



ANNUAL REVIEWS **Further**

Click [here](#) to view this article's online features:

- Download figures as PPT slides
- Navigate linked references
- Download citations
- Explore related articles
- Search keywords

Ecological Opportunity and Adaptive Radiation

James T. Stroud^{1,2} and Jonathan B. Losos³

¹Department of Biological Sciences, Florida International University, Miami, Florida 33199

²Fairchild Tropical Botanic Garden, Coral Gables, Florida 33156;
email: jamesTstroud@gmail.com

³Museum of Comparative Zoology and Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, Massachusetts 02138; email: jlosos@oeb.harvard.com

Annu. Rev. Ecol. Evol. Syst. 2016. 47:507–32

First published online as a Review in Advance on September 30, 2016

The *Annual Review of Ecology, Evolution, and Systematics* is online at ecolsys.annualreviews.org

This article's doi:
10.1146/annurev-ecolsys-121415-032254

Copyright © 2016 by Annual Reviews.
All rights reserved

Keywords

ecological opportunity, adaptive radiation, key innovations, adaptive landscape, character displacement, niche construction

Abstract

The process of adaptive radiation—the proliferation of species from a single ancestor and diversification into many ecologically different forms—has been of great interest to evolutionary biologists since Darwin. Since the middle of the last century, ecological opportunity has been invoked as a potential key to understanding when and how adaptive radiation occurs. Interest in the topic of ecological opportunity has accelerated as research on adaptive radiation has experienced a resurgence, fueled in part by advances in phylogenetic approaches to studying evolutionary diversification. Nonetheless, what the term actually means, much less how it mechanistically leads to adaptive diversification, is currently debated; whether the term has any predictive value or is a heuristic useful only for post hoc explanation also remains unclear. Recent recognition that evolutionary change can occur rapidly and on a timescale commensurate with ecological processes suggests that it is time to synthesize ecological and evolutionary approaches to the study of community assembly and evolutionary diversification.

Ecological opportunity:

the availability of ecologically accessible resources that may be evolutionarily exploited

Adaptive radiation:

evolutionary divergence of members of a single phylogenetic lineage into a variety of different adaptive forms (Futuyma 1998)

Diversification:

the evolution of phenotypically and reproductively distinct species in a clade

Key innovation: the evolution of a trait that allows a species to interact with the environment in a novel way

1. INTRODUCTION

Ecological opportunity, “loosely defined as a wealth of evolutionarily accessible resources little used by competing taxa” (Schluter 2000, p. 69), has long been thought to be an important—perhaps necessary—prerequisite for adaptive radiation. This view can be traced back to Darwin and was canonized by Simpson (1953); more recently, Schluter (2000) highlighted it as the centerpiece of understanding when and how adaptive radiation occurs. Certainly, the importance of ecological opportunity—as recognized by the absence of ecologically similar species—seems indisputable given the widespread occurrence of adaptive radiations after mass extinction events and on islands, lakes, and mountaintops.

But what actually is ecological opportunity? How is it identified and quantified? How mechanistically does it lead to evolutionary radiation, and why in some cases and not others? And, in cases where radiation has not occurred, can one assess whether the cause was lack of opportunity or some other explanation?

1.1. A Historical Perspective: From Conception to Modern Utilization

The notion of ecological opportunity as a prerequisite for adaptive radiation stems from the pioneering work of George G. Simpson (1953), in which he defined ecological opportunity as an environment experienced by an ancestral species that was previously “occupied by organisms for some reason competitively inferior to the entering group or must be empty” (p. 207). Other prominent evolutionary biologists of the time—notably David Lack (1947)—held similar views; namely they supported the idea that a release from the biotic constraints of competing taxa in depauperate environments allows for the proliferation of species, increased ecological specialization, and associated phenotypic diversification.

Simpson (1953) suggested that ecological opportunity could become available in a number of ways. The geographic colonization of isolated areas lacking competitors has provided ample examples of the production of adaptive radiations. Indeed, many classic examples of adaptive radiation, spurred by the ecological opportunity of colonization, are from islands (see Section 2). The absence of mainland species on islands provides opportunities for increased ecological specialization into niches not filled by competing taxa. A similar adaptive landscape may be encountered in a post-mass extinction environment, where ecologically similar competitors may again be absent. Conversely, the appearance of new resources rather than the absence of other species may be the source of opportunity. Finally, Simpson (1953) attached great importance to the evolution of unique phenotypes that allowed a species to interact with the environment in a novel way (i.e., key innovations; Miller 1949, Liem 1973). Simpson suggested that such traits have the potential to allow a species to diversify into a variety of niches not previously accessible and not occupied by competitor species.

Interest in the role of ecological opportunity in adaptive radiation has erupted in the last two decades (**Figure 1**). The term ecological opportunity, which in an evolutionary sense refers to the availability of ecologically accessible resources that may be evolutionarily exploited, was rarely used in the literature prior to 1995, but since then its use has increased almost exponentially. Certainly, a major cause of this increasing interest in ecological opportunity was Schluter’s book (Schluter 2000), which emphasized the importance of ecological radiation as a major contributor to evolutionary diversification. Of particular importance to the study of both adaptive radiation and ecological opportunity has been the explosion of molecular phylogenetic studies, a trend that began shortly before Schluter’s book was published (e.g., Givnish & Sytsma 1997). The proliferation of densely sampled, time-calibrated phylogenies of many groups has not only set the stage for modern

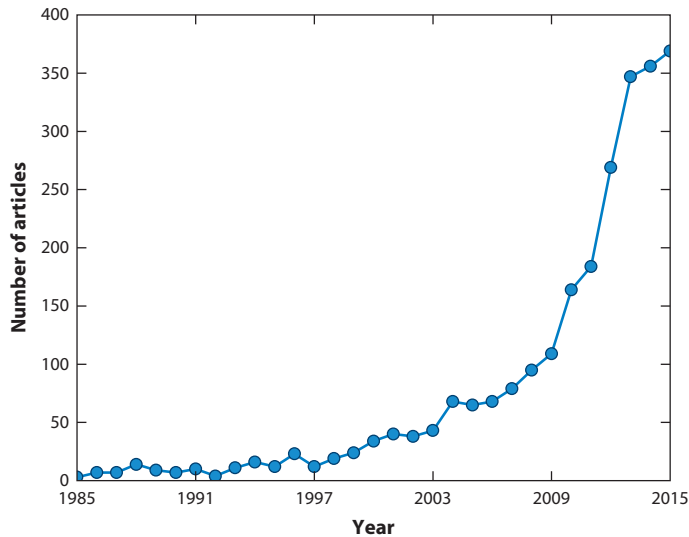


Figure 1

Trends in research on ecological opportunity as reflected in the number of articles published annually. The literature search was conducted using the Google Scholar database on January 20, 2016. To find articles that include “ecological opportunity,” the following specifications were used: Find articles with the exact phrases “ecological opportunity” and “adaptive radiation” and with at least one of the words “ecology” or “evolution.” Searches were run individually for every year 1950–2015; owing to a lack of studies before 1985, data from only 1985–2015 are shown.

investigation into the causes of evolutionary diversification but has also revealed macroevolutionary patterns consistent with a determinative role for ecological opportunity (discussed in Section 2).

2. WHAT IS THE EVIDENCE THAT ECOLOGICAL OPPORTUNITY PROMOTES ADAPTIVE RADIATION?

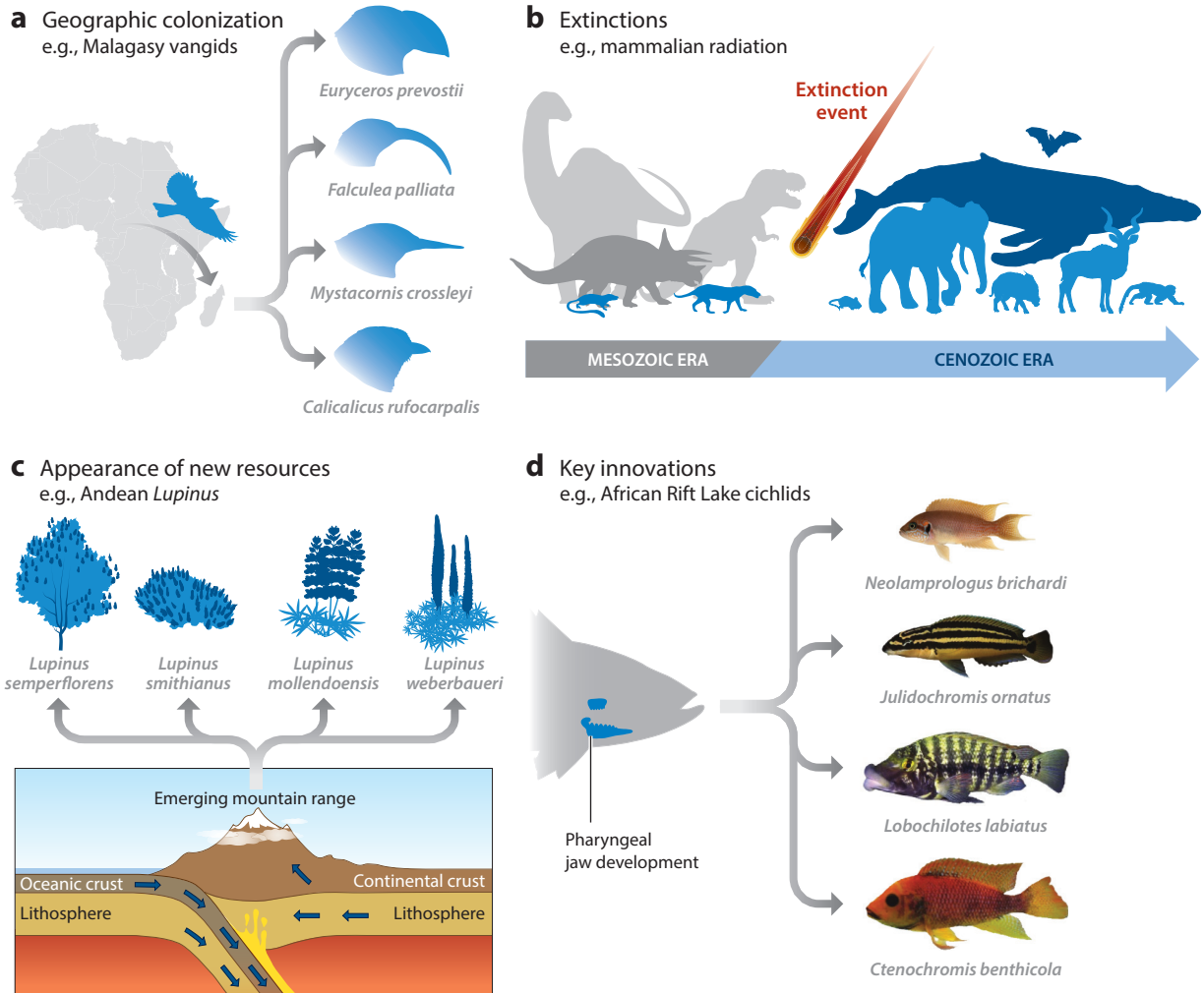
Simpson (1953) suggested that for an adaptive radiation to occur, an ancestral species must have geographical, ecological, and evolutionary access to ecological opportunity. By geographical access, he meant simply that a species must find itself in a location where ecological opportunity occurs. Ecological access requires the availability of resources not usurped by competitively superior species, and evolutionary access means that a species has the ability to utilize the resources. In Simpson’s view, adaptive radiation results when the ancestral species already has access along two of these axes and then an event occurs that provides access to the third. For example, adaptive radiation might ensue when a species colonizes an area in which resources are available and it has the evolutionary capability of diversifying to take advantage of them. Alternatively, it may also result when the species is already present in an area but an extinction event removes an ecologically dominant incumbent (see **Figure 2**).

2.1. Geographic Colonization

The fact that many textbook examples of adaptive radiation occur on islands is testament to the importance of geographical access as a precursor to adaptive radiation. The remote Hawaiian archipelago has been particularly fruitful, generating exceptional radiations of birds (Lovette et al. 2002), plants (Givnish et al. 2009), insects (Roderick & Gillespie 1998), and arachnids

Disparity: the difference among taxa of a phenotypic trait or traits

(Gillespie 2004, 2015). In landlocked lakes, the aquatic equivalent of islands, the same patterns can be observed: the African Rift Lakes have produced multiple spectacular radiations of cichlid fishes (Sturmbauer et al. 2011, Brawand et al. 2014, Seehausen 2014), and on the Indonesian island of Sulawesi, an adaptive radiation of silversides in Lake Matano has produced exceptional morphological diversity (Pfaender et al. 2010, 2016). One trait that both emergent islands and newly formed lakes have in common is the absence of competitors, such as those that may be present in areas from which an ancestral species has arrived (Carlquist 1974, Givnish 1997a, Leigh et al. 2007). A release from competitors provides the opportunity for radiating species to utilize ecological niches from which they were previously blocked. Similarly, a release from predators may also allow the use of habitats or resources previously inaccessible, thus spurring adaptive diversification (Schluter 1988, Benkman 1991, Heinen et al. 2013, Runemark et al. 2014). As a result of the release from the biotic pressures experienced on the mainland, island radiations often have much higher ecological and phenotypic disparity than their mainland sister clades (Schluter 2000, Lovette et al. 2002). Some have argued that adaptive radiation may primarily be an island phenomenon (Webb et al. 2002); however many extensive evolutionary radiations have also



occurred in mainland situations when ecological circumstances have permitted (Mouton & Van Wyk 1997, Tanentzap et al. 2015).

2.2. Extinctions and the Appearance of New Resources

In some cases, ancestral species encounter ecological opportunity within the ancestral range. Two ways in which this may occur are in the aftermath of an extinction event and following the appearance of new habitats or resources.

The presence of an incumbent clade usually prevents another clade from diversifying (Rosenzweig & McCord 1991), just as an incumbent species often prevents colonization of an ecologically similar species (the priority effect) (MacArthur 1972, Chase 2007). The evolutionary inhibition of incumbents has been demonstrated in laboratory microbial experiments in which the extent of adaptive radiation was significantly constrained when ecologically similar species were present (Brockhurst et al. 2007).

Given the inhibitory role of incumbents, it is not surprising that the elimination of competing taxa presents an ecological opportunity for those lineages able to survive an extinction event (Erwin 2007, 2015; Chen & Benton 2012). The paleontological record abounds with examples of surviving taxa rapidly radiating after extinction events, sometimes to the extent that the ensuing morphological disparity matches, or even exceeds, that observed in their extinct predecessors (Hull 2015). Surviving lineages of a clade may diversify only into niches that were previously occupied by members of that clade prior to the extinction event (Foote 1996, Ciampaglio 2002), such as in the rapid recovery of ammonoid diversity following the Permian–Triassic mass extinction (McGowan 2004); more commonly, however, lineages may radiate into niches previously filled by competitors that succumbed to extinction (Foote 1999, Friedman 2010).

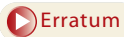
Mass extinction events, which remove entire or large proportions of taxonomic groups, provide ample evidence for the ecological opportunity hypothesis. For example, the Cretaceous–Paleogene mass extinction of the nonavian dinosaurs and other archosaurs resulted in an explosive radiation

Figure 2

(a) The colonization of an isolated area (e.g., an island or lake) can provide a release from competition and predation pressures, allowing a clade to diversify into a variety of ecological niches from which they were previously blocked. For example, the colonization of Madagascar resulted in a spectacular adaptive radiation of vangid birds (Yamagishi et al. 2001, Jönsson et al. 2012). Additional examples include Galápagos finches (Grant & Grant 2008), Hawaiian lobeliads (Givnish et al. 2009), African Rift Lake cichlids (Seehausen 2006), Hawaiian spiders (Gillespie 2004, 2015), Sulawesi silversides (Pfaender et al. 2010), and Caribbean *Anolis* lizards (Losos 2009). (b) Lineages able to survive extinction events may be presented with access to ecological space previously occupied by members of their own clade or by other competitors. For example, following the Cretaceous–Paleogene mass extinction of the nonavian dinosaurs and other archosaurs, ancestral birds and placental mammals gained access to previously unavailable ecological space and rapidly radiated (Hull 2015); mammals did exist, and some groups originated, prior to the extinction event, but this is simplified for heuristic purposes. An additional example is the Permian–Triassic radiation of ammonoids (McGowan 2004). (c) The appearance of new resources may provide ecological opportunities for species that can utilize them. For example, the emergence of the Andean mountains in the tropics provided new cool, high-elevation habitats that spurred adaptive radiations when colonized by ancestral *Espeletia* and *Lupinus* species (Monasterio & Sarmiento 1991, Hughes & Eastwood 2006). Additional examples include flowering plants and phytophagous insects (Labandeira & Sepkoski 1993, Bronstein et al. 2006) and the appearance of grasses and grazing horses (MacFadden 2005). (d) The evolution of a feature that allows a lineage to interact with the environment in a novel way may provide the ability to utilize formerly unavailable resources. The evolution of the pharyngeal jaw, for example, allowed African Rift Lake cichlids to diversify into a wide range of trophic specialists (Liem 1973, Mabuchi et al. 2007), although this has recently been debated (Seehausen 2006). Additional examples include the evolution of flight in bats and birds (Cracraft 1990, Sears et al. 2006, Simmons et al. 2008), phytophagy in insects (Farrell 1998), nectar spurs in *Aquilegia* columbines (Hodges & Arnold 1995, Ree 2005), and toepads in lizards (Williams & Peterson 1982). It should be noted that in all panels (a–d) the structures of all radiating groups are not meant to accurately reflect true phylogenetic relationships.

of birds and placental mammals (Smith et al. 2010, Erwin 2015; see Hull 2015 for a comprehensive review), which took advantage of the release from competitive and predatory pressures.

Access to new ecological opportunity can also occur in situ if new resources appear within an area. The evolution of flowering plants, for example, may have spurred the diversification of phytophagous insects (Labandeira & Sepkoski 1993, Bronstein et al. 2006; however, see McKenna et al. 2009). Similarly, soon after the Miocene appearance of grasslands in North America, horses (family Equidae) radiated rapidly, diversifying in body size, limb morphology, and dentition suitable for grazing on abrasive vegetation (MacFadden 2005). Alternatively, the appearance of novel environmental conditions, such as the emergence of new mountain ranges, can be the catalyst for ecological opportunity. For example, during the uplift of the Andean mountains, high-elevation páramo habitats formed as a type of above-treeline alpine tundra (Madriñán et al. 2013). Elevational tracking of this novel habitat zone by ancestral *Espeletia* and *Lupinus* plant species spurred their diverse radiations (Monasterio & Sarmiento 1991, Rauscher 2002, Hughes & Eastwood 2006, Hughes & Atchison 2015).



2.3. Key Innovations

In addition to geographical and ecological access, for a clade to radiate, it must have the evolutionary capability to diversify to take advantage of the available resources. Discussion of this topic has focused on the evolution of so-called key innovations—features that allow a lineage to interact with the environment in a novel way and thus may provide the ability, hitherto unavailable, to utilize available resources (Hunter 1998, Galis 2001, Rabosky 2014). For example, the evolution of flight in birds, bats, and pterosaurs presumably provided access to aerial prey resources, leading to subsequent diversification and specialization to different aspects of the aerial realm (Wellborn & Langerhans 2015). In other cases, the key innovation can provide access by minimizing the restricting effect of predators. For example, the evolution of brightly colored phenotypes to advertise toxicity in tropical dendrobatid poison frogs decreased the need for predator-induced hiding behavior and therefore allowed species to utilize habitats and resources that were previously inaccessible (Santos et al. 2003, Summers 2003, Arbuckle & Speed 2015).

Many possible examples of a key innovation leading to adaptive radiation have been suggested, but making a compelling case for a cause-and-effect relationship between the evolution of a trait and subsequent diversification is difficult in any particular instance. One solution is to investigate potential key innovations that have evolved several times to test for a general relationship between evolution of a trait and subsequent diversification (Mitter et al. 1988, de Queiroz 2002): Examples include hypocone dentition in mammals (Hunter & Jernvall 1995), toepads in lizards (Williams & Peterson 1982, Larson & Losos 1996), nectar spurs in *Aquilegia* columbines (Hodges & Arnold 1995, Ree 2005), phytophagy in insects (Mitter et al. 1988, Farrell 1998), and pharyngeal jaws in fish (Mabuchi et al. 2007). Of course, key innovations are often not a single trait but a complex of several traits—the evolution of one trait may set the stage for subsequent evolution of other traits, the combination of which in turn triggers radiation (Donoghue 2005, Marazzi et al. 2012, Werner et al. 2014). For example, features that appeared during the evolution of wings in birds evolved across multiple nodes in the phylogeny; thus, the wing as a key innovation was not a singular evolutionary event but the culmination of many evolutionary changes over millions of years (Cracraft 1990).

Two caveats must be kept in mind when evaluating claims of key innovations relative to adaptive radiation. First, key innovations may not lead to adaptive radiation. Many clades have evolved features that allow them to interact with the environment in a fundamentally different way yet have not diversified to an appreciable extent. For example, archer fish (*Toxotes* spp.) have evolved

the ability to shoot water from their mouths up to 3 m to dislodge insects perched on overhanging vegetation (Schuster et al. 2006, Burnette & Ashley-Ross 2015), yet the archer fish family (Toxotidae) comprises only seven morphologically similar species (Allen 2004). Similarly, *Aneides* salamanders evolved a novel foot structure that provides great climbing ability, but *Aneides* contains only six very similar species (Baum & Larson 1991). Among mammals, the elongated digits that evolved in parallel in the Madagascan aye-ayes (*Daubentonia madagascariensis*), Papuan striped possums (*Dactylopsila* sp.), and—although now extinct—the early Tertiary apatemyids (*Heterohyus* sp.) provide the ability to locate and extract prey from crevices in the manner of a woodpecker (Koenigswald & Schierning 1987, Erickson 1991, Rawlins & Handasyde 2002), yet each also failed to radiate to any great extent. The possession of a key innovation may not lead to adaptive radiation because either ecological opportunity is not available—the innovation may provide access to a very narrow spectrum of resources—or the clade does not have the evolutionary flexibility to diversify (see Section 5 for a discussion of the failure for adaptive radiation to occur).

The second caveat is that many studies have considered any trait that subsequently leads to species diversification to be a key innovation (e.g., von Hagen & Kadereit 2003, Ree 2005, Erkens et al. 2012, Silvestro et al. 2014). However, the observation that a clade is species rich does not indicate that it is adaptively diverse, much less that the trait allowed clade members to interact with the environment in a new way. More generally, some adaptive radiations contain few species, and some species-rich clades exhibit little diversity in ecological form (i.e., little adaptive disparity) (Givnish 1997b, Losos & Mahler 2010). Consequently, adaptive disparity and species richness are not necessarily related, and different terms are needed for traits that promote one type of diversification or the other (although in some cases, a trait may have both effects). The term key innovations (Miller 1949) refers to those traits that lead to interacting with the environment in a different way; another term is needed to refer to traits that increase the rate of species diversification.

2.4. Testing the Ecological Opportunity Hypothesis with Phylogenies

In several ways, molecular phylogenetics has been instrumental in the resurgence of the study of adaptive radiation and the role that ecological opportunity plays (Glor 2010). For example, molecular studies have revealed that many biotas previously assumed to be comprised of multiple ancestral lineages are the result of diversification of a single clade: Examples include the Malagasy vangas (Vangidae) (Yamagishi et al. 2001, Jönsson et al. 2012, Reddy et al. 2012); Australian, African, and global corvids (Barker et al. 2004; Jönsson et al. 2011, 2015); Lake Victoria cichlids (Meyer et al. 1990); and Hawaiian lobeliads (Givnish et al. 2009). Molecular phylogenies have also proved useful in clarifying temporal patterns of diversification, which is important for understanding the pace of diversification (Rabosky 2009) and recognizing potential catalysts for adaptive radiation (Donoghue 2005, Glor 2010).

In addition, molecular phylogenies have been used to directly test the hypothesis that rates of adaptive radiation are related to available ecological opportunity. Early studies simply looked at the rate of species proliferation through time as a clade diversified; assuming that as a clade became more species rich ecological opportunity would decrease, these studies tested the prediction that this declining opportunity would lead to a slowdown in the pace of diversification through time¹ (Schluter 2000, Freckleton & Harvey 2006). More recently, researchers have directly tested the

¹Note that adaptive radiation does not necessarily entail a burst of diversification at the outset; whether such a temporal pattern occurs as part of a radiation is a hypothesis to be tested, rather than part of the definition (Givnish 2015).

Ecological character displacement:

the divergence of sympatric species to minimize ecological overlap

relationship between opportunity, estimated as the interspecific morphological variety inferred to have existed at a given time, and the rate of ecomorphological diversification (Mahler et al. 2010). The prediction in these studies is that morphological diversification should decrease as clades become more ecomorphologically diverse (Arakaki et al. 2011, Burbrink et al. 2012, Svensson & Calsbeek 2012, Hughes et al. 2013, Slater 2015). When applying this framework to Greater Antillean *Anolis* lizard radiations, for example, time-calibrated phylogenetic methods revealed both a rapid early accumulation of lineages and bursts of phenotypic evolution (Mahler et al. 2010), a pattern which is common across taxonomic groups and geographic regions [e.g., the fossil record (Foote 1997), birds (Rabosky & Lovette 2008), plants (Agrawal et al. 2009), fishes (Near et al. 2012), and mammals (Schenk et al. 2013)].

3. HOW DOES ECOLOGICAL OPPORTUNITY MECHANISTICALLY LEAD TO ADAPTIVE RADIATION?

Picture a pregnant rodent washing ashore on a lush tropical island full of plant and arthropod life but lacking herbivores, granivores, and carnivores. The many available resources in the form of foods and habitats would constitute a wealth of ecological opportunity, and with luck and avoidance of inbreeding depression, the resulting population would quickly become well established. Scenarios such as this embody the first step in adaptive radiation: A population finding itself in the presence of great opportunity. But how does this ecological cornucopia translate into evolutionary diversification?

Adaptive radiation entails evolution in two dimensions: the proliferation of an initial ancestral species into multiple descendant species and the divergence of these species to adapt to an array of different ecological conditions, which we henceforth refer to as niches (for discussion of this long-lived and contentious term, see Chase & Leibold 2003). Whether speciation and adaptive divergence are a sequential or simultaneous process is an outstanding question in macroevolution.

3.1. The Classic Scenario: Interspecific Competition

The classic scenario postulates speciation occurring first, followed by subsequent divergence. The archipelago model—exemplified by adaptive radiation in Darwin’s finches (Grant & Grant 2008, 2014)—is a prime example: An ancestral finch initially colonizes one of the Galápagos Islands, and subsequently individuals from that population colonize another island. In allopatry, the two populations diverge to the extent that they are substantially or completely reproductively isolated (or would be were they to occur in sympatry); such reproductive isolation can evolve for many reasons, such as genetic drift or as an incidental by-product of divergence resulting from different adaptive or sexual selection pressures (Gittenberger 1991, Schluter 2000, Price 2008, Wagner et al. 2012). At the same time, some degree of adaptive divergence occurs as the two nascent species adapt to differences between the islands. Subsequently, colonization from one island to the other brings the two nascent species into sympatry. Once the populations reach their carrying capacities, they may compete for resources. Given the different types of available resources, the two populations may take advantage of this ecological opportunity and diverge in resource use; this resource partitioning permits the species to minimize interspecific competition. Assuming that the populations can coexist long enough (i.e., that competitive exclusion does not lead to the extinction of one; MacArthur & Levins 1967, Slatkin 1980, Gomulkiewicz & Holt 1995), natural selection may then cause the species to phenotypically diverge to adapt to their new resource utilization regime. This is the process of ecological character displacement (Brown & Wilson 1956). Natural selection against hybrids can also lead to the perfection of reproductive isolation if

it was incomplete prior to sympatry (the process of reinforcement; Blair 1955). Multiple cycles of such divergence in isolation followed by character displacement in sympatry can lead to a diverse adaptive radiation.

Not too long ago, both character displacement and reinforcement were thought by some to be unlikely on both theoretical and empirical grounds, but in recent years, these concerns have diminished and both are now generally considered to commonly occur (Schluter 2000; Dayan & Simberloff 2005; Grether et al. 2009; Pfennig & Pfennig 2012a,b; Stuart & Losos 2013). Evidence from both laboratory studies of microorganisms (Rainey & Travisano 1998, Tyerman et al. 2008, Bailey & Kassen 2012, Le Gac et al. 2012) and field studies of trait shifts in nature (Grant & Grant 2006, Pfennig et al. 2006, Goldberg et al. 2012, Stuart et al. 2014) continue to provide compelling evidence for character displacement when ecological opportunity is present. Additionally, an increased movement of species to regions outside of their native range in the Anthropocene is providing ample opportunities to observe species in the early stages of secondary contact and co-existence, setting the stage for many new observational studies of character displacement (Weber & Strauss 2016).

Reinforcement:

the evolution of traits that minimize hybridization between incipient species

3.2. Predation

Traditionally, interspecific competition has been considered the driving force behind adaptive radiation in the presence of ecological opportunity, but other mechanisms may be important as well. In particular, predation (defined here as consumption of one individual by another, thus including herbivory and parasitism) can cause populations to shift their resource use; in the presence of ecological opportunity, predation, in theory, may be a potent force driving adaptive radiation (Langerhans 2007).

Predation can play a role at several different stages of adaptive radiation. On one hand, allopatric populations may diverge adaptively not due to differences in resource availability but as a result of experiencing different predation pressures. Damselfly larvae, mosquitofish, sticklebacks, and zooplankton, for example, exhibit divergence in behavior, habitat use, and morphology depending on the types of predators to which they are exposed (Marchinko 2009, Strobbe et al. 2011, Walsh & Post 2011, Giery & Layman 2015, Giery et al. 2015).

On the other hand, divergence can also occur between sympatric species as they adapt in different ways to predation by a common predator (Allen et al. 2013). Such predator-driven divergence can be particularly potent when it drives species into different niches and lifestyles that also lead to differences in resource use. For example, the evolution of body armor can affect locomotion, which in turn may alter how animals can forage and acquire resources (Langerhans 2009, Broeckhoven et al. 2015).

Predator-driven prey divergence can result in evolutionary patterns similar to those resulting from interspecific competition. When prey species share a predator, an increase in the population size of one prey species may lead to a larger population of the predator, which in turn would lead to a reduction in the population of the second prey species. The result is that the population sizes of the two species would be negatively related, just as occurs with interspecific competition (Holt 1977). And just as with interspecific competition, prey species may diverge in habitat or resource use to minimize vulnerability to the shared predator, leading to the same pattern of character displacement as produced by interspecific competition. The process by which predation may lead to the same types of ecological and evolutionary response as competition has been termed competition for enemy free space (Jeffries & Lawton 1984) or apparent competition (Holt 1977).

In these ways, predation-driven divergent selection could lead to adaptive radiation in the presence of ecological opportunity. However, the extent to which predation drives adaptive radiation

remains unresolved, and few examples have been documented (Vamosi 2005, Langerhans 2007, Anderson & Langerhans 2015).

In addition to predation, other interspecific interactions may be important in stimulating adaptive radiation. For example, mutualisms can promote the coexistence of closely related species and may also lead to new ecological opportunities (Anacker & Strauss 2014, Weber & Strauss 2016).

3.3. Sympatric Speciation

The archipelago model of adaptive radiation by allopatric speciation, as described earlier for Darwin's finches, is easy to envision. Speciation may occur on different islands, with species subsequently coming into contact via dispersal. However, any biogeographic or historical setting in which populations become geographically isolated may produce allopatric speciation and serve as the first stage in the adaptive radiation process. Allopatric speciation has historically been considered the predominant process by which speciation occurs, at least in animals if not plants, and is one reason that the traditional view of adaptive radiation invokes speciation in allopatry followed by divergence in sympatry.

The alternative view of the relationship between ecological opportunity and adaptive radiation envisions speciation and adaptive diversification occurring in concert in one place without an allopatric stage. In this view, the ancestral population first expands its resource use, taking advantage of the variety of available resources, the lack of predators, or a combination of both (Parent & Crespi 2009, Yoder et al. 2010, Wellborn & Langerhans 2015). This niche expansion in the absence of other species is referred to as ecological release² (Yoder et al. 2010, Wellborn & Langerhans 2015). Subsequent to niche expansion, disruptive selection operates on the population, favoring individuals better suited to utilize specific resources but working against those intermediate individuals not well adapted to any particular resource (Wellborn & Langerhans 2015). As subpopulations become well adapted to specific resources, the relative fitness of intermediate phenotypes decreases. As a result, strong selection favors individuals that mate assortatively, leading to increasing reproductive isolation between the subpopulations, which may eventually become different species (Dieckmann & Doebeli 1999, Kondrashov & Kondrashov 1999, Doebeli & Dieckmann 2000). This represents one model of sympatric speciation, now often studied under the rubric of ecological speciation (Nosil 2012).

For much of the latter half of the twentieth century, sympatric speciation was theorized to be very unlikely, the primary criticism being that interbreeding between diverging subpopulations would tend to homogenize the populations and prevent the establishment of assortative mating (Mayr 1963, Felsenstein 1981, Coyne & Price 2000, Coyne & Orr 2004). However, in recent years the development of new theoretical frameworks and the discovery of a slew of suggestive examples have caused the pendulum to swing in the opposite direction, and such sympatric ecological speciation is now considered a likely phenomenon by many (e.g., Bolnick & Fitzpatrick 2007, Givnish 2010, Bird et al. 2012, Nosil 2012, Mullen & Shaw 2014). In some respects, a model of adaptive radiation via sympatric speciation is more parsimonious than invoking the existence of an intermediate allopatric stage for speciation, for which there often is no independent evidence. This argument has been made particularly forcefully for adaptive radiations occurring on small islands or in lakes where the opportunity for allopatry is not obvious (Schliewen et al. 1994, Barluenga et al. 2006, Kautt et al. 2012, Martin & Feinstein 2014; but see Martin et al. 2015). Nonetheless, whether adaptive radiation often proceeds by sympatric speciation remains much

²Ecological release can occur in the classic allopatric scenario as well but is not a necessary part of it.

debated (Schluter 2009, Nosil 2012, Henning & Meyer 2014, Martin et al. 2015), and the number of putative cases of adaptive radiation via this mechanism of speciation remains low.

4. IS ECOLOGICAL OPPORTUNITY NECESSARY FOR ADAPTIVE RADIATION?

Clearly, ecological opportunity often leads to adaptive radiation. But is it a necessary prerequisite? Can adaptive radiation occur in the absence of preexisting ecological opportunity by members of a diversifying clade either by wresting previously unavailable resources away from other taxa that had been using them or by creating their own opportunity?

4.1. Competitive Replacement

The paleontological literature is replete with proposed examples of one clade diversifying at the expense of an existing clade, outcompeting the extant species, and forcing the clade into evolutionary decline—a process known as active clade replacement (see discussions in Benton 1996, Jablonski 2008). The paleontological signal of such a situation would be negatively correlated species diversity patterns, one clade rising in diversity while the other diminishes. Perhaps the most convincing examples are those observed among post-Paleozoic cyclostome and cheilostome bryozoans (Lidgard et al. 1993, Sepkoski et al. 2000) and the multiple diversifying clades of canids and felids driving two canid subfamilies extinct in North America (Silvestro et al. 2015).

However, most purported examples of competitive clade replacement have fallen out of favor (Wilson et al. 2012, Benton et al. 2014) and have been replaced with a scenario whereby the incumbent taxon declined prior, and for reasons unrelated, to the diversification of a second group (Rosenzweig & McCord 1991; Benton 1996, 2009; Van Valkenburgh 1999; Brusatte et al. 2008). Subsequent diversification of the second clade after extinction of the first exemplifies postextinction radiation driven by new ecological opportunities, an alternative process known as passive clade replacement (Alroy 1999, Archibald 2011). For example, the idea that the rise of the dinosaurs came at the expense of other archosaurs has lost support, replaced by the idea that the process represented an extended passive replacement facilitated by dramatic changes in the global environment (Brusatte et al. 2008, 2010; Benton et al. 2014).

Moreover, the scenario of one clade competitively diversifying and eliminating a second clade is called into question by the rarity in which even a single species competitively eliminates another throughout its range today. The vast number of human introductions of species to areas outside their native ranges has brought together countless numbers of competitively similar species with no prior history of interaction (Sax et al. 2007). Nonetheless, few examples exist of one species causing the global (as opposed to local) extinction of another species as a result of interspecific competition (Davis 2013). By contrast, exposure to novel predation and disease from introduced species has led to the global extinction of many native species (Sax et al. 2007, Roy et al. 2012, Kraus 2015). Although novel competition can cause rapid reductions in population sizes of native species and may result in extinctions over a longer temporal scale than those caused by predation or disease (McCune et al. 2013), given the relative rarity of documented competition-caused extinctions of even single species, the possibility of an adaptively radiating clade eliminating an entire clade of established species by way of resource usurpation appears unlikely.

4.2. Self-Propagating Radiations: Niche Construction

Ecological opportunity thus seems usually necessary as a prerequisite for adaptive radiation. In most well-studied cases, such opportunity exists prior to adaptive radiation, but an alternative is

that clades create their own opportunity as they radiate (for extensive discussion, see Odling-Smee et al. 2003, 2013). Ecological opportunity may be generated through the creation of novel niches, whereby the environment is modified in some way—either biotically or abiotically—such that access to newly created resources results (Matthews et al. 2014).

This process of niche construction may occur in three ways. First, as a clade radiates, the presence of multiple species provides ecological opportunities that other clade members can exploit (which we term intracade niche construction). The traditional view is that as a clade diversifies, ecological resources become increasingly limited, and the rate of diversification slows down. But an alternative possibility is that the more species there are in a clade, the more opportunities there are for predators or mutualists (Whittaker 1977, Tokeshi 2009, Erwin 2015, Wellborn & Langerhans 2015). Most ecological communities comprise species from many clades; however, when members of a single radiation are extremely diverse ecologically (usually on islands or in lakes), this accelerating effect of species richness may have an evolutionary component: As the clade radiates, it may create additional opportunities, spurring further radiation, thus creating further opportunities, and so on.

This scenario is most likely when adaptive radiations transcend trophic levels, producing radiation members that consume other members of the same clade (Schluter 2000). Indeed, the evolution of a species that preys on other members of a clade has been observed in a diversity of taxa. For example, freshwater fishes specialized to feed on the scales of other species in its clade have evolved in both Bahamian pupfish (*Cyprinodon* sp.) (Martin & Wainwright 2013a, Martin & Feinstein 2014) and African Rift Lake cichlid (Koblmüller et al. 2007, Takahashi et al. 2007, Seehausen 2014) radiations. Additionally, intracade niche construction has also been demonstrated experimentally in microbial laboratory systems. For example, by manipulating the spatial complexity of the structural environment available to an ancestral population of *Escherichia coli*, initial colonizers of uninhabited fragment patches modified the environment such that divergence occurred when subsequent colonizers were able to exploit a different, previously unavailable niche, and adaptive radiation ensued (Habets et al. 2006). Similarly, several microbial studies have shown the evolution of a trophic specialist able to metabolize the waste product of an ancestral species (Kassen 2009). Though these examples demonstrate that adaptive radiations may create additional opportunity during diversification, the general pattern in adaptive radiations of decreasing diversification and ecomorphological adaptation rates through time (e.g., Agrawal et al. 2009, Mahler et al. 2010; but see Slater 2015) suggests that the creation of additional ecological opportunities within a clade during radiation is rare.

Second, concurrent radiation of two clades may create opportunities for ecological diversification in one or both clades (which we term intercade niche construction). For example, a radiation driven by competition or predation may create new ecological opportunities for a coexisting clade that may adapt and use constituent members of the first as a resource. This may spur further complementary radiation of the first clade as members seek to avoid being exploited by the second. The diversification of apple maggot flies, for example, is paralleled by radiation of the parasites that prey upon them (Forbes 2009), much like the diversification of herbivorous weevils occurring in concert with that of flowering plants (McKenna et al. 2009). In some cases, coevolutionary dynamics are more complicated, involving one clade escaping harmful biotic pressures of another and diversifying as selection is relaxed, thus providing the basis of the escape and radiation theory of plant–herbivore coevolution (Ehrlich & Raven 1964) and Vermeij’s (1987) theory of evolution and escalation.

A third way in which radiations may create their own ecological opportunity is through the construction of new ecological opportunities via alterations to the physical environment: so-called ecosystem engineering (Jones et al. 1994, 1997; Wright & Jones 2006; Odling-Smee et al. 2013).

Ecosystem engineers affect resource availability for other species by physically modifying the abiotic or biotic characteristics of an environment (Jones et al. 1994, Erwin 2008). Ecosystem engineering could lead to adaptive radiation if an ecosystem engineer (or clade of ecosystem engineers) creates new environments that prompt adaptive radiation in another clade. For example, the unique and complex habitat structure built by coral reefs provided ecological opportunities for specialization and subsequent diversification of tetraodontiform fishes (Alfaro et al. 2007), just as the evolution of burrowing led to increased bioturbation and subsequent diversification of marine animal lineages able to exploit the newly constructed habitat zone during the Cambrian explosion (Erwin 2008). These examples, of course, do not illustrate a clade creating its own ecological opportunities. However, the possibility exists that a radiating clade could take advantage of opportunities created by an ecosystem-engineering clade member. One example may be the exceptional adaptive radiation of lobeliad plants in Hawaii. After initial radiation by a colonial species, forest plants that evolved to withstand hot, unshaded habitats may have acted as ecological engineers by subsequently providing suitable environmental conditions, which allowed for the evolution of shade-tolerant subcanopy species (Givnish et al. 2004, 2009).

5. WHY DOES ECOLOGICAL OPPORTUNITY NOT ALWAYS RESULT IN ADAPTIVE RADIATION?

Although ecological opportunity is the usual stimulus for adaptive radiation, some clades fail to radiate in the presence of an apparent ecological opportunity (Wilson 1992, Losos 2010, Losos & Mahler 2010; depauperons, *sensu* Donoghue & Sanderson 2015, Weber & Strauss 2016). For example, in the Galápagos, Darwin's finches are the only clade of birds that radiated to any appreciable extent (Valente et al. 2015). Similarly, on Caribbean islands, *Anolis* lizards have repeatedly and independently diversified on separate islands, yet few other clades of lizards have followed suit, despite having had the same temporal opportunity to do so (Crother & Guyer 1996, Thorpe et al. 2008). The presence of ecological opportunity, therefore, does not guarantee that a clade will radiate.

A clade might fail to radiate in the apparent presence of ecological opportunity for several reasons. One explanation is that ecological opportunity is not actually present. For example, the failure of some clades to radiate on islands may be because earlier-arriving clades already usurped available resources. This can be observed in the case of muroid rodents, in which early colonizers had inhibitory effects on the ultimate diversity of later colonizers within independent radiations around the world (Schenk et al. 2013). Alternatively, ecological opportunity itself may have been misidentified. The spectrum of available resources that constitutes an ecological opportunity for one species may not be the same as that required by a different species. For example, the diversity of seed sizes and shapes available on the Galápagos may have constituted opportunity to an ancestral finch species, but the range of insects may not have provided diverse opportunities for insectivorous birds, thus explaining differences in evolutionary diversification among Galápagos birds (Arbogast et al. 2006, Grant & Grant 2008, Rundell & Price 2009, Valente et al. 2015).

Even in the presence of ecological opportunity, however, adaptive radiation may not occur for two reasons. First, as discussed previously, adaptive radiation requires both proliferation of species and diversification in resource use to fill different ecological niches. Consequently, if a species is unable to speciate then adaptive radiation cannot occur (Wellborn & Langerhans 2015). For example, many types of organisms—e.g., birds, lizards, snails, and snakes—almost never speciate on islands smaller than a threshold size (Coyne & Price 2000, Losos & Schluter 2000, Losos & Parent 2009, Kisel & Barraclough 2010, Pyron & Burbrink 2014). Failure of these species—such as *Pinaroloxias inornata*, the single species of Darwin finch on Cocos Island, or *Anolis* lizards in the

Evolvability: the capacity to generate heritable phenotypic variation

Lesser Antilles—to radiate on these ecologically diverse islands may be a result of their inability to speciate.

The second reason that a clade may fail to radiate in the presence of ecological opportunity is that it does not have the ability to evolve readily into diverse forms (Schluter 2000): Clades that lack such evolvability will change more slowly or not at all, whereas those that can readily change will be capable of adapting to new circumstances (Arbogast et al. 2006, Adamowicz et al. 2008, Wellborn & Langerhans 2015). Several factors could account for differences in evolvability. For example, species with greater modularity—i.e., in which different aspects of the phenotype can evolve independently—may be able to diversify to a greater extent than species in which phenotypic components are less independent (Vermeij 1973, Kirschner et al. 1998, Rutherford & Lindquist 1998, Clune et al. 2013). Phenotypic and behavioral plasticity may also be important factors in determining levels of evolvability (Baldwin 1896, Draghi & Whitlock 2012, Snell-Rood 2013). Plasticity may allow a species to exist in conditions that otherwise would be unsuitable, possibly providing sufficient time for subsequent genetic adaptations to the new ecological environment (reviewed in West-Eberhard 2003). However, identifying the evolutionary role of evolvability may be difficult as evolvability itself may evolve rather than being characteristic of entire clades (Zaman et al. 2014). For example, hybridization events increased evolvability in populations of African Lake Malawi cichlids (Parsons et al. 2011).

The observation that some clades radiate more than others (Carlquist 1974) suggests that evolvability and propensity to speciate may be important in determining whether adaptive radiation occurs. Some clades appear to have high evolvability and propensity to radiate regardless of the environment: Examples include the Hawaiian honeycreepers and Darwin's finches, which have both radiated extensively on their respective archipelagos, as have their sister taxa on the mainland (Burns et al. 2002, Lovette et al. 2002). By contrast, the Hawaiian thrushes and the Galápagos mockingbirds have not radiated on either islands or the mainland despite existing there for the same length of time (Lovette et al. 2002, Arbogast et al. 2006, Grant & Grant 2008). Cichlid fishes show a similar pattern, with some clades frequently radiating and others not (Seehausen 2014). These consistent differences among clades suggest either intrinsic differences in evolvability or speciation propensity may exist among taxa.

Alternatively, some clades that appear to readily radiate on islands remain relatively depauperate elsewhere. For example, *Tetragnatha* spiders and aglycyderid weevils radiated to a greater extent on Hawaii than on the mainland (Paulay 1994, Gillespie 2015), and cichlid fish diversity in Africa is much higher in lakes than rivers (Genner et al. 2015). In these cases, extrinsic circumstances appear to be more important than intrinsic propensities in determining whether radiation results.

6. CAN ECOLOGICAL OPPORTUNITY BE IDENTIFIED AND QUANTIFIED A PRIORI OR IS IT ONLY OF HEURISTIC VALUE?

Ecological opportunity is usually recognized after it has occurred: A clade that has experienced an adaptive radiation is identified and then its history is assessed to see whether ecological opportunity was present early on. Alternatively, an event that generates ecological opportunity—such as a mass extinction event or creation of a new island or mountain range—is identified, and clades are studied to see if any have subsequently radiated. The consensus is clear that ecological opportunity usually precedes adaptive radiation.

Nonetheless, two key questions remain. First, as discussed in the previous section, in those cases in which a clade did not radiate, is a lack of ecological opportunity the explanation? Second, looking forward, can we identify species or clades currently experiencing ecological opportunity and thus might radiate in the future? In other words, can ecological opportunity be identified

independently of the occurrence of adaptive radiation? Can ecological opportunity be measured? Does the concept have predictive value, or is it just a useful heuristic for explaining adaptive radiation after the fact (Losos 2010)?

Answering these questions requires a means of measuring ecological opportunity, which is not straightforward. In this respect, the concept of ecological opportunity is plagued by the same difficulties as the empty niche concept (Chase & Leibold 2003, Losos & Mahler 2010). Not only are both hard to identify in the absence of species that fill or take advantage of them but also it is difficult to demonstrate that resources are ever truly underutilized by some member of a community, even if not by a member of a focal clade.

Ecological opportunity represents an adaptive landscape with many vacant peaks (Simpson 1953). An approach to testing for the existence of ecological opportunity might involve estimating selection on an adaptive landscape (Fear & Price 1998, Schluter 2000, Arnold et al. 2001; the concept of adaptive landscapes is extensively reviewed in Svensson & Calsbeek 2012). If it is possible to generate a variety of phenotypes (e.g., through hybridization), it may be possible to detect unoccupied adaptive peaks, which suggests the existence of ecological opportunity. In other words, an alternative way of making a living may exist but awaits the evolution of a species able to utilize that ecological space.

Of course, estimating the adaptive landscape, particularly in the context of evolutionary radiation, is fraught with difficulty. In particular, the presence of other species—e.g., competitors and predators—will alter the shape of the landscape, potentially causing peaks to appear or vanish compared with a landscape estimated in their absence. Consequently, in the context of investigating whether ecological opportunity exists, the landscape will need to be investigated in the presence of other species. This approach, however, will only suffice to identify currently existing opportunity. Once speciation occurs, the landscape may shift as a result of the presence of a new species. Hence, estimating how long opportunity persists during the course of a radiation will be very difficult and is beyond any work that has been conducted to date.

Only a few studies have quantified the adaptive landscape experienced by the constituent species of a given adaptive radiation (e.g., Case 1979, Schluter & Grant 1984, Pfaender et al. 2016). For example, Martin & Wainwright (2013b) estimated the adaptive landscape by measuring selection on a variable population produced by hybridizing three sympatric Bahamian pupfish species (*Cyprinodon* sp.). Their study confirmed the existence of two peaks corresponding to the phenotypes of two of the three species. Similarly, Arnegard et al. (2014) explored the adaptive landscape of threespine stickleback fish (*Gasterosteus aculeatus* complex) using hybrids of two species adapted to different habitats. Hybrids were distributed across a great range of morphologies, with subsequent fitness (using growth rate as a proxy) highest at two adaptive peaks corresponding to morphologies most similar to those of the two parental species. Extending this approach to test for the existence of unoccupied adaptive peaks available to species potentially experiencing ecological opportunity is a logical next step in this research direction.

FUTURE ISSUES

Simpson's (1953) suggestion that ecological opportunity is the impetus for adaptive radiation is well supported. Now would seem to be the time for the field to move beyond documenting whether a relationship exists between ecological opportunity and adaptive radiation and to investigate the underlying mechanistic basis for the relationship. Additionally, investigations should assess when and why the two are sometimes uncoupled, either because adaptive radiation can occur without preexisting opportunity or, conversely, because radiations sometimes fail to follow from the existence of opportunity.

New tools—from comparative genomics to the ability to estimate adaptive landscapes and conduct evolutionary experiments in the field—now provide the means to further refine these questions. However, at the most fundamental level, a detailed understanding of the natural history of study organisms will remain crucially important to the development and interpretation of studies attempting to synthesize ecological and evolutionary processes (Greene 1986; Grant & Grant 2008, 2014). We expect the study of ecological opportunity to continue to blossom in the near future.

1. Adaptive landscapes. Recent empirical studies have estimated the shape of adaptive landscapes with multiple coexisting species in an adaptive radiation. Further studies attempting to quantify ecological opportunity (i.e., the presence of multiple adaptive peaks on a landscape) will continue to develop our understanding of the nature of ecological opportunity and will allow us to see how opportunity varies among taxa, among areas, and through time. Further, a more detailed macroevolutionary theory of how the adaptive landscape itself evolves will be important to further bridge the pattern–process divide (Arnold et al. 2001, Svensson & Calsbeek 2012, Wellborn & Langerhans 2015).
2. Genomics. Investigations into the genomic structure of several well-studied adaptive radiations have already begun [e.g., Darwin’s finches (Almen et al. 2015), African Rift Lake cichlids (Wagner et al. 2013, Brawand et al. 2014), *Heliconius* butterflies (Heliconius Genome Consort. et al. 2012, Supple et al. 2013)]. Studies that synthesize research on adaptive landscapes with genomics will have the potential to present a clearer understanding of the phenotype–fitness and genotype–phenotype relationships and will be instrumental in understanding the genetic basis of how and why ecological opportunity is exploited and adaptive radiation occurs.
3. Niche construction. As a better understanding of eco-evolutionary feedback relationships continues to develop, opportunities to provide empirical tests for hypotheses of niche construction will arise. Although long supported in the paleontological literature (Odling-Smee et al. 2013), and more recently by microbial laboratory studies (Rainey & Travisano 1998, Habets et al. 2006), little evidence from contemporary ecological studies exists (although see Matthews et al. 2016).
4. Species introductions. The global movement of species in the Anthropocene has provided unparalleled opportunities to observe novel ecological and evolutionary scenarios (Wellborn & Langerhans 2015, Weber & Strauss 2016). For example, movement of species to areas with no, or few, ecological competitors may provide situations analogous to conditions first experienced by ancestral species in an adaptive radiation, potentially presenting the introduced species with ecological opportunity. It may be particularly valuable to compare introduced species from clades that have radiated elsewhere with those from clades that have failed to radiate to examine the role of inherent evolvability of a clade when responding to ecological opportunity. Advances in methods for predicting areas vulnerable to invasion may also provide opportunities to identify ecological opportunity a priori. For example, the identification of young diversifying clades—particularly those in areas strongly associated with the production of adaptive radiations such as islands—may suggest that ecological opportunity is still present and therefore more vulnerable to ecological and evolutionary exploitation by novel colonists.

5. Global extinctions. As we enter the Sixth Mass Extinction in the Anthropocene (Ceballos et al. 2015), we are being presented with the first large-scale opportunity to study ecological and evolutionary responses to biodiversity loss (Wellborn & Langerhans 2015). Despite an undoubted catastrophe for global biodiversity, this evolutionary research opportunity is unprecedented. For example, current global amphibian rates of extinction are four orders of magnitude higher than expected background rates (Alroy 2015). The IUCN Red List currently classifies >30% of frogs and toads (Anura) and >49% of newts and salamanders (Urodela)—representing a wide range of ecological diversity—as either extinct or threatened with extinction (Catenazzi 2015). Their loss may present ecological opportunities for lineages able to exploit resources vacated by the loss of a large proportion of an entire taxonomic group. Of course, this discussion assumes that whatever is causing extinction—often habitat loss—would not preclude the addition of new species to a community. In many cases, this will not be true.
6. Latitudinal diversity gradient. The latitudinal gradient in species diversity is well supported (Hillebrand 2004); however, the underlying mechanisms which have led to it remain unclear. Given the existence of a positive relationship between ecological opportunity and speciation rates, identifying a relationship between ecological opportunity and latitude may present one way to understand the evolution of the latitudinal diversity gradient (Schluter 2016). A more comprehensive understanding of the geographic nature of ecological opportunity would help us understand how it may be important in shaping global patterns of species diversity.

DISCLOSURE STATEMENT

The authors are not aware of any affiliations, memberships, funding, or financial holdings that might be perceived as affecting the objectivity of this review.

ACKNOWLEDGMENTS

We thank Kenneth Feeley, Sean Giery, Evan Rehm, Rachel Hillyer, Sterling Nesbitt, Timothy Perez, Marjorie Weber, Thomas Givnish, and Sally Otto for their thoughtful comments and discussions on previous drafts of this manuscript. J.T.S. was supported by a Florida International University Doctoral Evidence Acquisition Fellowship.

LITERATURE CITED

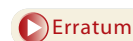
- Adamowicz SJ, Purvis A, Wills MA. 2008. Increasing morphological complexity in multiple parallel lineages of the Crustacea. *PNAS* 105(12):4786–91
- Agrawal AA, Fishbein M, Haltschke R, Hastings AP, Rabosky DL, Rasmann S. 2009. Evidence for adaptive radiation from a phylogenetic study of plant defenses. *PNAS* 106(43):18067–72
- Alfaro ME, Santini F, Brock CD. 2007. Do reefs drive diversification in marine teleosts? Evidence from the pufferfishes and their allies (order Tetraodontiformes). *Evolution* 61(9):2104–26
- Allen GR. 2004. *Toxotes kimberleyensis*, a new species of archerfish (Pisces: Toxotidae) from fresh waters of Western Australia. *Rec. Aust. Mus.* 56:225–30
- Allen WL, Baddeley R, Scott-Samuel NE, Cuthill IC. 2013. The evolution and function of pattern diversity in snakes. *Behav. Ecol.* 24(5):1237–50

- Almen MS, Lamichhaney S, Berglund J, Grant BR, Grant PR, et al. 2015. Adaptive radiation of Darwin's finches revisited using whole genome sequencing. *BioEssays* 38(1):14–20
- Alroy J. 1999. The fossil record of North American mammals: evidence for a Paleocene evolutionary radiation. *Syst. Biol.* 48(1):107–18
- Alroy J. 2015. Current extinction rates of reptiles and amphibians. *PNAS* 112(42):13003–8
- Anacker BL, Strauss SY. 2014. The geography and ecology of plant speciation: range overlap and niche divergence in sister species. *Proc. R. Soc. B* 281:20132980
- Anderson CM, Langerhans RB. 2015. Origins of female genital diversity: Predation risk and lock-and-key explain rapid divergence during an adaptive radiation. *Evolution* 69(9):2452–67
- Arakaki M, Christin P-A, Nyffeler R, Lendel A, Eggli U, et al. 2011. Contemporaneous and recent radiations of the world's major succulent plant lineages. *PNAS* 108(20):8379–84
- Arbogast BS, Drovetski SV, Curry RL, Boag PT, Seutin G, et al. 2006. The origin and diversification of Galápagos mockingbirds. *Evolution* 60(2):370–82
- Arbuckle K, Speed MP. 2015. Antipredator defenses predict diversification rates. *PNAS* 2015:1–6
- Archibald JD. 2011. *Extinction and Radiation: How the Fall of Dinosaurs Led to the Rise of Mammals*. Baltimore, MD: Johns Hopkins Univ. Press
- Arnegard ME, McGee MD, Matthews B, Marchinko KB, Conte GL, et al. 2014. Genetics of ecological divergence during speciation. *Nature* 511(7509):307–11
- Arnold SJ, Pfrender ME, Jones AG. 2001. The adaptive landscape as a conceptual bridge between micro- and macroevolution. *Genetica* 112–113:9–32
- Bailey SF, Kassen R. 2012. Spatial structure of ecological opportunity drives adaptation in a bacterium. *Am. Nat.* 180(2):270–83
- Baldwin JM. 1896. A new factor in evolution. *Am. Nat.* 30(354):441–51
- Barker FK, Cibois A, Schikler P, Feinstein J, Cracraft J. 2004. Phylogeny and diversification of the largest avian radiation. *PNAS* 101(30):11040–45
- Barluenga M, Stölting KN, Salzburger W, Muschick M, Meyer A. 2006. Sympatric speciation in Nicaraguan crater lake cichlid fish. *Nature* 439(7077):719–23
- Baum DA, Larson A. 1991. Adaptation reviewed: a phylogenetic methodology for studying character macroevolution. *Syst. Zool.* 40:1–18
- Benkman CW. 1991. Predation, seed size partitioning and the evolution of body size in seed-eating finches. *Evol. Ecol.* 5(2):118–27
- Benton MJ. 1996. On the nonprevalence of competitive replacement in the evolution of tetrapods. In *Evolutionary Paleobiology*, ed. JW Valentine, D Jablonski, pp. 185–210. Chicago: Univ. Chicago Press
- Benton MJ. 2009. The Red Queen and the Court Jester: species diversity and the role of biotic and abiotic factors through time. *Science* 323(5915):728–32
- Benton MJ, Forth J, Langer MC. 2014. Models for the rise of the dinosaurs. *Curr. Biol.* 24(2):R87–95
- Bird CE, Fernandez-Silva I, Skillings DJ, Toonen RJ. 2012. Sympatric speciation in the post “modern synthesis” era of evolutionary biology. *Evol. Biol.* 39(2):158–80
- Blair WF. 1955. Size differences as a possible isolation mechanism in *Microbyla*. *Am. Nat.* 89:297–301
- Bolnick DI, Fitzpatrick BM. 2007. Sympatric speciation: models and empirical evidence. *Annu. Rev. Ecol. Evol. Syst.* 38(1):459–87
- Brawand D, Wagner CE, Li YL, Malinsky M, Keller I, et al. 2014. The genomic substrate for adaptive radiation in African cichlid fish. *Nature* 513:375–81
- Brockhurst MA, Colegrave N, Hodgson DJ, Buckling A. 2007. Niche occupation limits adaptive radiation in experimental microcosms. *PLOS ONE* 2(2):e193
- Broeckhoven C, Diedericks G, Mouton PIFN. 2015. What doesn't kill you might make you stronger: functional basis for variation in body armour. *J. Anim. Ecol.* 84:1213–21
- Bronstein JL, Alarcón R, Geber M. 2006. The evolution of plant-insect mutualisms. *New Phytol.* 172(3):412–28
- Brown WL, Wilson EO. 1956. Character displacement. *Syst. Zool.* 5(2):49–64
- Brusatte SL, Benton MJ, Ruta M, Lloyd GT. 2008. Superiority, competition, and opportunism in the evolutionary radiation of dinosaurs. *Science* 321(5895):1485–88
- Brusatte SL, Nesbitt SJ, Irmis RB, Butler RJ, Benton MJ, Norell MA. 2010. The origin and early radiation of dinosaurs. *Earth-Sci. Rev.* 101(1):68–100

- Burbrink FT, Ruane S, Pyron RA. 2012. When are adaptive radiations replicated in areas? Ecological opportunity and unexceptional diversification in West Indian dipsadine snakes (Colubridae: Alsophiini). *J. Biogeogr.* 39(3):465–75
- Burnette MF, Ashley-Ross MA. 2015. One shot, one kill: the forces delivered by archer fish shots to distant targets. *Zoology* 118(5):302–11
- Burns KJ, Hackett SJ, Klein NK. 2002. Phylogenetic relationships and morphological diversity in Darwin's finches and their relatives. *Evolution* 56(6):1240–52
- Carlquist S. 1974. *Island Biology*. New York/London: Columbia Univ. Press
- Case TJ. 1979. Character displacement and coevolution in some *Cnemidophorus* lizards. *Fortschr. Zool.* 25:235–82
- Catenazzi A. 2015. State of the world's amphibians. *Annu. Rev. Environ. Resour.* 40(1):91–119
- Ceballos G, Ehrlich PR, Barnosky AD, García A, Pringle RM, Palmer TM. 2015. Accelerated modern human-induced species losses: entering the sixth mass extinction. *Sci. Adv.* 1:e1400253
- Chase JM. 2007. Drought mediates the importance of stochastic community assembly. *PNAS* 104(44):17430–34
- Chase JM, Leibold MA. 2003. *Ecological Niches: Linking Classical and Contemporary Approaches*. Chicago: Univ. Chicago Press
- Chen Z-Q, Benton MJ. 2012. The timing and pattern of biotic recovery following the end-Permian mass extinction. *Nat. Geosci.* 5(6):375–83
- Ciampaglio CN. 2002. Determining the role that ecological and developmental constraints play in controlling disparity: examples from the crinoid and blastozoan fossil record. *Evol. Dev.* 4(3):170–88
- Clune J, Mouret JB, Lipson H. 2013. The evolutionary origins of modularity. *Proc. R. Soc. B* 280(1755):20122863
- Coyne JA, Orr HA. 2004. *Speciation*. Sunderland, MA: Sinauer
- Coyne JA, Price TD. 2000. Little evidence for sympatric speciation in island birds. *Evolution* 54(6):2166–71
- Cracraft J. 1990. The origin of evolutionary novelties: pattern and process at different hierarchical levels. In *Evolutionary Innovations*, ed. MH Nitecki, pp. 21–44. Chicago: Univ. Chicago Press
- Crother BI, Guyer C. 1996. Caribbean historical biogeography: Was the dispersal-vicariance debate eliminated by an extraterrestrial bolide? *Herpetologica* 52:440–65
- Davis MA. 2013. Invasive species. In *Grzimek's Animal Life Encyclopedia: Extinction*, ed. N MacLeod, pp. 779–87. Detroit, MI: Gale Publ.
- Dayan T, Simberloff D. 2005. Ecological and community-wide character displacement: the next generation. *Ecol. Lett.* 8(8):875–94
- de Queiroz A. 2002. Contingent predictability in evolution: key traits and diversification. *Syst. Biol.* 51(6):917–29
- Dieckmann U, Doebeli M. 1999. On the origin of species by sympatric speciation. *Nature* 400(6742):354–57
- Doebeli M, Dieckmann U. 2000. Evolutionary branching and sympatric speciation caused by different types of ecological interactions. *Am. Nat.* 156(4):77–101
- Donoghue MJ. 2005. Key innovations, convergence, and success: macroevolutionary lessons from plant phylogeny. *Paleobiology* 31(Suppl. 2):77–93
- Donoghue MJ, Sanderson MJ. 2015. Confluence, synnovation, and depauperons in plant diversification. *New Phytol.* 207:260–74
- Draghi JA, Whitlock MC. 2012. Phenotypic plasticity facilitates mutational variance, genetic variance, and evolvability along the major axis of environmental variation. *Evolution* 66(9):2891–902
- Ehrlich PR, Raven PH. 1964. Butterflies and plants: a study in coevolution. *Evolution* 18(4):586–608
- Erickson CJ. 1991. Percussive foraging in the aye-aye, *Daubentonia madagascariensis*. *Anim. Behav.* 41(5):793–801
- Erkens RHJ, Chatrou LW, Couvreur TLP. 2012. Radiations and key innovations in an early branching angiosperm lineage (Annonaceae; Magnoliales). *Bot. J. Linn. Soc.* 169(1):117–34
- Erwin DH. 2007. Increasing returns, ecological feedback and the Early Triassic recovery. *Palaeoworld* 16(1–3):9–15
- Erwin DH. 2008. Macroevolution of ecosystem engineering, niche construction and diversity. *Trends Ecol. Evol.* 23(6):304–10

- Erwin DH. 2015. Novelty and innovation in the history of life. *Curr. Biol.* 25(19):R930–40
- Farrell BD. 1998. “Inordinate fondness” explained: Why are there so many beetles? *Science* 281(5376):555–59
- Fear KK, Price T. 1998. The adaptive surface in ecology. *Oikos* 82(3):440–48
- Felsenstein J. 1981. Skepticism towards Santa Rosalia, or why are there so few kinds of animals? *Evolution* 35(1):124–38
- Foote M. 1996. Ecological controls on the evolutionary recovery of post-Paleozoic crinoids. *Science* 274(5292):1492–95
- Foote M. 1997. The evolution of morphological diversity. *Annu. Rev. Ecol. Syst.* 28:129–52
- Foote M. 1999. Morphological diversity in the evolutionary radiation of Paleozoic and post-Paleozoic crinoids. *Paleobiology* 25(2):1–115
- Forbes AA. 2009. Sequential sympatric speciation across tropic levels. *Science* 323:776–79
- Freckleton RP, Harvey PH. 2006. Detecting non-Brownian trait evolution in adaptive radiations. *PLOS Biol.* 4(11):2104–11
- Friedman M. 2010. Explosive morphological diversification of spiny-finned teleost fishes in the aftermath of the end-Cretaceous extinction. *Proc. R. Soc. B* 277(1688):1675–83
- Futuyma DJ. 1998. *Evolutionary Biology*. Sunderland, MA: Sinauer. 3rd ed.
- Galis F. 2001. Key innovations and radiations. In *The Character Concept in Evolutionary Biology*, ed. GP Wagner, pp. 581–605. San Diego: Academic
- Genner MJ, Ngatunga BP, Mzighani S, Smith A, Turner GF. 2015. Geographical ancestry of Lake Malawi’s cichlid fish diversity *Biol. Lett.* 11(6):20150232
- Giery ST, Layman CA. 2015. Interpopulation variation in a condition-dependent signal: predation regime affects signal intensity and reliability. *Am. Nat.* 186(2):187–95
- Giery ST, Layman CA, Langerhans RB. 2015. Anthropogenic ecosystem fragmentation drives shared and unique patterns of sexual signal divergence among three species of Bahamian mosquitofish. *Evol. Appl.* 8:679–91
- Gillespie RG. 2004. Community assembly through adaptive radiation in Hawaiian spiders. *Science* 303(5656):356–59
- Gillespie RG. 2015. Island time and the interplay between ecology and evolution in species diversification. *Evol. Appl.* 9(1):53–73
- Gittenberger E. 1991. What about non-adaptive radiation? *Biol. J. Linn. Soc.* 43(4):263–72
- Givnish TJ. 1997a. Adaptive plant evolution on islands: classical patterns, molecular data, new insights. In *Evolution on Islands*, ed. PR Grant, pp. 281–304. Oxford, UK: Oxford Univ. Press
- Givnish TJ. 1997b. Adaptive radiation and molecular systematics: Issues and approaches. See Givnish & Sytsma 1997, pp. 1–54
- Givnish TJ. 2010. Ecology of plant speciation. *Taxon* 59(5):1326–66
- Givnish TJ. 2015. Adaptive radiation versus ‘radiation’ and ‘explosive diversification’: why conceptual distinctions are fundamental to understanding evolution. *New Phytol.* 207(2):297–303
- Givnish TJ, Millam KC, Mast AR, Paterson TB, Theim TJ, et al. 2009. Origin, adaptive radiation and diversification of the Hawaiian lobeliads (Asterales: Campanulaceae). *Proc. R. Soc. B Biol. Sci.* 276(1656):407–16
- Givnish TJ, Montgomery RA, Goldstein G. 2004. Adaptive radiation of photosynthetic physiology in the Hawaiian lobeliads: light regimes, static light responses, and whole-plant compensation points. *Am. J. Bot.* 91(2):228–46
- Givnish TJ, Sytsma KJ. 1997. *Molecular Evolution and Adaptive Radiation*. Cambridge, UK: Cambridge Univ. Press
- Glor RE. 2010. Phylogenetic insights on adaptive radiation. *Annu. Rev. Ecol. Evol. Syst.* 41(1):251–70
- Goldberg EE, Lande R, Price TD. 2012. Population regulation and character displacement in a seasonal environment. *Am. Nat.* 179(6):693–705
- Gomulkiewicz R, Holt RD. 1995. When does evolution by natural selection prevent extinction? *Evolution* 49(1):201–7
- Grant PR, Grant BR. 2006. Evolution of character displacement in Darwin’s finches. *Science* 313(5784):224–26
- Grant PR, Grant BR. 2008. *How and Why Species Multiply. The Radiation of Darwin’s Finches*. Princeton, NJ: Princeton Univ. Press

- Grant PR, Grant BR. 2014. *40 Years of Evolution: Darwin's Finches on Daphne Major Island*. Princeton, NJ: Princeton Univ. Press
- Greene HW. 1986. Natural history and evolutionary biology. In *Predator-Prey Relationships: Perspectives and Approaches from the Study of Lower Vertebrates*, ed. ME Feder, GV Lauder, pp. 99–108. Chicago: Univ. Chicago Press
- Grether GF, Losin N, Anderson CN, Okamoto K. 2009. The role of interspecific interference competition in character displacement and the evolution of competitor recognition. *Biol. Rev.* 84(4):617–35
- Habets MGJL, Rozen DE, Hoekstra RF, de Visser JAGM. 2006. The effect of population structure on the adaptive radiation of microbial populations evolving in spatially structured environments. *Ecol. Lett.* 9(9):1041–48
- Heinen JL, Coco MW, Marcuard MS, White DN, Peterson MN, et al. 2013. Environmental drivers of demographics, habitat use, and behavior during a post-Pleistocene radiation of Bahamas mosquitofish (*Gambusia hubbsi*). *Evol. Ecol.* 27(5):971–91
- Heliconius Genome Consort., Dasmahapatra KK, Walters JR, Briscoe AD, Davey JW, et al. 2012. Butterfly genome reveals promiscuous exchange of mimicry adaptations among species. *Nature* 487(7405):94–98
- Henning F, Meyer A. 2014. The evolutionary genomics of cichlid fishes: explosive speciation and adaptation in the postgenomic era. *Annu. Rev. Genom. Hum. Genet.* 15:417–41
- Hillebrand H. 2004. On the generality of the latitudinal diversity gradient. *Am. Nat.* 163(2):192–211
- Hodges SA, Arnold ML. 1995. Spurring plant diversification: are floral nectar spurs a key innovation? *Proc. R. Soc. B* 262(1365):343–48
- Holt RD. 1977. Predation, apparent competition and the structure of prey communities. *Theor. Popul. Biol.* 12(2):197–229
- Hughes CE, Atchison GW. 2015. The ubiquity of alpine plant radiations: from the Andes to the Hengduan Mountains. *New Phytol.* 207(2):275–85
- Hughes CE, Eastwood R. 2006. Island radiation on a continental scale: exceptional rates of plant diversification after uplift of the Andes. *PNAS* 103(27):10334–39
- Hughes M, Gerber S, Wills MA. 2013. Clades reach highest morphological disparity early in their evolution. *PNAS* 110(34):13875–79
- Hull P. 2015. Life in the aftermath of mass extinctions. *Curr. Biol.* 25(19):R941–52
- Hunter JP. 1998. Key innovations and the ecology of macroevolution. *Trends Ecol. Evol.* 13(1):31–36
- Hunter JP, Jernvall J. 1995. The hypocone as a key innovation in mammalian evolution. *PNAS* 92(23):10718–22
- Jablonski D. 2008. Biotic interactions and macroevolution: extensions and mismatches across scales and levels. *Evolution* 62(4):715–39
- Jeffries MJ, Lawton JH. 1984. Enemy free space and the structure of ecological communities. *Biol. J. Linn. Soc.* 23:269–86
- Jones CG, Lawton JH, Shachak M. 1994. Organisms as ecosystem engineers. *Oikos* 69(3):373–86
- Jones CG, Lawton JH, Shachak M. 1997. Positive and negative effects of organisms as physical ecosystem engineers. *Ecology* 78(7):1946–57
- Jönsson KA, Fabre PH, Fritz SA, Etienne RS, Ricklefs RE, et al. 2012. Ecological and evolutionary determinants for the adaptive radiation of the Madagascan vangas. *PNAS* 109(17):6620–25
- Jönsson KA, Fabre P-H, Kennedy JD, Holt BG, Borregaard MK, et al. 2015. A supermatrix phylogeny of corvid passerine birds (Aves: Corvides). *Mol. Phylogenet. Evol.* 94(Part A):87–94
- Jönsson KA, Fabre P-H, Ricklefs RE, Fjeldsø J. 2011. Major global radiation of corvid birds originated in the proto-Papuan archipelago. *PNAS* 108(6):2328–33
- Kassen R. 2009. Toward a general theory of adaptive radiation: insights from microbial experimental evolution. *Ann. N. Y. Acad. Sci.* 1168:3–22
- Kautt AF, Elmer KR, Meyer A. 2012. Genomic signatures of divergent selection and speciation patterns in a “natural experiment”, the young parallel radiations of Nicaraguan crater lake cichlid fishes. *Mol. Ecol.* 21(19):4770–86
- Kirschner M, Kirschner M, Gerhart J, Gerhart J. 1998. Evolvability. *PNAS* 95(15):8420–27
- Kisel Y, Barraclough TG. 2010. Speciation has a spatial scale that depends on levels of gene flow. *Am. Nat.* 175(3):316–34



- Koblmüller S, Egger B, Sturmbauer C, Sefc KM. 2007. Evolutionary history of Lake Tanganyika's scale-eating cichlid fishes. *Mol. Phylogenet. Evol.* 44(3):1295–305
- Koenigswald WV, Schiønering H-P. 1987. The ecological niche of an extinct group of mammals, the early Tertiary apatemyids. *Nature* 326:595–97
- Kondrashov AS, Kondrashov FA. 1999. Interactions among quantitative traits in the course of sympatric speciation. *Nature* 400(6742):351–54
- Kraus F. 2015. Impacts from invasive reptiles and amphibians. *Annu. Rev. Ecol. Evol. Syst.* 46(1):75–97
- Labandeira CC, Sepkoski JJ. 1993. Insect diversity in the fossil record. *Science* 261(5119):310–15
- Lack D. 1947. *Darwin's Finches*. Cambridge, UK: Cambridge Univ. Press
- Langerhans RB. 2007. Evolutionary consequences of predation: avoidance, escape, reproduction, and diversification. In *Predation in Organisms: A Distinct Phenomenon*, ed. AMT Elewa, pp. 177–220. Heidelberg, Ger.: Springer-Verlag
- Langerhans RB. 2009. Trade-off between steady and unsteady swimming underlies predator-driven divergence in *Gambusia affinis*. *J. Evol. Biol.* 22(5):1057–75
- Larson A, Losos JB. 1996. Phylogenetic systematics of adaptation. In *Adaptation*, ed. MR Rose, GV Lauder, pp. 187–220. San Diego: Academic
- Le Gac M, Pluain J, Hindre T, Lenski RE, Schneider D. 2012. Ecological and evolutionary dynamics of coexisting lineages during a long-term experiment with *Escherichia coli*. *PNAS* 109(24):9487–92
- Leigh EG Jr., Hladik A, Hladik CM, Jolly A. 2007. The biogeography of large islands, or how does the size of the ecological theater affect the evolutionary play? *Rev. Écol.* 62:105–68
- Lidgard S, McKinney FK, Taylor PD. 1993. Competition, clade replacement, and a history of cyclostome and cheilostome bryozoan diversity. *Paleobiology* 19(3):352–71
- Liem KF. 1973. Evolutionary strategies and morphological innovations: cichlid pharyngeal jaws. *Syst. Biol.* 22:425–41
- Losos JB. 2009. *Lizards in an Evolutionary Tree: Ecology and Adaptive Radiation of Anoles*. Berkeley: Univ. Calif. Press
- Losos JB. 2010. Adaptive radiation, ecological opportunity, and evolutionary determinism. American Society of Naturalists E. O. Wilson Award address. *Am. Nat.* 175(6):623–39
- Losos JB, Mahler DL. 2010. Adaptive radiation: the interaction of ecological opportunity, adaptation, and speciation. In *Evolution since Darwin: The First 150 Years*, ed. MA Bell, DJ Futuyma, WF Eanes, JS Levinton, pp. 381–420. Sunderland, MA: Sinauer
- Losos JB, Parent CE. 2009. The speciation-area relationship. In *The Theory of Island Biogeography Revisited*, ed. JB Losos, RE Ricklefs, pp. 415–38. Princeton, NJ: Princeton Univ. Press
- Losos JB, Schluter D. 2000. Analysis of an evolutionary species-area relationship. *Nature* 408(6814):847–50
- Lovette IJ, Bermingham E, Ricklefs RE. 2002. Clade-specific morphological diversification and adaptive radiation in Hawaiian songbirds. *R. Soc. Proc. B* 269(1486):37–42
- Mabuchi K, Miya M, Azuma Y, Nishida M. 2007. Independent evolution of the specialized pharyngeal jaw apparatus in cichlid and labrid fishes. *BMC Evol. Biol.* 7:10
- MacArthur RH. 1972. *Geographical Ecology: Patterns in the Distribution of Species*. New York: Harper & Row
- MacArthur RH, Levins R. 1967. The limiting similarity, convergence, and divergence of coexisting species. *Am. Nat.* 101(921):377–85
- MacFadden BJ. 2005. Fossil horses—evidence for evolution. *Science* 307(5716):1728–30
- Madriñán S, Cortés AJ, Richardson JE. 2013. Páramo is the world's fastest evolving and coolest biodiversity hotspot. *Front. Genet.* 4:192
- Mahler DL, Revell LJ, Glor RE, Losos JB. 2010. Ecological opportunity and the rate of morphological evolution in the diversification of Greater Antillean anoles. *Evolution* 64(9):2731–45
- Marazzi B, Ané C, Simon MF, Delgado-Salinas A, Luckow M, Sanderson MJ. 2012. Locating evolutionary precursors on a phylogenetic tree. *Evolution* 66(12):3918–30
- Marchinko KB. 2009. Predation's role in repeated phenotypic and genetic divergence of armor in threespine stickleback. *Evolution* 63(1):127–38
- Martin CH, Cutler JS, Friel JP, Denning Touokong C, Coop G, Wainwright PC. 2015. Complex histories of repeated gene flow in Cameroon crater lake cichlids cast doubt on one of the clearest examples of sympatric speciation. *Evolution* 69(6):1406–22

- Martin CH, Feinstein LC. 2014. Novel trophic niches drive variable progress towards ecological speciation within an adaptive radiation of pupfishes. *Mol. Ecol.* 23(7):1846–62
- Martin CH, Wainwright PC. 2013a. On the measurement of ecological novelty: scale-eating pupfish are separated by 168 my from other scale-eating fishes. *PLOS ONE* 8(8):e71164
- Martin CH, Wainwright PC. 2013b. Multiple fitness peaks on the adaptive landscape drive adaptive radiation in the wild. *Science* 339(6116):208–11
- Matthews B, Aebischer T, Sullam KE, Seehausen O. 2016. Experimental evidence of an eco-evolutionary feedback during adaptive divergence. *Curr. Biol.* 26:1–7
- Matthews B, De Meester L, Jones CG, Ibelings BW, Bouma TJ, et al. 2014. Under niche construction: an operational bridge between ecology, evolution, and ecosystem science. *Ecol. Monogr.* 84(2):245–63
- Mayr E. 1963. *Animal Species and Evolution*. Cambridge, MA: Harvard Univ. Press
- McCune JL, Harrower WL, Avery-Gomm S, Brogan JM, Csergő AM, et al. 2013. Threats to Canadian species at risk: an analysis of finalized recovery strategies. *Biol. Conserv.* 166:254–65
- McGowan AJ. 2004. Ammonoid taxonomic and morphologic recovery patterns after the Permian Triassic. *Geology* 32(8):665–68
- McKenna DD, Sequeira AS, Marvaldi AE, Farrell BD. 2009. Temporal lags and overlap in the diversification of weevils and flowering plants. *PNAS* 106(17):7083–88
- Meyer A, Kocher TD, Basasibwaki P, Wilson AC. 1990. Monophyletic origin of Lake Victoria cichlid fishes suggested by mitochondrial DNA sequences. *Nature* 34:550–53
- Miller AH. 1949. Some ecologic and morphologic considerations in the evolution of higher taxonomic categories. In *Ornithologie als Biologische Wissenschaft, 28 Beiträge als Festschrift zum 60. Geburtstag von Erwin Stresemann*, ed. E Mayr, E Schüz, pp. 84–88. Heidelberg, Ger.: Carl Winter
- Mitter C, Farrell B, Wiegmann B. 1988. The phylogenetic study of adaptive zones: Has phytophagy promoted insect diversification? *Am. Nat.* 132(1):107–28
- Monasterio M, Sarmiento L. 1991. Adaptive radiation of *Espeletia* in the cold Andean tropics. *Trends Ecol. Evol.* 6(12):387–91
- Mouton PIFN, Van Wyk JH. 1997. Adaptive radiation in cordyliform lizards: an overview. *Afr. J. Herpetol.* 46(2):78–88
- Mullen SP, Shaw KL. 2014. Insect speciation rules: unifying concepts in speciation research. *Annu. Rev. Entomol.* 59:339–61
- Near TJ, Dornburg A, Kuhn KL, Eastman JT, Pennington JN, et al. 2012. Ancient climate change, antifreeze, and the evolutionary diversification of Antarctic fishes. *PNAS* 109(9):3434–39
- Nosil P. 2012. *Ecological Speciation*. Oxford, UK: Oxford Univ. Press
- Odling-Smee FJ, Laland KN, Feldman MW. 2003. *Niche Construction: The Neglected Process in Evolution*. Princeton, NJ: Princeton Univ. Press
- Odling-Smee J, Erwin DH, Palkovacs EP, Feldman MW, Laland KN. 2013. Niche construction theory: a practical guide for ecologists. *Q. Rev. Biol.* 88(1):3–28
- Parent CE, Crespi BJ. 2009. Ecological opportunity in adaptive radiation of Galápagos endemic land snails. *Am. Nat.* 174(6):898–905
- Parsons KJ, Son YH, Albertson RC. 2011. Hybridization promotes evolvability in African cichlids: connections between transgressive segregation and phenotypic integration. *Evol. Biol.* 38(3):306–15
- Paulay G. 1994. Biodiversity on oceanic islands: its origin and extinction. *Am. Zool.* 34:134–44
- Pfaender J, Hadiaty RK, Schliewen UK, Herder F. 2016. Rugged adaptive landscapes shape a complex, sympatric radiation. *Proc. R. Soc. B* 283(1822):20152342
- Pfaender J, Schliewen UK, Herder F. 2010. Phenotypic traits meet patterns of resource use in the radiation of “sharpfin” sailfin silverside fish in Lake Matano. *Evol. Ecol.* 24(5):957–74
- Pfennig DW, Pfennig KS. 2012a. *Evolution's Wedge: Competition and the Origins of Diversity*. Berkeley/Los Angeles: Univ. Calif. Press
- Pfennig DW, Pfennig KS. 2012b. Development and evolution of character displacement. *Ann. N. Y. Acad. Sci.* 1256(1):89–107
- Pfennig DW, Rice AM, Martin RA. 2006. Ecological opportunity and phenotypic plasticity interact to promote character displacement and species coexistence. *Ecology* 87(3):769–79

- Price TD. 2008. *Speciation in Birds*. Boulder, CO: Roberts
- Pyron RA, Burbrink FT. 2014. Ecological and evolutionary determinants of species richness and phylogenetic diversity for island snakes. *Glob. Ecol. Biogeogr.* 23(8):848–56
- Rabosky DL. 2009. Ecological limits on clade diversification in higher taxa. *Am. Nat.* 173(5):662–74
- Rabosky DL. 2014. Automatic detection of key innovations, rate shifts, and diversity-dependence on phylogenetic trees. *PLOS ONE* 9(2):e89543
- Rabosky DL, Lovette IJ. 2008. Density-dependent diversification in North American wood warblers. *Proc. R. Soc. B* 275:2363–71
- Rainey PB, Travisano M. 1998. Adaptive radiation in a heterogeneous environment. *Nature* 394(6688):69–72
- Rauscher JT. 2002. Molecular phylogenetics of the *Espeletia* complex (Asteraceae): evidence from nrDNA ITS sequences on the closest relatives of an Andean adaptive radiation. *Am. J. Bot.* 89(7):1074–84
- Rawlins DR, Handasyde KA. 2002. The feeding ecology of the striped possum *Dactylopsila trivirgata* (Marsupialia: Petauridae) in far north Queensland, Australia. *J. Zool.* 257(2):195–206
- Reddy S, Driskell A, Rabosky DL, Hackett SJ, Schulenberg TS. 2012. Diversification and the adaptive radiation of the vangas of Madagascar. *Proc. R. Soc. B* 279(1735):2062–71
- Ree RH. 2005. Detecting the historical signature of key innovations using stochastic models of character evolution and cladogenesis. *Evolution* 59(2):257–65
- Roderick GK, Gillespie RG. 1998. Speciation and phylogeography of Hawaiian terrestrial arthropods. *Mol. Ecol.* 7(4):519–31
- Rosenzweig ML, McCord RD. 1991. Incumbent replacement: evidence for long-term evolutionary progress. *Paleobiology* 17(3):202–13
- Roy HE, Adriaens T, Isaac NJB, Kenis M, Martin GS, et al. 2012. Invasive alien predator causes rapid declines of native European ladybirds. *Divers. Distrib.* 18:717–25
- Rundell RJ, Price TD. 2009. Adaptive radiation, nonadaptive radiation, ecological speciation and nonecological speciation. *Trends Ecol. Evol.* 24(7):394–99
- Runemark A, Brydegaard M, Svensson EI. 2014. Does relaxed predation drive phenotypic divergence among insular populations? *J. Evol. Biol.* 27(8):1676–90
- Rutherford SL, Lindquist S. 1998. Hsp90 as a capacitor for morphological evolution. *Nature* 396(6709):336–42
- Santos JC, Coloma LA, Cannatella DC. 2003. Multiple, recurring origins of aposematism and diet specialization in poison frogs. *PNAS* 100(22):12792–97
- Sax DF, Stachowicz JJ, Brown JH, Bruno JF, Dawson MN, et al. 2007. Ecological and evolutionary insights from species invasions. *Trends Ecol. Evol.* 22(9):465–71
- Schenk JJ, Rowe KC, Stepan SJ. 2013. Ecological opportunity and incumbency in the diversification of repeated continental colonizations by muroid rodents. *Syst. Biol.* 62(6):837–64
- Schliewen UK, Tautz D, Pääbo S. 1994. Sympatric speciation suggested by monophyly of crater lake cichlids. *Nature* 368(6472):629–32
- Schluter D. 1988. Character displacement and the adaptive divergence of finches on islands and continents. *Am. Nat.* 131(6):799–824
- Schluter D. 2000. *The Ecology of Adaptive Radiation*. Oxford, UK: Oxford Univ. Press
- Schluter D. 2009. Evidence for ecological speciation and its alternative. *Science* 323(5915):737–41
- Schluter D. 2016. Speciation, ecological opportunity, and latitude. *Am. Nat.* 187(1):1–18
- Schluter D, Grant PR. 1984. Determinants of morphological patterns in communities of Darwin's finches. *Am. Nat.* 123(2):175–96
- Schuster S, Wöhl S, Griebisch M, Klostermeier I. 2006. Animal cognition: How archer fish learn to down rapidly moving targets. *Curr. Biol.* 16(4):378–83
- Sears KE, Behringer RR, Rasweiler JJ, Niswander LA. 2006. Development of bat flight: morphologic and molecular evolution of bat wing digits. *PNAS* 103(17):6581–86
- Seehausen O. 2006. African cichlid fish: a model system in adaptive radiation research. *Proc. R. Soc. B* 273(1597):1987–98
- Seehausen O, Wagner CE. 2014. Speciation in freshwater fishes. *Annu. Rev. Ecol. Evol. Syst.* 45:621–51
- Sepkoski JJ, McKinney FK, Lidgard S. 2000. Competitive displacement among post-Paleozoic cyclostome and cheilostome bryozoans. *Paleobiology* 26(1):7–18

- Silvestro D, Antonelli A, Salamin N, Quental TB. 2015. The role of clade competition in the diversification of North American canids. *PNAS* 112(28):8684–89
- Silvestro D, Zizka G, Schulte K. 2014. Disentangling the effects of key innovations on the diversification of Bromelioideae (Bromeliaceae). *Evolution* 68(1):163–75
- Simmons NB, Seymour KL, Habersetzer J, Gunnell GF. 2008. Primitive early Eocene bat from Wyoming and the evolution of flight and echolocation. *Nature* 451(7180):818–21
- Simpson GG. 1953. *The Major Features of Evolution*. New York: Columbia Univ. Press
- Slater GJ. 2015. Not-so-early bursts and the dynamic nature of morphological diversification. *PNAS* 112(12):3595–96
- Slatkin M. 1980. Ecological character displacement. *Ecology* 61(1):163–77
- Smith FA, Boyer AG, Brown JH, Costa DP, Dayan T, et al. 2010. The evolution of maximum body size of terrestrial mammals. *Science* 330(6008):1216–19
- Snell-Rood EC. 2013. An overview of the evolutionary causes and consequences of behavioural plasticity. *Anim. Behav.* 85(5):1004–11
- Strobbe F, McPeck MA, De Block M, Stoks R. 2011. Fish predation selects for reduced foraging activity. *Behav. Ecol. Sociobiol.* 65(2):241–47
- Stuart YE, Campbell TS, Hohenlohe PA, Reynolds RG, Revell LJ, Losos JB. 2014. Rapid evolution of a native species following invasion by a congener. *Science* 346(6208):463–66
- Stuart YE, Losos JB. 2013. Ecological character displacement: glass half full or half empty? *Trends Ecol. Evol.* 28(7):402–8
- Sturmbauer C, Husemann M, Danley PD. 2011. Explosive speciation and adaptive radiation of East African cichlid fishes. In *Biodiversity Hotspots: Distribution and Protection of Conservation Priority Areas*, ed. FE Zachos, JC Habel, pp. 333–62. Berlin/Heidelberg: Springer-Verlag
- Summers K. 2003. Convergent evolution of bright coloration and toxicity in frogs. *PNAS* 100(22):12533–34
- Supple M, Papa R, Counterman B, McMillan WO. 2013. The genomics of an adaptive radiation: insights across the *Heliconius* speciation continuum. In *Ecological Genomics: Ecology and the Evolution of Genes and Genomes*, ed. CR Landry, N Aubin-Horth, pp. 249–71. Dordrecht, Neth.: Springer
- Svensson EI, Calsbeek R. 2012. *The Adaptive Landscape in Evolutionary Biology*. Oxford, UK: Oxford Univ. Press
- Takahashi R, Watanabe K, Nishida M, Hori M. 2007. Evolution of feeding specialization in Tanganyikan scale-eating cichlids: a molecular phylogenetic approach. *BMC Evol. Biol.* 7:195
- Tanentzap AJ, Brandt AJ, Smissen RD, Heenan PB, Fukami T, Lee WG. 2015. When do plant radiations influence community assembly? The importance of historical contingency in the race for niche space. *New Phytol.* 207:468–79
- Thorpe RS, Surget-Groba Y, Johansson H. 2008. The relative importance of ecology and geographic isolation for speciation in anoles. *Philos. Trans. R. Soc. B* 363(1506):3071–81
- Tokeshi M. 2009. *Species Coexistence: Ecological and Evolutionary Perspectives*. Oxford, UK: Blackwell
- Tyerman JG, Bertrand M, Spencer CC, Doebeli M. 2008. Experimental demonstration of ecological character displacement. *BMC Evol. Biol.* 8:34
- Valente LM, Phillimore AB, Etienne RS. 2015. Equilibrium and non-equilibrium dynamics simultaneously operate in the Galápagos islands. *Ecol. Lett.* 18(8):844–52
- Vamosi SM. 2005. On the role of enemies in divergence and diversification of prey: a review and synthesis. *Can. J. Zool.* 83(7):894–910
- Van Valkenburgh B. 1999. Major patterns in the history of carnivorous mammals. *Annu. Rev. Earth Planet. Sci.* 27:463–93
- Vermeij GJ. 1973. Adaptation, versatility, and evolution. *Syst. Biol.* 22(4):466–77
- Vermeij GJ. 1987. *Evolution and Escalation: An Ecological History of Life*. Princeton, NJ: Princeton Univ. Press
- von Hagen KB, Kadereit JW. 2003. The diversification of *Halenia* (Gentianaceae): ecological opportunity versus key innovation. *Evolution* 57(11):2507–18
- Wagner CE, Harmon LJ, Seehausen O. 2012. Ecological opportunity and sexual selection together predict adaptive radiation. *Nature* 487(7407):366–69

- Wagner CE, Keller I, Wittwer S, Selz OM, Mwaiko S, et al. 2013. Genome-wide RAD sequence data provide unprecedented resolution of species boundaries and relationships in the Lake Victoria cichlid adaptive radiation. *Mol. Ecol.* 22(3):787–98
- Walsh MR, Post DM. 2011. Interpopulation variation in a fish predator drives evolutionary divergence in prey in lakes. *Proc. R. Soc. B* 278(1718):2628–37
- Webb CO, Ackerly DD, McPeck MA, Donoghue MJ. 2002. Phylogenies and community ecology. *Annu. Rev. Ecol. Syst.* 33:475–505
- Weber MG, Strauss SY. 2016. Coexistence in close relatives: beyond competition and reproductive isolation in sister taxa. *Annu. Rev. Ecol. Syst.* 47:359–81
- Wellborn GA, Langerhans RB. 2015. Ecological opportunity and the adaptive diversification of lineages. *Ecol. Evol.* 5(1):176–95
- Werner GDA, Cornwell WK, Sprent JI, Kattge J, Kiers ET. 2014. A single evolutionary innovation drives the deep evolution of symbiotic N₂-fixation in angiosperms. *Nat. Commun.* 5:4087
- West-Eberhard MJ. 2003. *Developmental Plasticity and Evolution*. Oxford, UK: Oxford Univ. Press
- Whittaker RH. 1977. Evolution of species diversity in land communities [birds and vascular plants]. *Evol. Biol.* 10:1–67
- Williams EE, Peterson JA. 1982. Convergent and alternative designs in the digital adhesive pads of scincid lizards. *Science* 215(4539):1509–11
- Wilson EO. 1992. *The Diversity of Life*. Cambridge, MA: Belknap
- Wilson GP, Evans AR, Corfe IJ, Smits PD, Fortelius M, Jernvall J. 2012. Adaptive radiation of multituberculate mammals before the extinction of dinosaurs. *Nature* 483:457–60
- Wright JP, Jones CG. 2006. The concept of organisms as ecosystem engineers ten years on: Progress, limitations, and challenges. *BioScience* 56(3):203–9
- Yamagishi S, Honda M, Eguchi K, Thorstrom R. 2001. Extreme endemic radiation of the Malagasy vangas (Aves: Passeriformes). *J. Mol. Evol.* 53:39–46
- Yoder JB, Clancey E, Des Roches S, Eastman JM, Gentry L, et al. 2010. Ecological opportunity and the origin of adaptive radiations. *J. Evol. Biol.* 23(8):1581–96
- Zaman L, Meyer JR, Devangam S, Bryson DM, Lenski RE, Ofria C. 2014. Coevolution drives the emergence of complex traits and promotes evolvability. *PLOS Biol.* 12(12):e1002023