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 Plana (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 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.

### References and Notes

2. Most forest censuses cover a single hectare or less and include only larger trees. Except in Europe and North America, such small samples capture less than half the local species, with many represented by a single individual.
17. Table 3 includes 4956 species, but some occur at more than one site, and taxonomy has not been fully aligned across sites. We estimate 4500 different species at the 10 sites.
19. The Sarawak plot covers 52 ha. For comparison, the number 1167 was taken from a 50-ha subset.
24. The rate constant is the derivative of population size *N* with respect to time, *t*, due to mortality alone. It is approximated by *dhb/dbh* Relative growth rate is *g* = *dhb/dbh* where *dhb* is the stem diameter at breast height.
25. Materials and methods are available as supporting material on Science Online.

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

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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 (*Mc1r*) in beach mice, which decreases receptor function. In genetic crosses, allelic variation at *Mc1r* explains 9.8% to 36.4% of the variation in seven pigmentation traits determining color pattern. The derived *Mc1r* 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 (*Mc1r*) 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 *Mc1r* 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).

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We sequenced the entire coding region of Mc1r (954 base pairs (bp)) in five Santa Rosa Island beach mice (P. p. leucopus) and five mainland mice (P. p. subgriseus) from colonies derived from wild populations (17). A single, fixed nucleotide polymorphism (SNP) occurs between the two subspecies, resulting in a charge-changing amino acid variant (R56C) in the first intracellular loop of MC1R (fig. S1). To determine the ancestry of this mutation, we genotyped this SNP in 14 other Peromyscus species (N = 45 individuals) (table S1). These additional species are all fixed for R56 as in the mainland “dark allele,” suggesting that the “light allele” is derived and not present in any other fully pigmented Peromyscus species.

To examine independently the relationship between this mutation (R56C) and the derived light-colored phenotype, we generated a large reciprocal F1 intercross between the Santa Rosa Island beach mouse and the mainland subspecies (Fig. 1). We characterized color phenotype for seven pigmentation traits, which provide an overall description of the continuous variation in color pattern, and genotyped the Mc1r allele in 459 F2 individuals [126 with two dark Mc1r alleles (RR), 215 with one dark and one light allele (RC), and 118 with two light alleles (CC)]. Based on the observed phenotypic variation among F2s, beach mouse color pattern has a multigenic architecture (Table 1). Pairwise correlation (r2) between traits ranged from 0.147 to 0.500 (P < 0.05 for all comparisons), revealing a shared genetic basis among traits but also indicating the role of loci expressed in distinct spatial regions.

Two lines of evidence suggest that both dominant and recessive alleles contribute to the adaptive light color phenotype. First, all F1 hybrids are intermediate in overall color pattern, with some traits resembling the mainland parent and other traits the beach mouse parent (Table 1). Second, dominance varies at the Mc1r locus itself. Phenotypic scores for pigmentation traits among F2s are sometimes above the mean score of 1, suggesting that the light Mc1r allele is partially recessive, and sometimes below the mean score, suggesting that the light Mc1r allele is partially dominant (Table 1).

We found a significant statistical association between the R56C polymorphism and all seven traits, although the percentage of phenotypic variance explained by Mc1r varied (9.8% to 36.4%) (Table 1). For some pigmentation traits, the association between phenotype and Mc1r genotype was notable: All RR F2s (N = 126) expressed the darkest phenotype on the rostrum and, conversely, only CC individuals expressed the lightest rostrum phenotype (N = 49). We calculated principal component scores for the combined color traits and evaluated whether there were

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**Table 1.** Statistical association between allelic variation at Mc1r and seven pigmentation traits in 459 F2 individuals. Phenotypes were categorically scored based on the pigmentation pattern on individual hairs in seven areas (traits). For all seven traits, each F2 was scored as follows: 0, unpigmented hair; 1, partially pigmented hair; and 2, fully pigmented hair. The total number of F2s with each phenotypic score is partitioned by Mc1r genotype for each trait: RR, homozygous for the dark Mc1r allele; RC, heterozygous; and CC, homozygous for the light Mc1r allele. The frequency of F2s with each phenotype score is shown for each trait, with the phenotype of all F2 individuals in bold. The mean phenotypic score (±SD) for each trait is provided and is also partitioned among each genotype. The percentage of variance explained by Mc1r genotype (PVE) and likelihood ratio chi-square (X2) values are shown. *All seven tests were statistically significant (P < 0.0001).
significant differences between the Mc1r alleles for PC1 scores (PC1 explained 71% of the variance in color traits); these analyses suggest that Mc1r accounts for 26% of PC1 ($P < 0.0001$). Together, these data indicate that Mc1r is a major effect locus on color pattern, having pleiotropic effects on pigmentation in spatially diverse areas of the body.

Functional tests of the light and dark Mc1r alleles demonstrate that the R65C amino acid mutation contributes to variation in adaptive pigmentation. We performed in vitro assays on HEK293T cells expressing light or dark Peromyscus coding region alleles at similar levels. Stimulation with an MC1R agonist (NDP-αMSH) and subsequent measurement of generated cyclic adenosine monophosphate (cAMP) by radioimmunoassay revealed that compared with dark receptor–expressing cells, cells that expressed the light MC1R had a statistically significant reduction in basal and stimulated cAMP formation (Fig. 2A) ($P < 0.05$, Student’s t test, $n = 4$; maximum responses: dark allele = 120 ± 15 pmol/mg protein, light allele = 28 ± 5 pmol/mg protein). MC1R-mediated cAMP formation is associated with melanin production (18); thus, reduced cAMP production may underlie the lack of dark pigmentation observed in mice expressing the light MC1R.

The light MC1R displays lower affinity binding to αMSH, and the decreased response to guanine nucleotide by the light receptor implies an altered ability to interact with G proteins (Fig. 2B). Specifically, whereas the dark receptor displays a substantial GppNHp-promoted shift in IC$_{50}$ (from 1.37 ± 0.01 nM in the absence to 15.4 ± 0.4 nM in the presence of GppNHp), the light allele has a decreased affinity for nucleotide and a nonsignificant IC$_{50}$ shift (without GppNHp, 222 ± 30 nM; with, 178 ± 14 nM). These results demonstrate that a single amino acid mutation in MC1R is responsible for reduced ligand binding and G protein coupling, consistent with the reduced ability of the light MC1Rs to promote cAMP formation in vivo.

**Fig. 2.** Cyclic AMP and ligand-binding assays showing functional differences between light and dark Mc1r alleles. All data points are means ± SEM. (A) HEK293T cells transfected with dark MC1R (filled circles) or light MC1R (open circles) have different cAMP responses to NDP-αMSH stimulation. Bars indicate basal level of cAMP production in cells transfected with dark MC1R (black) and light MC1R (white); lines indicate standard error. (B) αMSH competition for $[^{125}]$NDP-αMSH binding to HEK293T cells transfected with dark MC1R (filled symbols) or light MC1R (open symbols). Squares indicate assays without GppNHp and circles with GppNHp (100 μM).

**Fig. 3.** Frequency of Mc1r alleles in mainland and eight beach mouse subspecies from northern Florida. Images represent typical color patterns for each subspecies. Red areas represent the distribution of each beach mouse subspecies, and the gray area represents the range of mainland subspecies in Florida. Lines indicate sampling locales. Circles represent frequencies of the light Mc1r (white) and dark Mc1r (black) alleles identified. The number of alleles sampled is provided.
We next examined the frequency of McI\(r\) alleles in natural populations by genotyping the informative SNP in eight beach mouse subspecies and one mainland subspecies (Fig. 3 and table S2). In the Gulf Coast the light allele was at highest frequency (0.95) in the palest beach mouse subspecies, absent in the darkest subspecies, and at intermediate frequencies in the other subspecies (0.05 to 0.85) (Fig. 3). The distribution of McI\(r\) allele frequencies among Gulf Coast beach mice is not correlated with geographic distance. The light McI\(r\) allele was not detected in the mainland subspecies (0 of 40 alleles). In natural populations, different combinations of functionally distinct McI\(r\) alleles may contribute to variation in color patterning between mainland and Gulf Coast beach mouse subspecies, among the Gulf Coast subspecies, and within subspecies.

The derived McI\(r\) allele was absent from the similarly light-colored Atlantic coast beach mouse populations. We surveyed McI\(r\) allelic variation in two extant subspecies of beach mice and the extinct pallid beach mouse subspecies (~300 miles east of the nearest Gulf Coast beach mouse population) (Fig. 3). The absence of the derived McI\(r\) SNP from all three Atlantic coast subspecies suggests that the same mutation does not contribute to convergent phenotypic adaptation and that light coloration has evolved independently on the Atlantic coast.

This work has specific implications for understanding the evolutionary mechanisms responsible for adaptive phenotypic change. First, the identification and functional characterization of a single amino acid mutation’s effect on quantitative variation provides a convincing exception to a growing number of examples demonstrating that variation in morphology is governed by changes in gene regulatory regions (19, 20). Second, the observation that different combinations of alleles can produce similar pigmentation patterns suggests that distinct molecular mechanisms can underlie adaptive convergence even in similar selective environments (but see 21). Finally, McI\(r\) represents a large effect locus, containing a quantitative trait nucleotide (QTN) contributing to variation in fitness, consistent with the view that adaptation may often proceed by large steps.

References and Notes
17. Materials and methods are available as supporting material on Science Online.
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Supporting Online Material
www.sciencemag.org/cgi/content/full/313/5783/101/DC1
Materials and Methods
Fig. S1
Tables S1 and S2
References
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Decay of Endoplasmic Reticulum-Localized mRNAs During the Unfolded Protein Response

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The unfolded protein response (UPR) allows the endoplasmic reticulum (ER) to recover from the accumulation of misfolded proteins, in part by increasing its folding capacity. Inositol-requiring enzyme-1 (IRE1) promotes this remodeling by detecting misfolded ER proteins and activating a transcription factor, X-box-binding protein 1, through endonucleolytic cleavage of its messenger RNA (mRNA). Here, we report that IRE1 independently mediates the rapid degradation of a specific subset of mRNAs, based both on their localization to the ER membrane and on the amino acid sequence they encode. This response is well suited to complement other UPR mechanisms because it could selectively halt production of proteins that challenge the ER and clear the translocation and folding machinery for the subsequent remodeling process.

The ER is responsible for the structural maturation of proteins entering the secretory pathway (1). When the folding burden on the ER exceeds its capacity, a collection of transcriptional and translational mechanisms termed the UPR is initiated (2). The UPR is essential for survival during ER stress and for development in metazoans (3, 4), especially for the differentiation of secretory cells (5–7). A key sensor of the folding status of the ER is IRE1, a conserved transmembrane protein with an ER luminal sensor domain and cytosolic kinase and ribonuclease domains (8, 9). Accumulation of misfolded proteins in the ER leads to the autophosphorylation and activation of IRE1, which cleaves specific sites in the mRNA encoding the transcription factor X-box-binding protein 1 (XBP-1) (10, 11). This cleavage initiates an unconventional splicing reaction, leading to production of an active transcription factor. Although the sole output of IRE1 activation in yeast appears to be the splicing of the XBP-1 homolog HAC1 (12, 13), several studies in metazoans suggest that IRE1 has additional functions not mediated by XBP-1 (14–16). Overexpression of IRE1 can also promote cleavage of the 28S ribosomal RNA (rRNA) (17) and the mRNA encoding IRE1 itself (18), which raises the possibility that, under some circumstances, the IRE1 nuclease can act on a broader range of substrates.

To systematically test the XBP-1 dependence of IRE1 outputs, we compared the changes in expression profiles associated with the UPR in Drosophila S2 cells depleted of either IRE1 or XBP-1 by RNA interference (RNAi) (Fig. 1, A and B). We induced the UPR with the reducing agent dithiothreitol (DTT), which prevents disulfide-linked folding, and analyzed the changes in expression of ~5000 genes using spotted DNA microarrays.