalescent theory and the analysis of genetic polymorphisms. *Nat. Rev. Genet.* 3, 380–390
- [75] Wakeley, J. (2008) Coalescent Theory. Roberts and Company Publishers
- [76] Charlesworth, D. (2010) Don't forget the ancestral polymorphisms. *Heredity* 105, 509–510

- [77] Edwards, S.V. and Beerli, P. (2000) Perspective: gene divergence, population divergence, and the variance in coalescence time in phylogeographic studies. *Evolution* 54, 1839–1854
- [78] Shapiro, B.J. et al. (2016) What is speciation? PLoS Genet. 12, e1005860
- [79] Nosil, P. (2008) Speciation with gene flow could be common. Mol. Ecol. 17, 2103–2106
- [80] Cruickshank, T.E. and Hahn, M.W. (2014) Reanalysis suggests that genomic islands of speciation are due to reduced diversity, not reduced gene flow. *Mol. Ecol.* 23, 3133–3157
- [81] Yang, M. et al. (2017) Can genomic data alone tell us whether speciation happened with gene flow? Mol. Ecol.
- [82] Estoup, A. and Clegg, S.M. (2003) Bayesian inferences on the recent island colonization history by the bird zosterops lateralis lateralis. *Mol. Ecol.* 12, 657–674
- [83] Estoup, A. et al. (2004) Genetic analysis of complex demographic scenarios: spatially expanding populations of the cane toad, bufo marinus. Evolution 58, 2021–2036
- [84] Hickerson, M.J. and Meyer, C. (2008) Testing comparative phylogeographic models of
 marine vicariance and dispersal using a hierarchical bayesian approach. BMC Evol.
 Biol. 8, 322
- [85] Gray, M.M. *et al.* (2014) Demographic history of a recent invasion of house mice on the isolated island of gough. *Mol. Ecol.* 23, 1923–1939
- [86] Kuhner, M.K. et al. (1998) Maximum likelihood estimation of population growth rates based on the coalescent. Genetics 149, 429–434
- [87] Slatkin, M. and Hudson, R.R. (1991) Pairwise comparisons of mitochondrial DNA sequences in stable and exponentially growing populations. *Genetics* 129, 555–562

- [88] Drummond, A.J. et al. (2005) Bayesian coalescent inference of past population dynamics from molecular sequences. Mol. Biol. Evol. 22, 1185–1192
- [89] Schiffels, S. and Durbin, R. (2014) Inferring human population size and separation history from multiple genome sequences. *Nat. Genet.* 46, 919–925
- [90] Boitard, S. et al. (2016) Inferring population size history from large samples of Genome-Wide molecular data - an approximate bayesian computation approach. PLoS Genet. 12, e1005877
- [91] Mazet, O. et al. (2015) On the importance of being structured: instantaneous coalescence rates and human evolution—lessons for ancestral population size inference?

 Heredity 116, 362–371
- [92] Prado-Martinez, J. et al. (2013) Great ape genetic diversity and population history.
 Nature 499, 471–475
- [93] Bahlo, M. and Griffiths, R.C. (2000) Inference from gene trees in a subdivided population. *Theor. Popul. Biol.* 57, 79–95
- [94] Beerli, P. and Felsenstein, J. (2001) Maximum likelihood estimation of a migration matrix and effective population sizes in n subpopulations by using a coalescent approach.
 Proc. Natl. Acad. Sci. U. S. A. 98, 4563–4568
- [95] Hey, J. and Nielsen, R. (2004) Multilocus methods for estimating population sizes, migration rates and divergence time, with applications to the divergence of drosophila pseudoobscura and d. persimilis. *Genetics* 167, 747–760
- [96] Slatkin, M. (2005) Seeing ghosts: the effect of unsampled populations on migration rates estimated for sampled populations. *Mol. Ecol.* 14, 67–73
- [97] Alter, S.E. et al. (2007) DNA evidence for historic population size and past ecosystem impacts of gray whales. Proc. Natl. Acad. Sci. U. S. A. 104, 15162–15167

- [98] Kuhlwilm, M. et al. (2016) Ancient gene flow from early modern humans into eastern neanderthals. Nature 530, 429–433
- [99] Veeramah, K.R. and Hammer, M.F. (2014) The impact of whole-genome sequencing on the reconstruction of human population history. *Nat. Rev. Genet.* 15, 149–162
- [100] Beaumont, M. (2010) Approximate bayesian computation in evolution and ecology.

 Annu. Rev. Ecol. Evol. Syst. 41, 379–406
- [101] Pritchard, J.K. et al. (1999) Population growth of human Y chromosomes: a study of Y chromosome microsatellites. Mol. Biol. Evol. 16, 1791–1798
- [102] Sunnåker, M. et al. (2013) Approximate bayesian computation. PLoS Comput. Biol.
 9, e1002803. doi:10.1371/journal.pcbi.1002803
- [103] Kim, Y. and Stephan, W. (2002) Detecting a local signature of genetic hitchhiking along a recombining chromosome. *Genetics* 160, 765–777
- [104] Kern, A.D. and Schrider, D.R. (2016) Discoal: flexible coalescent simulations with selection. *Bioinformatics* 32, 3839–3841
- [105] Ewing, G.B. and Jensen, J.D. (2016) The consequences of not accounting for background selection in demographic inference. *Mol. Ecol.*
- [106] Tajima, F. (1989) Statistical method for testing the neutral mutation hypothesis by

 DNA polymorphism. Genetics 123, 585–595
- [107] Freedman, A.H. et al. (2016) Demographically-based evaluation of genomic regions under selection in domestic dogs. PLoS Genet. 12, e1005851
- [108] Barton, N.H. (1998) The effect of hitch-hiking on neutral genealogies. Genet. Res. 72, 123–133

- [109] Barton, N.H. (2000) Genetic hitchhiking. *Philos. Trans. R. Soc. Lond. B Biol. Sci.*355, 1553–1562
- [110] Jensen, J.D. *et al.* (2005) Distinguishing between selective sweeps and demography using DNA polymorphism data. *Genetics* 170, 1401–1410
- [111] Schrider, D.R. et al. (2016) Effects of linked selective sweeps on demographic inference and model selection. Genetics 204, 1207–1223
- [112] Stephan, W. (2016) Signatures of positive selection: from selective sweeps at individual loci to subtle allele frequency changes in polygenic adaptation. *Mol. Ecol.* 25, 79–88
- [113] Good, B.H. et al. (2014) Genetic diversity in the interference selection limit. PLoS
 Genet. 10, e1004222
- [114] Phung, T.N. et al. (2016) Determining the effect of natural selection on linked neutral divergence across species. PLoS Genet. 12, e1006199
- [115] Roux, C. et al. (2016) Shedding light on the grey zone of speciation along a continuum of genomic divergence. PLoS Biol. 14, e2000234
- [116] Avise, J.C. et al. (1987) Intraspecific phylogeography: The mitochondrial DNA bridge between population genetics and systematics. Annu. Rev. Ecol. Syst. 18, 489–522
- [117] Avise, J.C. (1998) The history and purview of phylogeography: a personal reflection.
 Mol. Ecol. 7, 371–379
- ⁷⁰⁵ [118] Arbogast, B.S. and Kenagy, G.J. (2001) Comparative phylogeography as an integrative approach to historical biogeography. *J. Biogeogr.* 28, 819–825
- [119] Smith, B.T. et al. (2012) Evaluating the role of contracting and expanding rainforest in initiating cycles of speciation across the isthmus of panama. Proc. Biol. Sci. 279, 3520–3526

- [120] Carstens, B.C. *et al.* (2005) Investigating the evolutionary history of the pacific northwest mesic forest ecosystem: Hypothesis testing within a comparative phylogeographic framework. *Evolution* 59, 1639–1652
- 713 [121] Hickerson, M.J. and Cunningham, C.W. (2005) Contrasting quaternary histories in 714 an ecologically divergent pair of low-dispersing intertidal fish (xiphister) revealed by 715 multi-locus DNA analysis. *Evolution* 59, 344–360
- [122] Hickerson, M.J. et al. (2006) Test for simultaneous divergence using approximate
 bayesian computation. Evolution 60, 2435–2453
- 718 [123] Carstens, B.C. et al. (2016) Community trees: Identifying codiversification in the 719 páramo dipteran community. Evolution 70, 1080–1093
- [124] Satler, J.D. and Carstens, B.C. (2016) Phylogeographic concordance factors quantify phylogeographic congruence among co-distributed species in the sarracenia alata pitcher plant system. *Evolution*
- [125] Losos, J.B. (2008) Phylogenetic niche conservatism, phylogenetic signal and the relationship between phylogenetic relatedness and ecological similarity among species. Ecol. Lett. 11, 995–1007
- [126] Ricklefs, R.E. (2007) History and diversity: Explorations at the intersection of ecology and evolution. Am. Nat. 170, S56–S70
- [127] Emerson, B.C. and Gillespie, R.G. (2008) Phylogenetic analysis of community assembly and structure over space and time. *Trends Ecol. Evol.*
- 730 [128] Vellend, M. (2005) Species diversity and genetic diversity: parallel processes and cor-731 related patterns. Am. Nat. 166, 199–215
- [129] Vellend, M. et al. (2014) Drawing ecological inferences from coincident patterns of
 population- and community-level biodiversity. Mol. Ecol. 23, 2890–2901

- 734 [130] Venkataraman, A. et al. (2015) Application of a neutral community model to assess

 735 structuring of the human lung microbiome. MBio 6
- [131] Hubbell, S.P. (2005) The neutral theory of biodiversity and biogeography and stephen
 jay gould. Paleobiology 31, 122–132
- [132] Etienne, R.S. and Olff, H. (2004) A novel genealogical approach to neutral biodiversity theory. *Ecol. Lett.* 7, 170–175
- [133] Jabot, F. and Chave, J. (2009) Inferring the parameters of the neutral theory of biodiversity using phylogenetic information and implications for tropical forests. *Ecol. Lett.* 12, 239–248
- [134] Burbrink, F.T. et al. (2015) Predicting community structure in snakes on eastern
 nearctic islands using ecological neutral theory and phylogenetic methods. Proceedings
 of the Royal Society B: Biological Sciences 282, 20151700
- [135] Davies, T.J. et al. (2011) NEUTRAL BIODIVERSITY THEORY CAN EXPLAIN
 THE IMBALANCE OF PHYLOGENETIC TREES BUT NOT THE TEMPO OF
 THEIR DIVERSIFICATION. Evolution 65, 1841–1850
- [136] Rosindell, J. et al. (2010) Protracted speciation revitalizes the neutral theory of biodiversity. Ecol. Lett. 13, 716–727
- [137] Olszewski, T.D. and Erwin, D.H. (2004) Dynamic response of permian brachiopod
 communities to long-term environmental change. Nature 428, 738–741
- [138] Wagner, P.J. et al. (2006) Abundance distributions imply elevated complexity of post paleozoic marine ecosystems. Science 314, 1289–1292
- [139] Kendall, D.G. (1948) On the generalized "Birth-and-Death" process. Ann. Math. Stat.
 19, 1–15

- [140] Kendall, D.G. (1948) On some modes of population growth leading to r. a. fisher's
 logarithmic series distribution. Biometrika 35, 6–15
- [141] Engen, S. and Lande, R. (1996) Population dynamic models generating the lognormal
 species abundance distribution. Math. Biosci. 132, 169–183
- [142] Engen, S. and Lande, R. (1996) Population dynamic models generating species abundance distributions of the gamma type. *J. Theor. Biol.* 178, 325–331
- ⁷⁶³ [143] McGill, B. (2003) Strong and weak tests of macroecological theory. Oikos 102, 679–685
- ⁷⁶⁴ [144] McGill, B.J. *et al.* (2007) Species abundance distributions: moving beyond single prediction theories to integration within an ecological framework. *Ecol. Lett.* 10, 995–1015
- ⁷⁶⁶ [145] Etienne, R.S. and Haegeman, B. (2012) A conceptual and statistical framework for adaptive radiations with a key role for diversity dependence. *Am. Nat.* 180, E75–E89
- ⁷⁶⁸ [146] Rabosky, D.L. (2013) Diversity-dependence, ecological speciation, and the role of competition in macroevolution. *Annu. Rev. Ecol. Evol. Syst.* 44
- 770 [147] Rabosky, D.L. and Lovette, I.J. (2008) Explosive evolutionary radiations: decreasing 771 speciation or increasing extinction through time? *Evolution* 62, 1866–1875
- 772 [148] Adler, P.B. et al. (2007) A niche for neutrality. Ecol. Lett. 10, 95–104
- [149] HilleRisLambers, J. et al. (2012) Rethinking community assembly through the lens of coexistence theory. Annual Review of
- ⁷⁷⁵ [150] Levine, J.M. and HilleRisLambers, J. (2009) The importance of niches for the mainte-⁷⁷⁶ nance of species diversity. *Nature* 461, 254–257
- 777 [151] Mayfield, M.M. and Levine, J.M. (2010) Opposing effects of competitive exclusion on 778 the phylogenetic structure of communities. *Ecol. Lett.* 13, 1085–1093

- 779 [152] Godoy, O. *et al.* (2014) Phylogenetic relatedness and the determinants of competitive 780 outcomes. *Ecol. Lett.* 17, 836–844
- [153] Kirchner, J.W. and Weil, A. (1998) No fractals in fossil extinction statistics. Nature
 395, 337–338
- ⁷⁸³ [154] Krehenwinkel, H. *et al.* (2016) A cost-efficient and simple protocol to enrich prey dna from extractions of predatory arthropods for large-scale gut content analysis by illumina sequencing. *Methods in Ecology and Evolution*
- ⁷⁸⁶ [155] Elbrecht, V. and Leese, F. (2015) Can dna-based ecosystem assessments quantify ⁷⁸⁷ species abundance? testing primer bias and biomass—sequence relationships with an ⁷⁸⁸ innovative metabarcoding protocol. *PloS one* 10, e0130324
- ⁷⁸⁹ [156] Coissac, E. *et al.* (2016) From barcodes to genomes: extending the concept of dna ⁷⁹⁰ barcoding. *Molecular ecology*
- [157] Nielsen, H.B. et al. (2014) Identification and assembly of genomes and genetic elements
 in complex metagenomic samples without using reference genomes. Nature biotechnol ogy 32, 822–828
- ⁷⁹⁴ [158] Angly, F.E. *et al.* (2014) Copyrighter: a rapid tool for improving the accuracy of microbial community profiles through lineage-specific gene copy number correction.

 ⁷⁹⁶ *Microbiome* 2, 11
- ⁷⁹⁷ [159] Saitoh, S. *et al.* (2016) A quantitative protocol for dna metabarcoding of springtails

 ⁷⁹⁸ (collembola) 1. *Genome* 59, 705–723
- ⁷⁹⁹ [160] Overcast, I.A. and Hickerson, M.J. (in prep.) Integrating community assembly models and comparative population genetics.
- [161] Kelleher, J. et al. (2016) Efficient coalescent simulation and genealogical analysis for large sample sizes. PLoS Comput Biol 12, e1004842

\mathbf{Boxes}

804 Box 1: Wetlab techniques

Next generation sequencing technology has ushered in a revolution in evolutionary biology and ecology. The large scale recovery from bulk samples (e.g. passive arthropod traps) of species richness, food web structure, cryptic species promise unprecedented new insights into ecosystem function and assembly [154, 9, 8, 3]. Two approaches, differing in cost and effectiveness, have emerged.

Metabarcoding describes the targeted PCR amplification and next generation sequencing 810 of short DNA barcode markers (typically 300-500 bp) from community samples [1, 4]. 811 The resulting amplicon sequences can be clustered into OTUs or grafted onto more well 812 supported phylogenies. Even minute traces of taxa in environmental samples can be detected 813 using metabarcoding [7]. Amplicon sequencing is cheap, requires a small workload and thus 814 allows rapid inventories of species composition and species interactions in whole ecosystems 815 [8, 11, 2]. However, the preferential amplification of some taxa during PCR can lead to 816 highly skewed abundance estimates [1, 155] from metabarcoding libraries. 817

Metagenomic approaches , in contrast, avoid marker specific amplification bias by se-818 quencing libraries constructed either from untreated genomic DNA [12, 10, 6], or after tar-819 geted enrichment of genomic regions [13]. While being more laborious, expensive and com-820 putationally demanding than metabarcoding, metagenomics thus offers improved accuracy 821 in detecting species composition and abundance [5]. Moreover, the assembly of high coverage 822 metagenomic datasets recovers large contiguous sequence stretches, even from rare members 823 in a community, offering high phylogenetic resolution at the whole community level [156]. 824 Due to large genome sizes and high genomic complexity, metazoan metagenomics is currently 825 mostly limited to the assembly of fairly short high copy regions. Particularly mitochondrial 826

and chloroplast genomes as well as nuclear ribosomal clusters are popular targets [12, 156].

In contrast, microbial metagenomic studies now routinely assemble complete genomes and
characterize gene content and metabolic pathways even from complex communities [157].

This allows unprecedented insights into functional genetic process underlying community
assembly and evolutionary change of communities to environmental stress.

Box 2: Bioinformatic advances

While species richness can be routinely identified by sequencing bulk samples using high throughput methods, estimating species abundance remains challenging [155] and severely limits the application of high throughput sequencing methods to many community-level studies. We propose two complementary approaches to estimate species abundance from high throughput data. The first approach estimates abundance free from any models of community assembly, the second jointly estimates the parameters of a specific assembly model of interest along with the parameters of a coalescent-based population genetic model.

Model-free abundance estimation. We propose a pipeline (Fig. I) where raw reads 840 are generated and assembled into a phylogeny using standard approaches, and potentially 841 aided by additionally available sequence data in a super tree or super matrix approach. The 842 numbers of sequences assigned to each terminal tip are then used in a Bayesian hierarchical 843 model which seeks to estimate the true number of organisms representing each terminal tip, 844 accounting for sequencing biases originating from primer affinity and copy number differences 845 between taxa. Information on phylogenetic relatedness can inform modeled correlations in 846 biases between taxa [158, e.g. copy number is known to be phylogenetically conserved 847 at least in microbes. This approach is particularly tailored to metabarcoding data. In a 848 potentially powerful extension, and thanks to the proposed Bayesian framework, information 849 from sequencing experiments that seek to calibrate metabarcoding studies [e.g., 154, 159] can 850 be used to build meaningfully informative priors and improve model accuracy. Through a 851

simulation study (described in the supplement) we show that true underlying abundances
can be accurately estimated (Fig. II).

Joint inference of community assembly and population genetic models. 854 individual-based, forward-time models of community assembly with backwards-time hierar-855 chical multi-taxa coalescent models permits inference about the values of the parameters in 856 both models. This framework is flexible enough to incorporate multiple refugia, coloniza-857 tion routes, ongoing migration and both neutral and deterministic processes of assembly on time scales of hundreds of thousands of years (Fig. III). A forthcoming implementation 859 [gimmeSAD π ; 160] simulates an individual-based forward time community dispersal model 860 [56] linked with the msPrime coalescent simulator [161]. This has been accomplished by 861 rescaling the time dependent local abundance distributions into time dependent effective 862 population size distributions while allowing for heterogeneity in migration and colonization 863 rates. This simulation model can be combined with random forest classifiers and hierarchi-864 cal ABC to enable testing alternative assembly models, including models that have not yet 865 reached their theoretical equilibria. 866

867 Glossary

868 ahistorical

869 coalesent

870 statistical equilibrium

Figures Figures

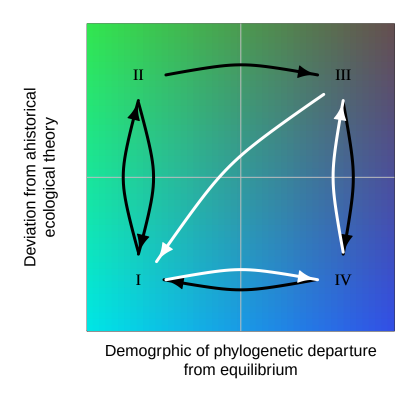


Figure 1: Hypothesized cycles between different states of equilibrium and non-equilibrium in ecological metrics (y-axis) and evolutionary metrics (x-axis). Panels I–IV are discussed in the next. Colors correspond to deviation from ahistorical ecological theory and evolutionary equilibrium. Black cycle corresponds to non-equilibrium initiated by ecological disturbance (with potential to continue to evolutionary non-equilibrium or relaxation to equilibrium). White cycle is initiated by evolutionary innovation.

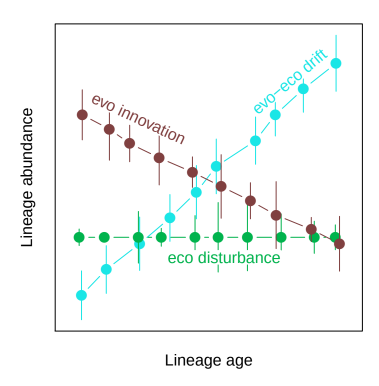


Figure 2: Hypothesized relationships between lineage age and abundance under different evo-ecological scenarios. Colors correspond to panels in Figure 1: teal is evo-ecological equilibrium; green is rapid transition to ecological non-equilibrium following short timescale distrubance; dark brown is non-equilibrium in both ecological and evolutionary metrics.

Box 2 figures

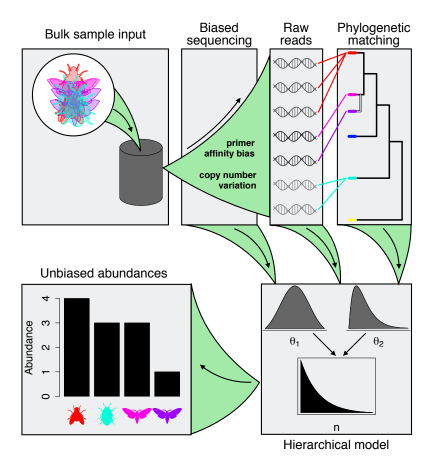


Figure I: Pipeline to estimate true abundances from metabarcoding data. The pipeline follows sequence generation, matching sequences to a phylogeny (generated from the sequences themselves, or better yet from highercoverage data) and finally Bayesian hierarchical modeling leading to abundance estimates.

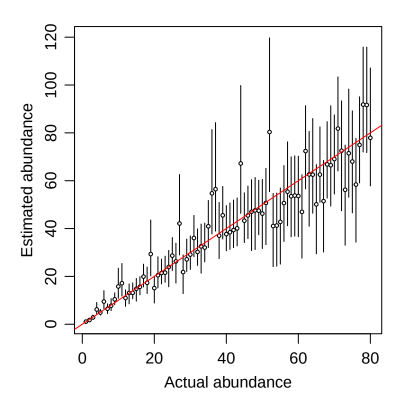


Figure II: Demonstration of agreement between actual and estimated abundances. Actual (simulated) abundances are on the x-axis, which the y-axis shows estimated abundances (error bars are 95% maximum credible intervals). The simulation study is described in the supplement.

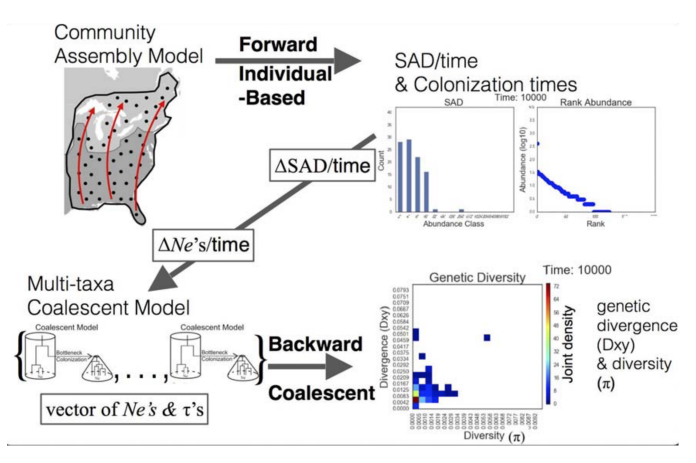


Figure III: The gimmeSAD π pipeline. The forward time models involves multi-regional expansion generating local abundance distributions over time with geterogeneity in colonization times. These temporally dynamic local abundances are re-scaled into local n_e distributions over time to generate multi-species genetic data the the coalescent, which is summarized here with a time-dependent joint spectrum of genetic diversity statistics.