

## EVOLUTION

# Agouti-related peptide 2 facilitates convergent evolution of stripe patterns across cichlid fish radiations

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The color patterns of African cichlid fishes provide notable examples of phenotypic convergence. Across the more than 1200 East African rift lake species, melanic horizontal stripes have evolved numerous times. We discovered that regulatory changes of the gene *agouti-related peptide 2* (*agrp2*) act as molecular switches controlling this evolutionarily labile phenotype. Reduced *agrp2* expression is convergently associated with the presence of stripe patterns across species flocks. However, cis-regulatory mutations are not predictive of stripes across radiations, suggesting independent regulatory mechanisms. Genetic mapping confirms the link between the *agrp2* locus and stripe patterns. The crucial role of *agrp2* is further supported by a CRISPR-Cas9 knockout that reconstitutes stripes in a nonstriped cichlid. Thus, we unveil how a single gene affects the convergent evolution of a complex color pattern.

Stephen Jay Gould famously posited that if it were possible to rerun the “tape of life,” outcomes would be different (1). The relative importance of determinism and contingency during evolution is still far from settled (2, 3). But for particular groups of organisms, one can now test Gould’s hypothesis. For instance, in less than 8 million to 12 million years, more than 1200 species of cichlid fishes have evolved to form repeated adaptive radiations in the East African Rift Valley lakes, such as Lakes Victoria, Tanganyika, and Malawi (Fig. 1B) (4–8). These adaptive radiations have given rise to a large diversity of species displaying various color patterns (Fig. 1, C to N), including the repeated occurrence of melanic horizontal stripes (Fig. 1A and supplementary text). Convergent evolution is prevalent in the East African cichlid radiations (9–11), providing a replicated natural experiment whereby distantly related species from independent adaptive radiations can be used to determine what mechanisms have generated these recurrent phenotypes (12–16). More specifically, we address whether horizontal stripes, a convergent phenotype, have an identical, similar, or different molecular bases across the independent adaptive radiations of cichlid fishes.

Previously (17)—using a genetic mapping panel of two Lake Victoria species, *Pundamilia nyererei*

(*Pnye*, nonstriped, Fig. 1E) and *Haplochromis sawagei* (*Hsau*, striped, Fig. 1C)—we found that horizontal stripes (Fig. 2C) are inherited as a recessive Mendelian trait mapping to chromosome 18 (Fig. 2A). This was confirmed by a second cross involving the same nonstriped species and another striped species, *H. chilotes* (*Hchi*, striped) (Fig. 2A and supplementary text). To more precisely isolate the causal genetic interval for stripe presence, we fine-mapped the trait using recombinant F<sub>2</sub> individuals of the *Pnye* × *Hsau* cross and reduced the causal interval from 600 to 25 kb (fig. S1). This interval contained the genes *agouti-related peptide 2* (*agrp2*), *v-type proton ATPase subunit d 2* (*atp6Vod2*), and an unknown gene (*unk*) (Fig. 2B). The resequencing of all coding regions revealed no fixed missense or nonsense mutations (fig. S2), suggesting that cis-regulatory variation determines stripe presence.

The teleost-specific *agrp2* (fig. S3) is a strong candidate gene for stripes because its paralogs have been previously associated with pigmentation phenotypes (18–20). To test for *agrp2* expression differences between nonstriped (*Pnye*) and striped (*Hsau*) Lake Victoria cichlids, we performed quantitative polymerase chain reaction (qPCR; Fig. 2D and fig. S5) on a number of adult tissues, including skin (supplementary text). Here, *agrp2* showed a significantly higher expression in the skin of *Pnye* (Fig. 2D and fig. S4). The lack of consistent expression variation between melanic and nonmelanic regions and generally across dorsoventral and anterior-posterior positions suggests that *agrp2* does not shape pigmentation patterns through local expression-level variation but rather acts as a general stripe pattern inhibitor (fig. S6). Whereas qPCR revealed no such expression differences for paralogs and neighboring genes (supplementary text),

qPCRs on F<sub>2</sub> *Pnye/Hsau* hybrid individuals confirmed that expression differences are linked to the *agrp2* locus and exhibit an allelic dosage effect as expected for cis-regulatory mutations (fig. S4).

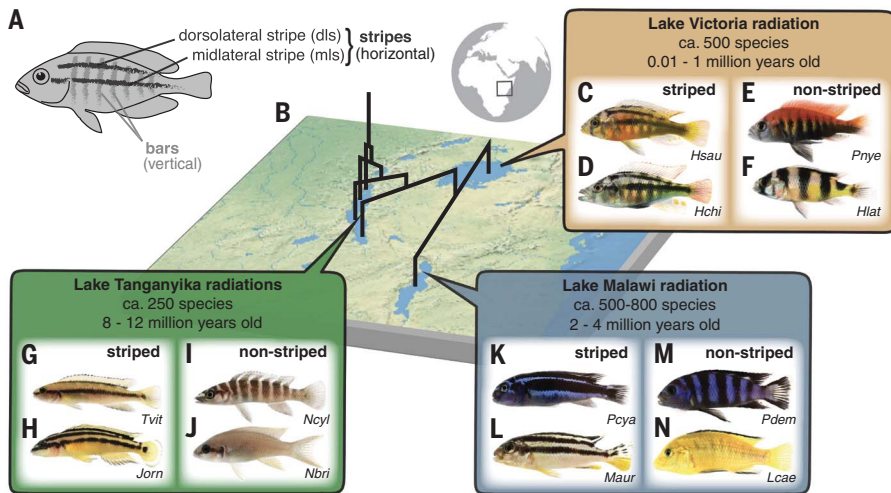
To identify causal mutations affecting both *agrp2* expression and stripe phenotype, we sequenced the *agrp2* locus in individuals ( $n \geq 10$  individuals per population) from natural populations of the three hybrid-cross species. We screened for alternatively fixed, fully associated variants with the stripe phenotype in pairwise comparisons of each striped species (*Hsau* or *Hchi*) versus the nonstriped *Pnye*. Our analyses indicated that a 1.1-kb interval within the first *agrp2* intron (Fig. 2B) that exhibited shared, alternatively fixed alleles is a strong candidate region for a regulatory element controlling *agrp2* expression (fig. S7). To test whether this 1.1-kb interval [*enhancer of agrp2 in Pnye* (*Pnye.enh.a*)] contains cis-regulatory elements that could influence interspecific differences between striped and nonstriped species, we tested the elements of both species in a green fluorescent protein (GFP) reporter assay in vivo (supplementary text). It showed that *Pnye.enh.a* efficiently modulates GFP expression [Tukey’s honest significant difference (HSD),  $P < 0.001$ ] and is significantly more potent than the homologous sequence of the striped species *Hsau* (Tukey’s HSD,  $P < 0.001$ ) (Fig. 2, E and F, and fig. S8). Together, these results indicate that higher expression of *agrp2*, and thereby the suppression of stripe patterns, is indeed enhanced by *Pnye.enh.a* (fig. S9).

Our results reveal *agrp2* as a major determinant of stripe presence that might be sufficient to suppress stripe patterns in *Pnye*. To further test this finding, we used CRISPR-Cas9 genome engineering to manipulate *agrp2* and to thereby potentially derepress stripe patterns. *Pnye* eggs were injected with Cas9 and *agrp2* guide RNAs, and we obtained four mutants, all of which had nonsense and frameshift mutations within *agrp2* (fig. S10 and table S1). These CRISPR-Cas9 mutants developed a continuous midlateral stripe (Fig. 2H and fig. S10) yet no dorsolateral stripe (supplementary text). Because horizontal stripes were never observed in noninjected *Pnye* individuals (>100 observations; Fig. 2G), this strongly suggests that although species such as *Pnye* have no stripes, the genomic and developmental machinery for stripe pattern formation is in place, and stripes can reappear in this nonstriped species by experimental manipulation of *agrp2*.

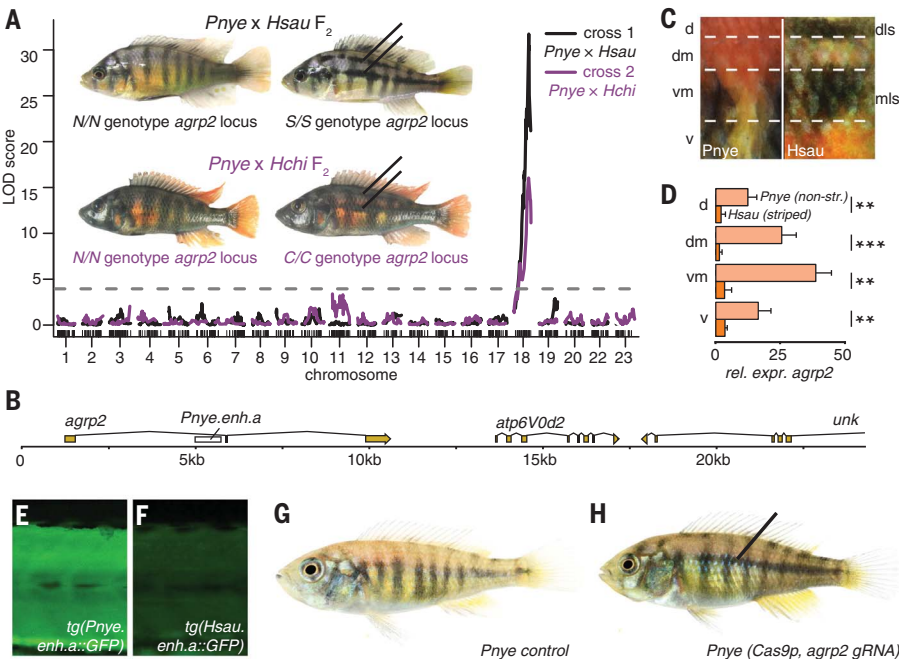
Next, we tested if the expression levels of *agrp2* and stripe patterns are generally associated across other cichlid species from the repeated species flocks of Lakes Victoria, Malawi, and Tanganyika, suggesting a shared molecular basis for convergent stripe phenotypes. Using qPCR on adult skins of striped and nonstriped species of each of the three major East African cichlid radiations (in total, 24 species; fig. S11), we revealed that nonstriped species commonly had higher *agrp2* expression levels than striped species (Fig. 3B). This association was confirmed

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**Fig. 1. Convergent evolution of horizontal stripes across African cichlid radiations.** (A) Schematic of melanic horizontal stripes and vertical bars. (B) Map of the African Great Lakes Victoria, Tanganyika, and Malawi and superimposed simplified phylogenetic tree of the adaptive radiations of Lakes Tanganyika (green; not all paraphyletic lineages shown), Malawi (blue), and Victoria (orange). (C to N) Twelve of the focal striped and nonstriped species of this study, including species from Lakes Victoria [(C) to (F)], Tanganyika [(G) to (J)], and Malawi [(K) to (N)]. *Hlat*, *Haplochromis latifasciatus*; *Jorn*, *Julidochromis ornatus*; *Lcae*, *Labidochromis caeruleus*; *Maur*, *Melanochromis auratus*; *Nbrl*, *Neolamprologus brichardi*; *Ncyl*, *Neolamprologus cylindricus*; *Tvit*, *Telmatochromis vittatus*.



**Fig. 2. *Agrp2* controls stripe loss in Lake Victoria cichlids.** (A) In two mapping crosses between *Pnye* and *Hsau* and *Pnye* and *Hchi*, horizontal stripes map to the same region on chromosome 18. (B)  $F_2$  recombinant fine-mapping isolates a 25-kb interval containing *agrp2*. (C and D) Skin biopsies show differential *agrp2* expressions between skins from *Hsau* (C) and *Pnye* (D). Error bars indicate means + SD. (E and F) An intronic 1.1-kb element of *Pnye* (*Pnye.enh.a*) is regulatory active (E) and shows stronger activity than the homologous sequence of the striped species *Hsau* (F) in a zebrafish larvae GFP reporter assay. (G and H) CRISPR-Cas9 mediated knockouts of *agrp2* in the normally nonstriped cichlid *Pnye* (G) develop stripes [(H); midlateral stripe indicated by line]. LOD score, logarithm of the odds score; *N/N*, homozygous for the *Pnye agrp2* allele; *S/S*, homozygous for the *Hsau agrp2* allele; *C/C*, homozygous for the *Hchi agrp2* allele; *tg*, transgenic construct; *gRNA*, guide RNA; *mls*, midlateral stripe; *dls*, dorsolateral stripe; *d*, dorsal; *dm*, dorsomedial; *vm*, ventromedial; *v*, ventral.

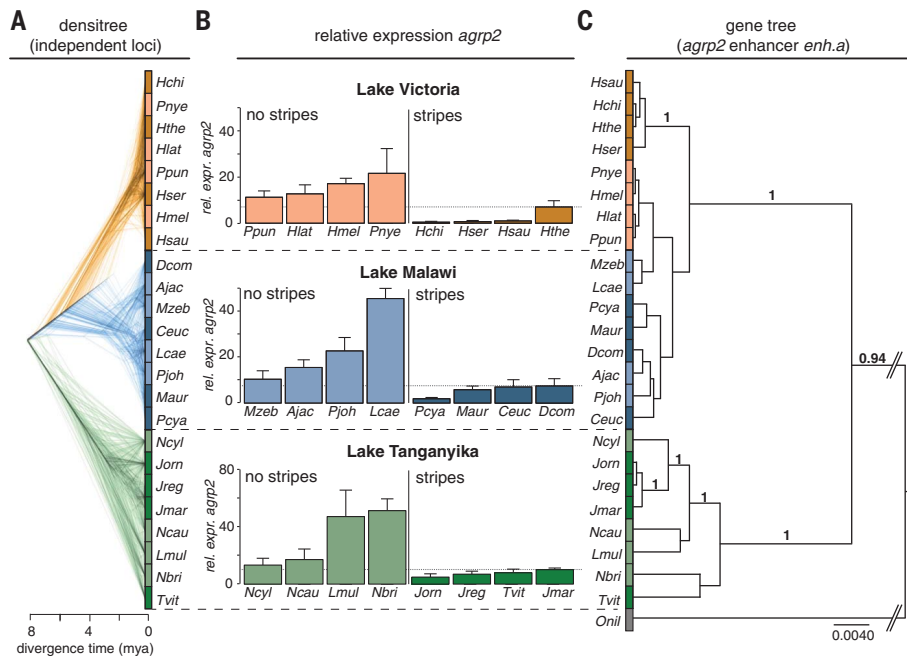
by comparative phylogenetic analyses (Fig. 3A), demonstrating a significant evolutionary association between low *agrp2* expression and stripe presence [phylogenetic analysis of variance (ANOVA); mean  $P < 0.001$ ; supplementary text].

To determine if this convergence at the phenotypic and *agrp2* gene expression level is also paralleled at the sequence level (16), we comparatively analyzed homologous *enh.a* sequences across cichlids from Lakes Victoria, Malawi, and Tanganyika. A tree of *enh.a* revealed substantial sequence variation and resolved striped Lake Victoria species as monophyletic, suggesting a single origin of the striped alleles, whereas striped species of other lakes were not monophyletic (Fig. 3C). None of the nine mutations within *enh.a* that showed complete association with stripes in Lake Victoria cichlids showed similar stripe association in cichlids of Lakes Malawi or Tanganyika (figs. S12 and S13). Consequently, independent mutations must be affecting *agrp2* expression and thereby stripe patterns across the three major cichlid radiations (Fig. 4D).

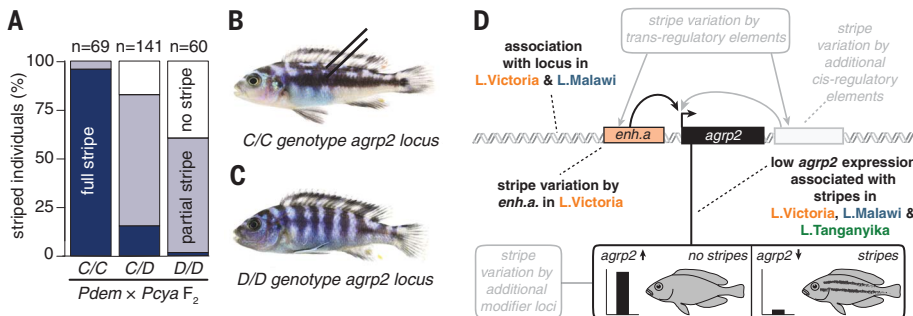
Lastly, we tested whether the same locus is responsible for stripe-pattern variation outside of Lake Victoria cichlids using a hybrid cross between the nonstriped Lake Malawi species *Pseudotropheus demasoni* (*Pdem*; Fig. 1M) and striped species *Ps. cyaneorhabdos* (*Pcy*; Fig. 1K) that also differed in their skin *agrp2* expression (fig. S14). We obtained 270  $F_2$  hybrid individuals that were genotyped at the *agrp2* locus and phenotyped regarding their stripe patterns (Fig. 4, B and C; fig. S15; and supplementary text). The results revealed significant linkage between the *agrp2* allele and stripe presence (Fisher's exact test,  $P = 7.6 \times 10^{-8}$ ; table S2). The allelic variation at the *agrp2* locus explains more than 50% of the phenotypic variance in stripe patterns [Cox-Snell or Nagelkerke pseudo- $R^2$  from ordered logistic regression; table S3]. Nevertheless, the phenotypic distribution of  $F_2$  individuals (49 nonstriped and 221 striped individuals) differed from the Mendelian 3:1 ratio observed in the Lake Victoria crosses (chi-square test,  $P < 0.001$ ), providing evidence for additional minor modifier loci. These results strongly suggest that *agrp2* acts as a major determinant of stripe pattern absence or presence in Lake Malawi (as in the younger Lake Victoria radiation), but additional minor stripe modifiers have evolved or were recruited in the older Lake Malawi radiation (Fig. 4D).

The repeated evolution of horizontal stripes in East African cichlid radiations is facilitated by cis-regulatory evolution of *agrp2*. Despite its described role and function in the brain (18), we have discovered a hitherto unknown function for this gene in the skin that highlights notable functional similarities between *Agp2* and the mammalian Agouti (*Asip*) as well as teleost *Asip1* (19, 20). From what is known about proteins of the Agouti family, *Agp2* likely acts as an antagonist for the melanocortin receptors *Mclr* and/or *Mc5r* (21). Low *Agp2* levels would trigger stripe melanophore proliferation, pigment





**Fig. 3. Regulatory changes of *agrp2* are predictive of convergent stripe evolution.** (A) Densitree representation of phylogeny for the 24 examined species, including divergence time estimates (8) mya, million years ago. (B) Skin gene expression analysis of *agrp2* across adaptive radiations highlights the strong evolutionary association of stripes with low *agrp2* expression levels (phylogenetic ANOVA,  $P < 0.001$ ). Error bars indicate means + SD (C) A gene (locus) tree of the cis-regulatory element *enh.a* supports a single origin of striped alleles in Lake Victoria. Numbers present posterior probabilities  $>0.9$ . *Ajac*, *Aulonocara jacobfreibergi*; *Ceuc*, *Cheilochromis euchilus*; *Dcom*, *Dimidiochromis compressiceps*; *Hmel*, *Haplochromis melanopterus*; *Hser*, *Haplochromis serranus*; *Hthe*, *Haplochromis thereuterion*; *Jmar*, *Julidochromis marlieri*; *Jreg*, *Julidochromis regani*; *Lmul*, *Lamprologus multifasciatus*; *Mzeb*, *Maylandia zebra*; *Ncau*, *Neolamprologus caudopunctatus*; *Onil*, *Oreochromis niloticus*; *Pjoh*, *Placidochromis johnstoni*; *Ppun*, *Pundamilia pundamilia*.



**Fig. 4. A shared genomic basis of stripes across cichlid radiations.** (A) Stripes are associated with *agrp2* alleles in Lake Malawi hybrid  $F_2$  individuals ( $Pdem \times Pcy$ ). Inheritance is not Mendelian, suggesting that additional genetic modifiers exist in Lake Malawi cichlids. (B and C)  $F_2$  hybrids homozygous for the *Pcy* *agrp2* allele (C/C, showing stripes, black lines) and *Pdem* allele (D/D, no stripes) exhibit clear stripe pattern differences. Parental species are shown in Fig. 1K (striped) and Fig. 1M (nonstriped). (D) Summary of the known (black) and unknown (gray) aspects of the genetic control of cichlid horizontal stripes.

dispersion, and/or pigment production (stripe patterns present), whereas high levels would block these processes (no stripe patterns) (21). Expression levels of *agrp2* thereby act as a switch controlling stripe presence and absence. In Lake Victoria, expression-level differences

seen to be caused by several mutations in a 1.1-kb intronic regulatory region (*enh.a*; Fig. 2B) that push the expression of *agrp2* levels above or below a threshold that determines the stripe phenotype. Such a threshold-based molecular on-off switch may have permitted the frequent

loss as well as reevolution of stripes within East African cichlids. Although the presence of stripes appears to be controlled by differential expression of the same gene (*agrp2*), causal genetic variants must differ among the independent radiations of Lakes Victoria, Malawi, and Tanganyika (Fig. 4D and figs. S12 and S13). The intermediate phenotypes obtained from the Malawi cross (Fig. 4A), together with the lack of the dorso-lateral stripe in the CRISPR-Cas9 mutants (Fig. 2H), provide evidence for additional modifier loci determining stripe presence (Fig. 4D). However, those seem generally less prominent in the young (<15,000 years old) Lake Victoria radiation compared to the older (2 million to 4 million years old) Lake Malawi radiation (supplementary text).

Regulatory variation of *agrp2* provides a molecular basis for the repeated evolution and loss of stripe patterns across cichlid species flocks. Recurrent regulatory evolution at the *agrp2* locus constitutes an example of regulatory tinkering (22, 23) that might have facilitated the ease and speed of the evolution of both converged and diverged phenotypes that characterizes the East African cichlid radiations. The simplicity of such a threshold mechanism might have permitted the phylogenetically observed rapid losses and reevolutions of stripe patterns. Therefore, Stephen Jay Gould's predictions (1) appear questionable at this evolutionary scale, and if one were to replay the evolution of cichlid adaptive radiations, the results might be surprisingly similar: striped and nonstriped cichlids evolving again and again through regulatory evolution at the *agrp2* locus.

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declare no competing interests. **Data and materials availability:** Additional data have been deposited on Dryad (24) and are available in the supplementary materials.

#### SUPPLEMENTARY MATERIALS

[www.sciencemag.org/content/362/6413/457/suppl/DC1](http://www.sciencemag.org/content/362/6413/457/suppl/DC1)  
Materials and Methods  
Supplementary Text  
Figs. S1 to S15  
Tables S1 to S5  
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### Showing their stripes

The adaptive radiation of East African cichlids has led to more than 1200 species across a number of lakes. Across these species, many convergent traits have emerged, including the presence or absence of horizontal stripes. Kratochwil *et al.* show that the appearance or loss of stripes is related to changes in the *agouti-related peptide 2* gene, which acts as a kind of on-off switch for stripe generation (see the Perspective by Gante). This action has enabled rapid and repeated evolution of stripes across this speciose radiation.

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