

The Importance of Demographic Niches to Tree Diversity

Richard Condit,^{1,2*} Peter Ashton,³ Sarayudh Bunyavejchewin,⁴ H. S. Dattaraja,⁵ Stuart Davies,³ Shameema Esufali,⁶ Corneille Ewango,⁷ Robin Foster,⁸ I. A. U. N. Gunatilleke,⁶ C. V. S. Gunatilleke,⁶ Pamela Hall,⁹ Kyle E. Harms,¹⁰ Terese Hart,¹¹ Consuelo Hernandez,¹² Stephen Hubbell,¹³ Akira Itoh,¹⁴ Somboon Kiratiprayoon,¹⁵ James LaFrankie,¹⁶ Suzanne Loo de Lao,² Jean-Remy Makana,¹¹ Md. Nur Supardi Noor,¹⁷ Abdul Rahman Kassim,¹⁷ Sabrina Russo,³ Raman Sukumar,⁵ Cristián Samper,¹⁸ Hebbalalu S. Suresh,⁵ Sylvester Tan,¹⁹ Sean Thomas,²⁰ Renato Valencia,¹² Martha Vallejo,²¹ Gorky Villa,¹² Tommaso Zillio^{1,2}

Most ecological hypotheses about species coexistence hinge on species differences, but quantifying trait differences across species in diverse communities is often unfeasible. We examined the variation of demographic traits using a global tropical forest data set covering 4500 species in 10 large-scale tree inventories. With a hierarchical Bayesian approach, we quantified the distribution of mortality and growth rates of all tree species at each site. This allowed us to test the prediction that demographic differences facilitate species richness, as suggested by the theory that a tradeoff between high growth and high survival allows species to coexist. Contrary to the prediction, the most diverse forests had the least demographic variation. Although demographic differences may foster coexistence, they do not explain any of the 16-fold variation in tree species richness observed across the tropics.

Comparative studies of tree demography typically consider the entire community as a unit, ignoring species differences (1), simply because most tree inventories include small samples of many species (2, 3). Comparative studies show that tropical forests typically have higher turnover than do temperate forests (4) and that higher tree turnover associates with higher tree diversity (5). These studies cannot, however, test ecological hypotheses about diversity, coexistence, and demography (6–10).

A tradeoff between rapid growth and long life span permits species coexistence and can foster diversity: Species reproducing early in life persist despite poor competitive ability by growing rapidly on disturbed sites where resources are abundant. Long-lived species coexist by outliving the weedy invaders, persisting where resources are scarce. This is a familiar and widely known tradeoff in plant and animal communities (9–11) called the successional-ecological hypothesis (7, 12). At a deterministic equilibrium, an indefinite number of species can coexist by this mechanism, each differing from all others along a continuum from short life span (with high growth) to long life span (and low growth). With stochastic demography, however, there is limiting similarity and the equilibrium species richness is finite (11, 13). This hypothesis is widely quoted as an explanation for tropical forest diversity (14–16). Here, we ask whether species differences along a demographic axis explain why some tropical forests have many more species than others.

If demographic niches are a key force controlling forest diversity, then more diverse forests have more demographic niches. More niches could come about either by spreading

demographic rates over a wider range or packing more in the same range. Here, we focus on the first prediction: Tropical forests gain diversity by having a wider range of demographic niches, as reflected by the range of mortality and growth rates across species.

We provide a direct test by quantifying mortality and growth of 4500 tree species in 10 different forests in America, Asia, and Africa (17). The 10 sites form a large-scale observation program, spanning a wide range of environmental conditions, designed to provide species-specific information for little-known tropical trees (18). At each site, a 20- to 52-ha tree census was set out in extensive, largely undisturbed forest (table S1). Species richness within the census plots differed by 16-fold, from 73 species per 50 ha in a dry forest at Mudumalai, India, to 1167 species per 50 ha in a wet dipterocarp forest in Sarawak, Malaysia (19).

Past studies on the demography of individual tree species were based on direct measures of rate constants. These excluded many rare species because their rate estimates are subject to high error (20, 21). To overcome this limitation, we did not simply record species' rates of mortality and growth; instead, we quantified the distribution of demographic rates across the entire community. A hierarchical Bayesian approach accomplishes this with explicit probability models covering both the observations of individual trees within species and the variation among species; all species, including rare ones, are included. For mortality, within-species distributions were modeled with the binomial distribution; for growth, we chose the log-normal based on the tendency for individual growth rates within a species to be

highly skewed to the right. By separating within-species variation, the hierarchical model allows focus on the question of how species differ (10, 22, 23).

At the community level, we had to describe the variation in species' demographic rates across species, and again, we used the log-normal to account for the skew to the right. Histograms of mortality rate m and growth rate g (24) are fitted well by the log-normal when rare species are excluded (Fig. 1). The log-normal requires two parameters, μ and σ , the mean and standard deviation of the natural logarithm of m or g , respectively. We were able to estimate values of μ and σ that best describe a community's demography with the use of the Gibbs sampler, simultaneously producing for every species estimates of mortality and growth rates that are adjusted for abundance. That is, for abundant species, the estimate is barely different from the observed rate, but for rare species, it is guided by the community-wide pattern (25).

Fitted log-normal distributions for the Lambir forest in Malaysia are plotted with observed histograms of sapling mortality and growth (Fig. 1). The fit is close for more common species (filled bars), demonstrating that the large number of zeroes in the mortality

¹National Center for Ecological Analysis and Synthesis, 735 State Street, Santa Barbara, CA 93101, USA. ²Center for Tropical Forest Science, Smithsonian Tropical Research Institute, Unit 0948, APO AA 34002–0948, USA. ³Center for Tropical Forest Science, Asia Program, Arnold Arboretum Asia Program, Harvard University Herbaria, 22 Divinity Avenue, Cambridge, MA 02138, USA. ⁴Thai National Park Wildlife and Plant Conservation Department, Research Office, 61 Phaholyothin Road, Chatuchak, Bangkok 10900, Thailand. ⁵Center for Ecological Sciences, Indian Institute of Science, Bangalore 560012, India. ⁶Department of Botany, Faculty of Science, University of Peradeniya, Peradeniya 20400, Sri Lanka. ⁷University of Missouri, 8001 Natural Bridge Road, St. Louis, MO 63121–4499, USA. ⁸Botany Department, The Field Museum, Roosevelt Road at Lake Shore Drive, Chicago, IL 60605–2496, USA. ⁹Department of Biology, Florida State University, 5051 Quail Valley Road, Tallahassee, FL 32309, USA. ¹⁰Department of Biological Sciences, Louisiana State University, Baton Rouge, LA 70803, USA. ¹¹Wildlife Conservation Society, 1725 Avenue Monjiba, Chanic Building, 2nd floor, Ngalinema, Boite Postale 240, Kinshasa I, Democratic Republic of Congo. ¹²Pontificia Universidad Católica de Ecuador, Post Office Box 17012184, Quito, Ecuador. ¹³Department of Plant Sciences, University of Georgia, 2502 Miller Plant Sciences Building, Athens, GA 30602, USA. ¹⁴Plant Ecology Lab, Faculty of Science, Osaka City University, Sugimoto 3-3-138, Sumiyoshi-ku, Osaka 558, Japan. ¹⁵Faculty of Science and Technology, Thammasat University (Rangsit), Klongluang, Patumtani 12121, Thailand. ¹⁶Center for Tropical Forest Science, Asia Program, National Institute of Education, Nanyang Technological University, 1 Nanyang Walk, 637616 Singapore. ¹⁷Forest Environment Division, Forest Research Institute Malaysia, Kepong, Kuala Lumpur 52109, Malaysia. ¹⁸National Museum of Natural History, Smithsonian Institution, 10th Street and Constitution Avenue, NW, Washington, DC 20560, USA. ¹⁹Forest Research Center, Sarawak Forest Department, Km10 Jalun Datak Amar Kalong Ningkan, 93250 Kuching, Sarawak, Malaysia. ²⁰University of Toronto, 33 Willcock Street, Toronto, Ontario M5S 3B3, Canada. ²¹Instituto Alexander von Humboldt, Carrera 13 #28-01, Piso 7, Edificio Palma Real, Bogotá, Colombia.

*To whom correspondence should be addressed: condit@ctfs.si.edu

histogram are sampling artifacts in rare species. Growth rates are also spread by rare species, though not as conspicuously. Fitted distributions for all the forests can be compared graphically (Fig. 2) or with estimated 2.5 and 97.5 percentiles (Table 1). In the supporting online material, tables of mortality and growth rates for all species are provided (database S1).

Most of the forests were dominated by species with sapling mortality rates near 1% year⁻¹ (Fig. 2). Even the high-mortality forests, such as Barro Colorado Island (BCI) and Hua Khae Khaeng National Park (HKK), had modes close to 1% year⁻¹ and low rates around 0.4% year⁻¹. The main feature separating these high-rate forests from the low-rate sites (such as Pasoh in Malaysia) is the long tail reaching 20% year⁻¹ mortality; at Pasoh, nearly all species had mortality rates of <3% year⁻¹. Thus, forests

fell broadly in two groups: BCI, HKK, and La Planada had upper sapling mortality rates above 20%, whereas Sinharaja, Lambir, Pasoh, and Yasuni had upper rates below 8%. The Congo sites had exceedingly low mortality stretching to a modest 10% at the upper end.

Distributions of growth rates were similar to distributions of mortality, but growth was about half as variable across species (Fig. 2). Conspicuously, sites with less variation in mortality also had less variation in growth (Fig. 3). These patterns held for larger trees, although mortality and growth rates were lower (table S2 and fig. S1).

Examples from individual species help illustrate. At BCI, a fast-growing understory treelet, *Palicourea guianensis*, had a population of 376 saplings in 1982, and every single one had died by 2005 (nevertheless, the population

grew to 851). Although *Palicourea*'s mortality rate is infinite by direct calculation, the Gibbs sampler produces an estimate of 33% year⁻¹. *Alloplectus schultzei*, a small, weedy treelet at La Planada, also suffered 33% year⁻¹ mortality, losing 284 of 335 individuals over 6 years. In contrast, of 1161 species at Lambir, none had mortality of 30% year⁻¹, and only two had rates above 20%; at Pasoh between 1987 and 2000, the very highest Gibbs-corrected mortality rate among 802 species was 14% year⁻¹, in *Melastoma malabathricum*.

At the other end of the distribution, *Cupania sylvatica*, a mid-sized tree of the BCI understory, lost only 10 of 1102 individuals between 1990 and 1995 (0.23% year⁻¹), and *Carapa guianensis* at La Planada, a large and valuable timber tree, lost only 11 of 894 (0.28% year⁻¹). In three census intervals at Pasoh, the lowest

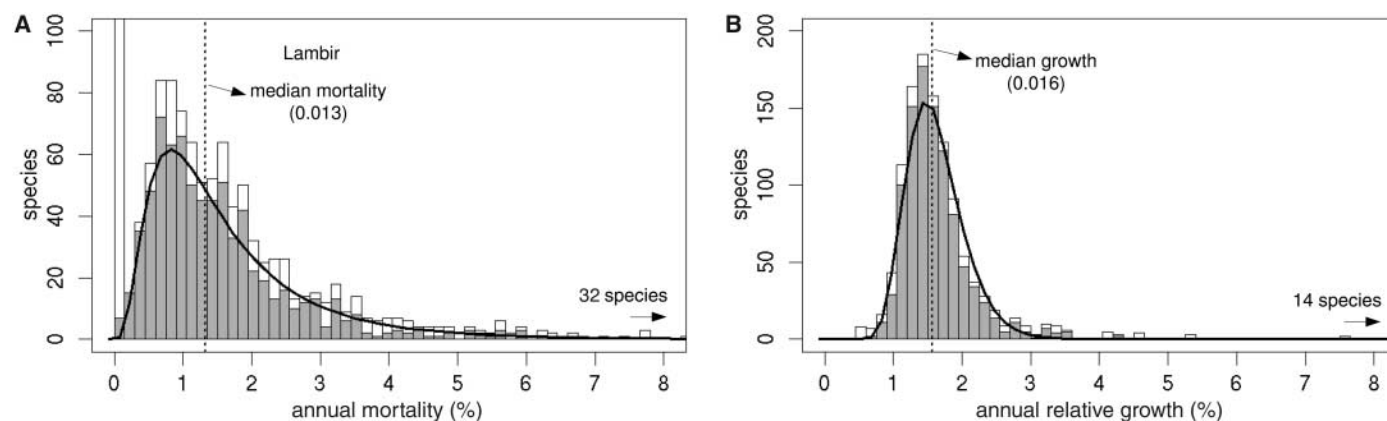


Fig. 1. Distribution of sapling demographic rates of all species in the Lambir plot. **(A)** Annual mortality, m , for all individuals with $dbh = 10$ to 99 mm. Filled bars show the histogram of observed mortality rates for the 746 species with ≥ 50 individuals; open bars add the 415 species with < 50 individuals. The open bar at $m = 0$ extends off the graph (162 species had no mortality; 6 of these had < 50 individuals). The horizontal axis is m , expressed as a percent. The solid line is the fitted log-normal, based on all 1161 species. The dashed vertical line indicates the mean of

the logarithm of the fitted distribution (parameter μ , Table 1), which is very close to the median. **(B)** Annual growth, g , for individuals 10 to 49 mm in diameter. Filled bars are the histogram for 995 species with ≥ 10 individuals; open bars for the remaining 154 species. The solid curve and dashed line are the fitted log-normal and the mean of the logarithm, respectively, as in (A). Both histograms are curtailed at 8% to accentuate details where most of the species fall. The number of species above 8% is indicated by arrows.

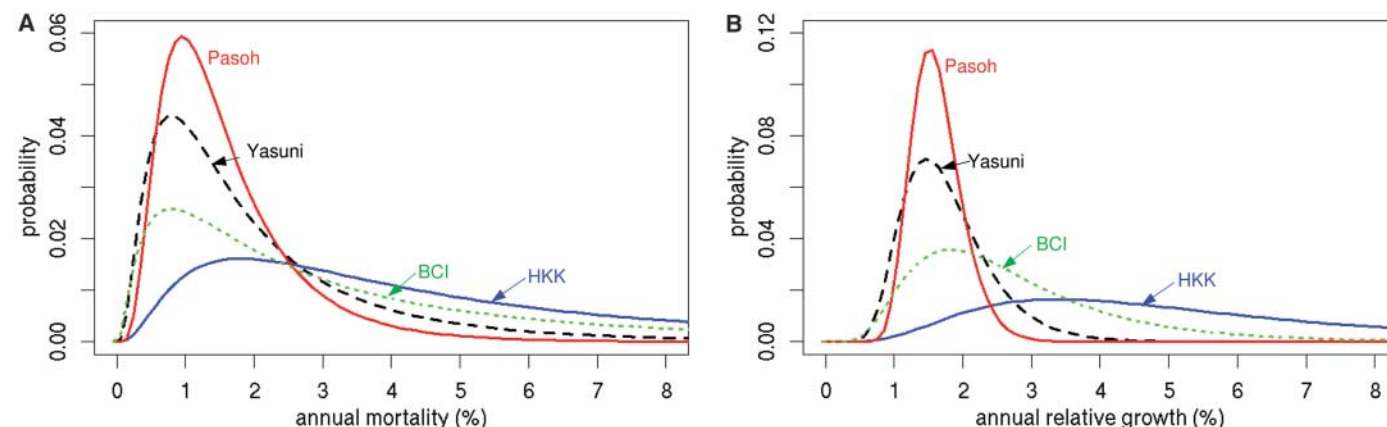


Fig. 2. Comparing the fitted distributions of sapling demography in four forests. **(A)** Annual mortality rate, m . **(B)** Annual growth rate, g . The lower end of the growth distribution in saplings is limited by measurement accuracy (30).

mortality rate recorded was 0.34% year⁻¹, in *Cynometra malaccensis*.

The Mudumalai forest stood out. Saplings had greatly elevated mortality and growth, with rates stretching much higher than any other site. Between 1988 and 1992, every species at Mudumalai had sapling growth of >6% year⁻¹, and only BCI and HKK had many rates this high. At Lenda and Sinharaja, no species grew by 6% year⁻¹. For larger trees (≥100 mm diameter), however, Mudumalai was in line with other forests, having modes of mortality and growth near 1% year⁻¹ (table S2). Indeed, trees at Mudumalai had among the lowest rates as well as the highest: *Anogeissus latifolia* had

116 deaths out of 2179 trees from 1988 to 2000, whereas *Kydia calycina* had 1272 of 1328 trees die over the same interval, many because of elephant herbivory (26). Their rates differ by 50-fold: 27% year⁻¹ in *Kydia* compared with 0.47% year⁻¹ in *Anogeissus*.

Three of the sites with long tails of elevated mortality and growth—BCI, HKK, and Mudumalai—have intense annual dry seasons (table S1). Mudumalai and HKK also burn in some years (26) (other plots do not burn and none suffer large-scale wind damage). It was not surprising that annual drought elevated mortality. Many species at these sites, however, had exceedingly low rates of mortality and evident-

ly did not suffer much from drought. Conversely, forests lacking the tail of high growth and mortality had no or modest annual dry seasons, including the three forests dominated by Dipterocarpaceae (Sinharaja, Pasoh, and Lambir). Seasonality, however, was not the only factor predicting high variation in demography; the ever-wet cloud forest at La Planada, Colombia, had a wide spread of growth and mortality, comparable to the seasonally dry sites.

Mudumalai and HKK have relatively open canopies compared with all of the other sites, a typical feature of dry forests, and many saplings at Mudumalai are sprouts from large root systems. These are likely reasons for elevated sapling growth at the two sites. In contrast, both BCI and La Planada have dense canopies and dark understories, so canopy openness does not obviously account for the high-growth species at those two sites.

Contrary to the prediction that demographic variability begets species richness, diverse forests had the least variation in demography (Fig. 3 and fig. S1). If anything, the most diverse forests had the fewest demographic niches. At Lambir, high species richness coupled with a low diversity of demographic rates meant that 135 tree species coexisting in close proximity had sapling mortality rates in a narrow window from 0.8 to 1.0% year⁻¹.

We do not question that demographic variability plays some role in species coexistence. In American forests, the familiar genus *Cecropia* is found exclusively in small forest clearings (or outside the forest), where it rapidly

Fig. 3. Range of sapling demographic rates for tree species within a community versus the number of species at the site. The range is the logarithm of the ratio between the 97.5 and 2.5 percentiles of the fitted distributions (Table 1). The range for mortality is given by filled circles; the range for growth is given by open triangles. Sites can be identified by the number of species; for example, Lambir is the most diverse and farthest to the right. Multiple censuses at BCI, Pasoh, and Mudumalai are included, and in each case, fall in a tight group.

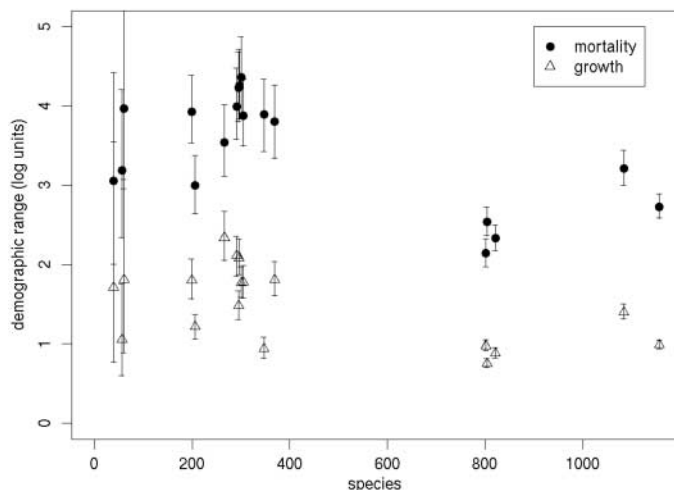


Table 1. Variation in sapling mortality and growth rates across species in tropical forests. For mortality, all individuals with *dbh* = 10 to 99 mm were included; for growth, those with *dbh* = 10 to 49 mm were included. Species number refers to those with at least one 10- to 99-mm sapling alive at the outset of a given census interval. Under mortality are percentiles of the distribution of mortality rate parameters (*m*) across

species: the median plus lower and upper percentiles (2.5 and 97.5) of the fitted log-normal. Similarly, under growth are percentiles for the distribution of growth rates (*g*) across species. Rates are expressed as percentages (100*m* or 100*g*)—that is, 5 = 5% = 0.05. For each of the percentiles, confidence limits are given, based on the Gibbs sampler (25). Information about the sites is presented in table S1.

Site	Years	No. of species	Annual mortality (%)			Relative growth (%)		
			Median	Lower	Upper	Median	Lower	Upper
BCI	82–85	284	3.14 ± 0.46	0.38 ± 0.10	26.0 ± 6.9	2.84 ± 0.16	1.35 ± 0.14	6.0 ± 0.7
BCI	85–90	282	2.56 ± 0.37	0.31 ± 0.08	21.5 ± 5.5	2.41 ± 0.18	0.85 ± 0.12	6.8 ± 1.0
BCI	90–95	282	2.85 ± 0.43	0.32 ± 0.09	25.3 ± 6.7	2.15 ± 0.13	0.89 ± 0.09	5.2 ± 0.7
BCI	95–00	285	3.35 ± 0.42	0.48 ± 0.11	23.3 ± 5.7	1.97 ± 0.12	0.81 ± 0.09	4.8 ± 0.6
BCI	00–05	285	2.91 ± 0.41	0.40 ± 0.10	21.4 ± 5.5	2.10 ± 0.16	0.73 ± 0.10	6.1 ± 0.9
Yasuni	96–03	1077	1.55 ± 0.10	0.31 ± 0.04	7.7 ± 0.9	1.67 ± 0.04	0.83 ± 0.04	3.4 ± 0.2
La Planada	97–03	218	3.22 ± 0.47	0.45 ± 0.13	22.9 ± 5.9	2.30 ± 0.17	0.93 ± 0.13	5.7 ± 0.8
Pasoh	87–90	802	1.04 ± 0.06	0.36 ± 0.04	3.0 ± 0.3	2.25 ± 0.05	1.38 ± 0.05	3.7 ± 0.2
Pasoh	90–95	801	1.35 ± 0.07	0.42 ± 0.05	4.3 ± 0.4	1.59 ± 0.03	1.02 ± 0.04	2.5 ± 0.1
Pasoh	95–00	804	1.69 ± 0.09	0.47 ± 0.06	6.0 ± 0.6	1.55 ± 0.03	1.07 ± 0.03	2.3 ± 0.1
Lambir	92–97	1161	1.32 ± 0.07	0.34 ± 0.03	5.2 ± 0.4	1.57 ± 0.03	0.96 ± 0.03	2.6 ± 0.1
HKK	93–99	256	4.11 ± 0.57	0.70 ± 0.19	24.1 ± 6.6	4.83 ± 0.45	1.53 ± 0.26	15.2 ± 2.7
Mudumalai	88–92	56	13.06 ± 3.48	2.65 ± 1.42	64.4 ± 36	7.87 ± 1.26	4.65 ± 1.52	13.3 ± 3.8
Mudumalai	92–96	52	17.06 ± 6.43	2.35 ± 1.51	124 ± 113	6.35 ± 1.67	2.57 ± 1.38	15.7 ± 11
Mudumalai	96–00	39	7.96 ± 2.70	1.73 ± 1.13	36.6 ± 25	5.71 ± 1.53	2.42 ± 1.69	13.4 ± 9.8
Sinharaja	95–01	205	1.35 ± 0.17	0.30 ± 0.07	6.0 ± 1.3	1.38 ± 0.07	0.75 ± 0.07	2.5 ± 0.2
Edoro	94–00	368	1.43 ± 0.20	0.21 ± 0.06	9.6 ± 2.5	1.41 ± 0.09	0.57 ± 0.07	3.5 ± 0.5
Lenda	94–00	346	1.26 ± 0.19	0.18 ± 0.06	8.8 ± 2.2	1.06 ± 0.04	0.66 ± 0.05	1.7 ± 0.1

colonizes and rapidly dies. The upper end of sapling mortality and growth distributions in America is set by gap specialists: *C. obtusifolia* at BCI (12% year⁻¹ mortality, 14% year⁻¹ growth), *C. sciadophylla* at Yasuni (5.0% year⁻¹ mortality, 6.3% year⁻¹ growth), and *C. monostachya* at La Planada (8.8% year⁻¹ mortality, 8.2% year⁻¹ growth). Diverse Southeast Asian forests lacked species with such high rates (27).

The most diverse tropical forests are the least diverse demographically. It remains plausible that demographic niches are packed more tightly in some forests than others, but this seems unlikely, because packing should depend only on population size and turnover, which do not vary much. Moreover, the successional-niche hypothesis is not favored by the strong peak in demographic rates near 1 to 2% year⁻¹; if demographic niches were crucial, then rates ought to be spread evenly over the entire range (28). Instead, the similarity in demography of many species suggests trait convergence (29). We believe that broad diversity differences are due to the source pool of different biogeographic regions, and that demographic differences play a minor role in species coexistence.

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- Analyses were supported by NSF grant DEB-9806828 of the Research Coordination Network Program to the Center for Tropical Forest Science. Data collection was funded by many organizations, principally the NSF, Andrew W. Mellon Foundation, Peninsula Community Foundation, Smithsonian Tropical Research Institute, Arnold Arboretum (Harvard), Indian Institute of Science, Forest Research Institute of Malaysia, Royal Thai Forest Department, National Institute of Environmental Studies (Japan), and John D. and Catherine T. MacArthur Foundation. We thank the hundreds of field workers who have measured and mapped trees.

Supporting Online Material

www.sciencemag.org/cgi/content/full/1124712/DC1

Materials and Methods

Fig. S1

Tables S1 to S4

References

Database S1

Computer Codes

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A Single Amino Acid Mutation Contributes to Adaptive Beach Mouse Color Pattern

Hopi E. Hoekstra,^{1*} Rachel J. Hirschmann,¹ Richard A. Bunday,² Paul A. Insel,² Janet P. Crossland³

Natural populations of beach mice exhibit a characteristic color pattern, relative to their mainland conspecifics, driven by natural selection for crypsis. We identified a derived, charge-changing amino acid mutation in the melanocortin-1 receptor (*Mcl1r*) in beach mice, which decreases receptor function. In genetic crosses, allelic variation at *Mcl1r* explains 9.8% to 36.4% of the variation in seven pigmentation traits determining color pattern. The derived *Mcl1r* allele is present in Florida's Gulf Coast beach mice but not in Atlantic coast mice with similar light coloration, suggesting that different molecular mechanisms are responsible for convergent phenotypic evolution. Here, we link a single mutation in the coding region of a pigmentation gene to adaptive quantitative variation in the wild.

The identification of the specific molecular changes underlying adaptive variation in quantitative traits in wild populations is of prime interest (1, 2). Pigmentation phenotypes are particularly amenable to genetic dissection because of their high heritability and

our knowledge of the underlying developmental pathway (3). In a series of classic natural history studies (4, 5), Sumner documented pigment variation in *Peromyscus polionotus*, including eight extremely light-colored "beach mouse" subspecies, which inhabit the primary dunes

and barrier islands of Florida's Gulf and Atlantic coasts (6). This light color pattern is driven by selection for camouflage (7, 8) as major predators of *P. polionotus* include visual hunters (9). Because the barrier islands on the Gulf Coast are <6000 years old (10), this adaptive color variation may have evolved rapidly.

We examined the contribution of the melanocortin-1 receptor gene (*Mcl1r*) to this adaptive color patterning. MC1R, a G protein-coupled receptor, plays a key role in melanogenesis by switching between the production of dark eumelanin and light pheomelanin (11). Mutations in *Mcl1r* have been statistically associated with Mendelian color polymorphisms in several mammalian species (e.g., 12–14) and in natural variants of avian plumage (15, 16).

¹Division of Biological Sciences and ²Department of Pharmacology, University of California, San Diego, La Jolla, CA 92093, USA. ³*Peromyscus* Genetic Stock Center, Department of Biological Sciences, University of South Carolina, Columbia, SC 29208, USA.

*To whom correspondence should be addressed. E-mail: hoekstra@ucsd.edu