Different sources of allelic variation drove repeated color pattern divergence in cichlid fishes

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Abstract
The adaptive radiations of East African cichlid fish in the Great Lakes Victoria, Malawi, and Tanganyika are well known for their diversity and repeatedly evolved phenotypes. Convergent evolution of melanic horizontal stripes has been linked to a single locus harboring the gene agouti-related peptide 2 (agrp2). However, where and when the causal variants underlying this trait evolved and how they drove phenotypic divergence remained unknown. To test the alternative hypotheses of standing genetic variation versus de novo mutations (independently originating in each radiation), we searched for shared signals of genomic divergence at the agrp2 locus. While we discovered similar signatures of differentiation at the locus level, the haplotypes associated with stripe patterns are surprisingly different. In Lake Malawi, the highest associated alleles are located within and close to the 5′ untranslated region of agrp2 and likely evolved through recent de novo mutations. In the younger Lake Victoria radiation, stripes are associated with two intronic regions overlapping with a previously reported cis-regulatory interval. The origin of these segregating haplotypes predate the Lake Victoria radiation since they are also found in more basal riverine and Lake Kivu species. This suggest that both segregating haplotypes were present as standing genetic variation at the onset of the Lake Victoria adaptive radiation with their more than 500 species and drove phenotypic divergence within the species flock. In summary, both new (Lake Malawi) or ancient (Lake Victoria) allelic variation at the same locus can fuel rapid and convergent phenotypic evolution.

Introduction
Understanding how genetic variation translates into phenotypic diversity is an important goal in evolutionary biology. Repeatedly evolved phenotypes are particularly interesting for the
study of the genetic basis of phenotypic diversity because they provide natural replicates that can inform if the same evolutionary mechanisms have recurrently generated these phenotypes (Kuraku and Meyer 2008; Protas and Patel 2008; Stern 2013; Elmer, et al. 2014; Kratochwil and Meyer 2015; Kratochwil, et al. 2018). Repeated evolution can result from either evolution though independent *de novo* mutations occurring in different species; or from preexisting variation that can be recruited via introgression or from standing genetic variation in a common ancestor (Stern 2013). Most *de novo* mutations are expected to be neutral or deleterious (Ohta 1992) whereas old standing genetic variation has likely already been purged from deleterious alleles due to previous selection. Adaptation from standing genetic variation is generally thought to be faster, as alleles reach fixation more quickly. Thereby standing genetic variation might facilitate rapid diversification (Barrett and Schluter 2008; Marques, et al. 2019). Ancestral variants can be more easily reassembled into new combinations whereas fixation of *de novo* mutations is predicted to result in a slower diversification process (Barrett and Schluter 2008; Hedrick 2013; Marques, et al. 2019). Accordingly, recent studies recognized the recruitment of alleles from standing genetic variation as an important evolutionary mechanism in driving the rapid phenotypic diversification found in adaptive radiations (Colosimo, et al. 2004; Hines, et al. 2011; Seehausen 2015; Lamichhaney, et al. 2016; Han, et al. 2017; Meier, Marques, et al. 2017; Bassham, et al. 2018; Malinsky, et al. 2018; Nelson and Cresko 2018; Salzburger 2018; York, et al. 2018; Lewis, et al. 2019; Svardal, et al. 2020; Kautt, et al. in press). However, with a few exceptions (Colosimo, et al. 2004; Hines, et al. 2011; Lamichhaney, et al. 2016; Meier, Marques, et al. 2017; Lewis, et al. 2019), most studies reporting evidence for the importance of ancestral standing genetic variation across whole genomes lacked knowledge of genotype-phenotype connections (Malinsky, et al. 2018; Nelson and Cresko 2018; Svardal, et al. 2020). As a consequence, the specific impact of old genetic variation on phenotypic diversification often remains elusive.

The adaptive radiations of cichlid fishes offer a great opportunity to investigate the contribution of standing genetic variation to rapid adaptive divergence, due to their exceptionally high diversity in species and the repeated evolution of multiple phenotypes (Meyer, et al. 1990; Meyer 1993; Stiassny and Meyer 1999; Kocher 2004; Genner and Turner 2005; Henning and Meyer 2014). Within the Great Lakes of the African Rift Valley, cichlids diversified into hundreds of endemic species in several lakes of different sizes and ages. In the three East African Great Lakes alone – Lake Victoria, Lake Tanganyika and Lake Malawi – more than 1,200 cichlid species evolved (Salzburger and Meyer 2004). Recent studies showed that the onset of the exceptionally rapid adaptive radiation in Lake Victoria, in which at least 500 species evolved within the past 15,000 years (Johnson, et al. 2000; Verheyen, et al. 2003; Wagner, et al. 2013), was fueled by high levels of genome-wide standing genetic variation (Seehausen 2004; Meier, Marques, et al. 2017). The Lake Victoria cichlid flock is
derived from divergent lineages of the geologically older Lake Kivu (Verheyen, et al. 2003) and adjacent rivers (Salzburger, et al. 2005; Meier, Marques, et al. 2017), which started diversifying about 100–200 thousand years ago (Verheyen, et al. 2003; Seehausen 2006; Genner, et al. 2007). The older radiation of Lake Malawi cichlids encompasses about 700 species (Turner, et al. 2001), which are believed to have evolved within the last 800 thousand years (Danley and Kocher 2001; Brawand, et al. 2014). Recently, whole genome resequencing revealed that standing genetic variation contributed to the high diversification rates of this adaptive radiation (Svardal, et al. 2020). Furthermore, standing genetic variation derived from ancestral lineages was also reported for Lake Tanganyika cichlids (Irisarri, et al. 2018), the oldest and phenotypically most diverse of the three East African cichlid fish adaptive radiations (Sturmbauer and Meyer 1992; Salzburger, et al. 2005; Koblmüller, et al. 2008).

Within and between these different adaptive radiations multiple phenotypes have evolved repeatedly (Stiassny and Meyer 1999). This is exemplified by melanic horizontal stripes, an adaptive phenotype that is often associated with shoaling behavior and a piscivorous feeding mode (Seehausen, et al. 2001). Previous work identified the gene agouti-related peptide 2 (agrp2, also called asip2b) as a major effect locus for stripe pattern divergence in African cichlids (Kratochwil, et al. 2018). The teleost-specific gene agrp2/asip2b and its paralogs have been previously associated with pigmentation phenotypes (Zhang, et al. 2010; Manceau, et al. 2011; Ceinos, et al. 2015). In zebrafish agrp2 is mainly expressed in the pineal gland. Biochemically it acts as an antagonist of melanocortin receptors (Zhang, et al. 2010). In cichlids, agrp2 has been demonstrated to also have a function in the skin, where it controls the presence of stripe patterns. While high expression of agrp2 inhibits stripe patterns, low expression permits their development. Yet, prior work could not identify the exact causal haplotypes and their evolutionary origin(s). The adaptive importance of horizontal stripes (Seehausen, et al. 2001), together with the detailed insights into the well resolved genotype to phenotype connection (Kratochwil, et al. 2018), make the agrp2 locus an ideal subject to investigate the role of preexisting standing genetic variation versus de novo mutations arising within the adaptive radiation in driving adaptive phenotypic divergence of rapidly evolving species flocks.

First, we addressed whether striped and non-striped fish of the parallel and independent radiations of Lake Victoria, Lake Malawi and Lake Tanganyika show the same signals of genomic divergence explaining the convergent evolution of stripe patterns. Next, we included genomic sequences of more basal lineages of the Lake Victoria radiation from Lake Kivu and adjacent rivers to trace back the evolutionary origin of the causal major effect allelic variants.
Results and Discussion

Stripe pattern convergence and diversification in African cichlid fish radiations

To reconstruct the evolutionary history of the haplotypes associated with stripe patterns, we investigated the genomic interval around the agrp2 gene with a combination of target enrichment (~30 kb agrp2 region ± 100kb) and whole-genome re-sequencing as the agrp2 locus was previously shown to be associated with horizontal stripes in cichlids of the three African Great Lakes (Henning, et al. 2014; Kratochwil, et al. 2018). Data were collected from 213 individuals from the three great African species flocks (number of individuals/species in Lake Malawi n=143/111, L. Tanganyika n=26/23, L. Victoria n=36/22; Supplementary Table S1).

We inferred a species tree of the sampled species based on 6,545 genome-wide randomly selected loci of 3 kb from 33 high-quality genomes. The phylogeny from this state-of-the-art, high-density data set agrees with previous reports based on mitochondrial (Meyer, et al. 1990), RAD-seq (Wagner, et al. 2013) and, most recently, genomic data (Malinsky, et al. 2018; Svardal, et al. 2020). All phylogenies show strong discordance between the stripe phenotype and phylogeny showing that stripes clearly evolved repeatedly (Fig. 1).

Stripes in Lake Malawi and Victoria radiations are associated with the same gene but different non-coding regions

Using whole-genome re-sequencing and target enrichment data, we calculated relative genetic differentiation ($F_{ST}$) between striped and non-stripped species for the cichlid radiations of Lakes Tanganyika, Malawi and Victoria over the 672,091 filtered bi-allelic single nucleotide polymorphisms (SNPs) called across the whole ~10 Mb scaffold 3 containing the agrp2 gene. Parallel evolution drives certain mutations to fixation independently in different populations and thereby acts on very local genomic regions. Therefore, we used the software Saguaro for $F_{ST}$ calculation which implements an algorithm that sets out to identify and pinpoint such regions using a Hidden Markov Model and a Neural Network, applied in an interleaved fashion. Saguaro then infers local relationships among individuals in the form of genetic distance matrices and assigns segments across the genomes to these topologies.

In the Lake Tanganyika radiations, we did not find regions of elevated $F_{ST}$ between striped and non-stripped species around agrp2 (Supplementary Fig. S1), although a link between agrp2 expression and stripes has been shown earlier (Kratochwil, et al. 2018).

The Lake Tanganyika species flock is more than 10 million years old and consists of several ancient independent radiations (Salzburger, et al. 2002; Clabaut, et al. 2005; Salzburger, et al. 2005; Koblmüller, et al. 2008; Takahashi and Koblmüller 2011) with a complex history of repeated colonization events (Nishida 1991; Salzburger, et al. 2002). Therefore, the missing
association of alleles within the agrp2 locus with stripes is likely explained by more complex genetic mechanisms of stripe formation and for example involves multiple cis-regulatory loci and/or trans-regulatory mechanisms as well as additional modifier loci.

Both adaptive radiations of Lakes Victoria and Malawi are composed of a single lineage of haplochromine cichlids which evolved within the last 2-4 million years in Lake Malawi and in Lake Victoria within 0.01 to 1 million years (Meyer, et al. 1990; Kocher 2004; Turner 2007; Brawand, et al. 2014). Among the 700 endemic Lake Malawi cichlids the agrp2 locus shows elevated differentiation among the littoral rock-dwelling mbuna which contains at least 200 species (Fig. 2A) (Danley and Kocher 2001). Within this lineage, the strongest differentiation between striped and non-striped species includes the 5′ untranslated region (UTR) of agrp2 ($F_{ST} = 0.85$ vs. scaffold mean $0.09$; Fig. 2A and 2B). In a gene tree inferred based on this region, the topology clearly separates striped mbuna from non-striped mbuna but not the Lake Victoria phenotypes (Fig. 2C). A single species, Petrotilapia nigra, is heterozygous for two of the three variants close to and within the 5′UTR (Fig. S2) and shows a very indistinct stripe pattern. Variants within region LM are unique to striped Lake Malawi mbuna and there is no association in non-mbuna nor Lake Victoria cichlids (Fig. 3). These variants therefore most likely constitute de novo mutations that evolved within the last ~300 kyr in the Lake Malawi mbuna radiation (Genner, et al. 2007). However, this association between the agrp2 locus and stripes vanishes when comparing the whole Lake Malawi dataset including non-mbuna species (Supplementary Fig. S1).

For the Lake Victoria radiation, the agrp2 locus was shown to be highly differentiated between striped and non-striped species (Fig. 2A). The two most differentiated regions ($F_{ST} = 0.87$ and $F_{ST} = 0.78$ vs. scaffold mean of 0.06) are directly upstream of the second exon and largely overlap (58.4% overlap) with a cis-regulatory active region (442,318–443,409) that was previously identified based on Sanger sequencing of three Lake Victoria species and experimentally tested using a transgenic reporter assay (Kratochwil, et al. 2018). Taken together the Lake Victoria regulatory interval (including both highly associated regions, LV 1 and LV 2, Fig. 2B) has a size of ~1.23 kb and is likely composed of several smaller cis-regulatory elements such as enhancers and/or silencers (Fig. 2B). In contrast to the topology of the region LM, the gene trees inferred from these two regions of highest differentiation (LV 1 and LV 2) clearly separate striped from non-striped Lake Victoria cichlids (Fig. 2D and 2E). This pattern supports the hypothesis that different regulatory regions at the same locus facilitate convergent evolution of stripe patterns across different cichlid radiations.

To further support the association of the identified cis-regulatory intervals found in Lakes Malawi and Victoria with stripe patterns, we employed a second, complementary approach, in which we assessed topology weights with TWISST (Van Belleghem, et al. 2017). The TWISST results strongly support a topology that groups species by stripe phenotype (Fig.
The adjacent gene, *atp6V0d2*, also exhibited pronounced topology grouping by stripe phenotype, but previous work did not reveal any fixed mis- or nonsense mutations nor differential expression (Kratochwil, et al. 2018).

Non-coding variants predict changes in transcription factor binding in highly divergent regions of both, Lake Malawi and Victoria

Both highly divergent regions are non-coding and might therefore contribute to variation in *agrp2* transcription and/or translation. The highly associated 90 bp region in Lake Malawi (LM, Fig. 2B and 2C) overlaps with the 5’ UTR of *agrp2*. 5’ UTRs can contain transcription factor binding sites (Barrett, et al. 2012; Lavallee-Adam, et al. 2017) but also have been shown to play important roles in post-transcriptional regulation (Araujo, et al. 2012) and could therefore lead to variation in transcript stability or translation rate. To provide additional evidence that the substitutions within and close to the 5’UTR of *agrp2* in Lake Malawi mbuna might influence *agrp2* transcription, we screened these regions for potential transcription factor binding sites (TFBSs) that are likely affected by the associated variants. For this, we used sequences from a representative non-striped and striped species (non-striped *Ps. demasoni*, and striped *Ps. cyaneorhabdos*) flanking the three variant sites within candidate region LM +/- 10 bp (Fig. 3 and Fig. S2, Table S2). The sequence flanking the first variant (position 438,598) contained 18 TFBSs of which 10 have a delta relative score of >0.1 (Materials and Methods, Table S2, Fig. S2) therefore suggesting a higher TF binding affinity in the non-striped (the species with high expression of the ‘stripe-repressor gene’ *agrp2*) than in the striped species. The TFs include *tfc3* that was associated with pigmentation previously (Dorsky, et al. 2000). For the second variant we did not identify TFBSs with a delta relative score of > 0.1. For the third variant in LM (position 438,687), 18 TFBSs were predicted within the 5’ UTR and five of these show a delta relative score of >0.1. These five transcription factors (TFs) have all (*snai2*, two variants of *tfap2e*, *tfap2a*, and *six1*) been linked to pigmentation (Sanchez-Martin, et al. 2002; Van Otterloo, et al. 2010; Yang, et al. 2019) and are expressed in the skin, melanophore or neural crest in zebrafish (https://zfin.org/). The neural crest is a highly migratory population of embryonic cells from which melanophores originate (Le Douarin and Kalcheim 1999). In conclusion, variants within the 5’ UTR of *agrp2* might have led to lower expression or transcript stability of *agrp2*. The resulting low expression of *agrp2* might in turn have triggered the *de novo* appearance of the stripe phenotype in Lake Malawi mbuna cichlids.

The most highly associated region in the Lake Victoria radiation (also when we included closely related riverine and Lake Kivu lineages; Fig. 4) is LV 1. We therefore also screened this region for potential TFBSs using the non-striped species *P. nyererei* (Pnye), and striped *H. sauvagei* (Hsau). Our analysis revealed high delta relative scores for several TFs with a known function in pigmentation pathways. For example, *tcfl5* at position 441,862...
belongs to a group of transcription factors involved in the Wnt signaling pathway. In zebrafish, Wnt signaling activates nacre, a zebrafish homolog of mitf, a key regulator of pigment synthesis, which in turn leads to pigment cell differentiation. Position 442,188 harbors a TFBS for zeb1, which in cichlids represses the expression of mitf (Albertson, et al. 2014). The sequence around variant position 442,399 has a TFBSs for sox18. The sequence around position 442,399 contains six more TFBSs belonging to the sox family of transcription factors with lower delta relative scores. Sox proteins including Sox18 regulate and interact during all stages of the melanocyte/melanophore life cycle (Harris, et al. 2010). Some transcription factor binding differences are shared between LM and LV (i.e. nfix, spi1, nr2c2(var.2), zeb1, rbpj, sox3, sox10) suggesting that transregulatory factors might be identical in both radiations while cis-regulatory elements are not.

These analyses support that divergence between striped and non-striped species in the radiations of East African cichlids is fueled by distinct cis-regulatory mechanisms controlling agrp2 expression, demonstrating that the recurrent involvement of the same gene does not necessarily mean that also the underlying causal mutations are the same.

The causal stripe haplotype in Lake Victoria evolved prior to the adaptive radiation

The finding of a single haplotype associated with stripes across all Lake Victoria species in our dataset is particularly interesting, as it suggests recruitment from ancestral standing genetic variation that was already present prior to the Lake Victoria cichlid radiation. To test this hypothesis, we analyzed the agrp2 locus in five species from ancestral lineages that are known (Verheyen, et al. 2003) to have diverged before the onset of the adaptive radiation in Lake Victoria (i.e. from Lake Kivu). The more distantly related lineages include non-striped species from Lake Kivu (‘Haplochromis’ gracilior) and Lake Edward (Thoracochromis pharyngalis). Two more closely related lineages include striped and non-striped species endemic to Lake Kivu (Haplochromis vittatus, and Haplochromis paucidens) and a striped riverine haplochromine species (Astatotilapia stappersii) from Kalambo River and Rusizi River (Greenwood 1979; Seehausen, et al. 2003; Meier, Marques, et al. 2017), which form a connection between Lake Kivu and Lake Tanganyika. Horizontal stripes are only present in the more closely related lineages (Fig. 1, and (Luc De, et al. 2001; McGee, et al. 2016; Meier, Marques, et al. 2017). To test if the causal alleles underlying stripe pattern divergence were already present in these more ancient haplochromine cichlid lineages, we analyzed the agrp2 locus in all striped ancestral species as well as in H. gracilior which was previously proposed as the source population of the Lake Victoria radiation (Verheyen, et al. 2003).

First, we calculated $F_{ST}$ between striped and non-striped phenotypes of Lake Victoria cichlids, and the ancestral lineages. From the most differentiated region (region LVRS, 538
bp, $F_{ST}=0.88$) we built a haplotype network. To reveal the evolutionary origin of the causal variants in the Lake Victoria superflock, we added species from Lake Malawi to the haplotype network (Fig. 4A). The Lake Victoria superflock is a group of 700 haplochromine cichlid species endemic to the region around Lake Victoria and nearby western rift lakes in East Africa (Meyer, et al. 1990; Verheyen, et al. 2003; Seehausen 2006; Genner, et al. 2007). The haplotype network shows that striped and non-striped Lake Malawi cichlids have different haplotypes than striped Lake Victoria species (Fig. 4A), as already suggested by the results above (Fig. 2D and 2E). Yet, striped Lake Victoria species share the same haplotype with the two striped species of the ancestral lineages of the Lake Victoria radiation (riverine A. stappersii and H. vittatus from the older Lake Kivu). We can therefore conclude that the cis-regulatory interval (Fig. 2A and 2B) must have evolved after their split from their common ancestor with Lake Malawi cichlids (2 – 4 Mya) but before their major radiation into the endemic species flocks of Lake Victoria and Lake Kivu (> 0.5 Mya, the age of Lake Kivu (Verheyen, et al. 2003)). To identify the ancestral lineage from which the haplotype at the agrp2 locus originated, we used ChromoPainter (Lawson, et al. 2012). For several Lake Victoria species, we calculated the per site probability of ancestry along haplotypes (Fig. 4B). Species from Lake Victoria acted as recipients with three ancestral striped and non-striped species acting as donors (i.e. ancestral haplotypes that are sources of recipient haplotypes). In total, we used three striped and non-striped recipient species each and ran two separate analyses for every striped and non-striped recipient haplotype: apart from the three ancestral lineages that acted as donors in the first analysis, all striped species had one non-striped within-lake donor acting as a control, while all non-striped species had one striped within-lake donor as control. Thereby, four donor species were competing in every analysis where a single recipient haplotype was tested – three ancestral donors and one within-lake control. If the cis-regulatory interval would have evolved within Lake Victoria, we would expect high per site probability of ancestry for these within-lake comparisons — this is not the case. This result underlines the role of old standing genetic variation from ancestral lineages in driving the repeated evolution of stripes in the Lake Victoria cichlid species flock. We found strong evidence that the cis-regulatory interval in striped Lake Victoria species (candidate regions LV 1 and LV 2, Fig. 2B) is most closely related to the riverine A. stappersii while other segments of the agrp2 locus are more closely related to the striped species from Lake Kivu (H. vittatus, Fig. 4B).

The region of highest differentiation between striped and non-striped species of the whole Lake Victoria superflock (Fig. 4A) overlaps with candidate region LV 1. This region shows a higher probability of ancestry from the striped donor species of the riverine haplochromine (A. stappersii) than from all other striped donors. Painwise comparisons
between the striped and non-striped species from Lake Victoria as well as its sister lineages revealed the same highly differentiated SNPs (Fig. 3). Therefore, incomplete lineage sorting due to ancestral standing genetic variation that was introduced into the lake by the haplochromine founders, is the most parsimonious explanation for the recurrent evolution of stripes in the Lake Victoria species flock.

Several recent studies across a wide range of study systems suggested that rapid speciation often involves ‘old genetic variants’ upon which selection can act (Han, et al. 2017; Meier, Marques, et al. 2017; Van Bellegem, et al. 2017; Cameron and Whitfield 2019; Edelman, et al. 2019; Jiggins 2019; Lewis, et al. 2019; Marques, et al. 2019). By a comprehensive analysis of the “stripe locus” with its well-resolved genotype-phenotype connection, we provide additional insights into how ancestral standing genetic variation at the root of adaptive radiations can facilitate rapid phenotypic divergence within species flocks.

By tracing the evolutionary history of highly associated variants, our study sheds light on the origin of the genetic basis of horizontal stripes, an adaptive phenotype that evolved repeatedly within the hundreds of species of the East African cichlid radiations (Seehausen, et al. 2001). Our findings show how different cis-regulatory regions of the same gene, agrp2, underlie rapid phenotypic divergence in the adaptive radiations of haplochromine cichlid fishes. We discovered that ancestral variants that form the genetic basis for stripe phenotypes in the Lake Victoria radiation predate the lake colonization and were introduced into it by the ancestors of this species flock and thereby allowed the repeated gain and loss of horizontal stripes within less than 100,000 years. In this radiation of more than 500 species, ancestral variants with an identified phenotypic effect (Kratochwil, et al. 2018) permitted the repeated phenotypic diversification and explosive speciation that characterizes the Lake Victoria cichlid fish adaptive radiation.

Materials and Methods

Experimental Model and Sampling Details
This study was performed in accordance with the rules of the animal research facility (T-16/13) of the University of Konstanz and the animal protection authorities of the State of Baden-Württemberg.

We obtained whole-genome re-sequencing data from mostly wild caught individuals from several different sources (see Supplementary Table S1). The whole genome samples from Malinsky, et al. 2018, Meier, Sousa, et al. 2017, Meier, Marques, et al. 2017, and McGee, et al. 2016 were obtained from wild caught individuals. The genomes from Brawand, et al. 2014 were inbred individuals from different laboratories.
Additionally, we sequenced 83 samples using target enrichment and 15 samples using whole genome re-sequencing. These samples were obtained from wild caught individuals from commercial breeders and maintained in the animal research facility of the University of Konstanz.

For most species we sampled one individual per species to obtain a comprehensive dataset that includes all major lineages. The reason why we have two samples of some species is that we used a combination of available genomes and target enrichment data for the agrp2 locus. This also allowed us to verify that different sequencing approaches did not introduce any biases during the downstream analyses (see Table S1).

Generally, we were very conservative with the phenotyping and classified a species as “striped” if either the male or the female possessed a horizontal stripe along the lateral side. All specimens that we sampled for this study were phenotyped using a photography chamber as described in Kratochwil, et al. (2018). To our knowledge there are few polymorphic species and these exceptions (Neolamprologus buescheri, Haplochromis phythophagus) we sampled ourselves and documented the phenotype accordingly. In our analyses these samples appear separately as if belonging to a striped and a non-stripped species.

**Species names and assignment**

Several of the analyzed species have different names across the literature. Because no commonly accepted taxonomy of cichlids is available we added a column „Current status“ in Table S1 which is based on the current classification of the ‘Catalog of Fishes of the California Academy’ (Fricke 2020).

**Target enrichment data**

Target enrichment data was produced using customized 120 nt baits with ~3x flexible tiling density by MYBaits for the 270 kb interval around agrp2. Baits were designed from the Pundamilia nyererei reference genome (Brawand, et al. 2014) which was curated by filling gaps of the genome assembly using Sanger sequencing reads so that the modified genome now contains the “stripe interval” (Kratochwil, et al. 2019). This version of the P. nyererei genome is available on Dryad: [https://doi.org/10.5061/dryad.bnzs7h467](https://doi.org/10.5061/dryad.bnzs7h467) (Kratochwil, et al. 2020).

DNA was either extracted from muscle tissue or from fin clips stored in EtOH following the DNeasy blood & tissue protocol (QIAGEN) or Genaxxon Genomic DNA Purification Mini Spin Column Kit (Genaxxon Bioscience GmbH), respectively. For library preparation, we used the Illumina TruSeq Nano HT Library preparation kit (Illumina Inc.) following the manufacturer’s guidelines. For the baits, we followed the MYBaits manual v3
and hybridized the probes at 65°C for 22 hours. Probes were sequenced in paired-end mode on a HiSeq 2500 system.

**Whole-genome re-sequencing**

DNA was extracted as explained above and DNA concentration was measured with fluorescence spectrophotometry by Qubit (Invitrogen). For library preparation, we used the Illumina TruSeq Nano HT Library preparation kit (Illumina Inc.) following the manufacturer’s guidelines. Samples were run on a Bioanalyzer 12000 Chip to assure a high quality of DNA libraries and afterwards amplified using PCR. Finally, size distributions of all libraries were checked on a Bioanalyzer HS chip before pooling them equimolarly. Sequencing was performed in paired-end mode (151PE) on a Hiseq X Ten platform (Illumina Inc.). Quality of the sequenced reads was assessed using MultiQC (Ewels, et al. 2016). The short-read data has been archived in the NCBI SRA database under the bioproject accession number PRJNA649899.

**Quality control and statistical analysis**

Illumina adapters were trimmed from the raw fastq reads using picard v2.17.11 (http://broadinstitute.github.io/picard) and reads were mapped to the curated *P. nyererei* reference genome (Kratochwil, et al. 2019; Kratochwil, et al. 2020) using bwa mem v0.7.12 (Li and Durbin 2009) and duplicate reads were marked with picard v2.17.11. Variants were called using the standard filter (\--min-mapping-quality 30 \--min-base-quality 20 \--min-supporting-allele-qsum 0 \--genotype-variant-threshold 0) and population options (population-based Bayesian inference model) in freebayes v1.1.0 (Garrison and Marth 2012). We decomposed multiple nucleotide polymorphisms in the VCF file into single nucleotide polymorphism (SNP) per line using a custom python script. The resulting VCF file was then hard-filtered using common hard filters from vcflib’s vcffilter ("QUAL > 1 & QUAL / AO > 10 & SAF > 0 & SAR > 0 & RPR > 1 & RPL > 1"), where "QUAL" refers to the quality of the variant site and thus removes really bad sites; “QUAL / AO > 10“ requires an additional contribution of each alternative allele observation of 10 log units (~ Q10 per read); „SAF > 0 & SAR > 0“ requires that alternative allele observations are present on both strands; and "RPR > 1 & RPL > 1“ requires that at least two reads with alternative allele observations are placed towards each side of the variant site. Additionally, we used VCFtools v0.1.15 (Danecek, et al. 2011) to remove indels, include only bi-allelic sites (\--max-alleles 2) and exclude sites that are missing in more than 5% of the samples (\--max-missing 0.05).
Finally, the filtered VCF file was normalized using vt normalize (Tan, et al. 2015). Mean effective sequencing depth, estimated from filtered VCF files using samtools flagstat, (Li, et al. 2009) can be found in Supplementary Table S1.

Next, we extracted phase informative reads using quality filters --base-quality 13 and --read-quality 10 before phasing with SHAPEIT v2.r790 (Delaneau, et al. 2011) and generated individual consensus fasta sequences with a custom python script. The consensus base was only kept when the site depth was above 5x coverage.

To calculate mean absolute genetic divergence ($d_{XY}$) and mean relative genetic differentiation ($F_{ST}$) between striped and non-striped species, we used the program Saguaro (Zamani, et al. 2013). With no prior assumption about the relatedness of the species, Saguaro creates local distance matrices for each region of the genome. This method infers local relationships among individuals in the form of genetic distance matrices and assigns segments across the genomes to these topologies. Thereby, it is possible that a single SNP that is alternatively fixed/highly associated between two populations results in a candidate region for high relative genetic differentiation.

To identify regions within the stripe interval that differ between stripe phenotypes, we ran TWISST (Van Belleghem, et al. 2017) for each lake separately. To reduce computation time, we used a subset of six species per lake (three striped and three non-striped) resulting in unrooted 15 topologies. For this, we followed the authors’ recommendations (https://github.com/simonhmartin/twisst/), which involved variant calling with GATK’s Haplotype Caller (Poplin, et al. 2018), filtering with VCFtools v0.1.15 (Danecek, et al. 2011), and phasing with beagle 4 (Browning and Browning 2007). Finally, we constructed neighbor joining trees for SNP windows (window size 50) in PhyML v3.1 (Guindon, et al. 2010). The curated P. nyererei reference genome assembly (Kratochwil, et al. 2019) had a 26936 bp zero coverage assembly gap on scaffold 3 that we removed from all plots as we also did not find this region in closely related species (Maylandia zebra, Astatotilapia calliptera).

Next, we inferred a gene tree for the loci with the highest $F_{ST}$ values (candidate regions LM, LV 1 and LV 2) using jModelTest v2.1.1 (Darriba, et al. 2012) to find the appropriate substitution model and BEAST 2 (Bouckaert, et al. 2014).

To compare the topology of the gene trees to a species tree we inferred the phylogenetic relationships using 33 genomes (Fig. 1). In brief, we mapped the 33 whole genome sequences to the Oreochromis niloticus genome assembly (NCBI: GCA_000188235.1) which is more complete (higher scaffold N50) than the P. nyererei genome. Variant calling and filtering steps were performed as described earlier. In the python script used to generate individual consensus fasta sequences we applied a maximum missingness filter of 0.75 that excludes sites on the basis of the proportion of missing data. Then, from each genome we extracted loci with a maximum physical extent of 3,000 bp each
of which a minimum of 2,000 sites had to be covered. These genome-wide loci were selected randomly, requiring a minimum distance of 100 kb between loci resulting in a total of 6,545 genome-wide loci. Next, we inferred single gene trees of all loci using IQ-tree 1.6.9 (Nguyen, et al. 2015) with the ModelFinder option (Kalyaanamoorthy, et al. 2017) for automatic selection of the appropriate model of evolution and with 100 rounds of ultra-fast bootstrapping (Hoang, et al. 2018) and estimation of the Shimodaira–Hasegawa-like approximate likelihood ratio test (Guindon, et al. 2010) respectively. Ultimately, we built the species tree using all 6,545 gene trees in ASTRAL-III (Zhang, et al. 2018). All trees were illustrated with FigTree v1.4.0.

To provide additional supporting evidence that the substitutions close to and within the 5'UTR of agrp2 in Lake Malawi mbuna could have an effect on agrp2 transcription, we screened the two divergent haplotypes for transcription factor binding sites (TFBSs) using JASPAR (Fornes, et al. 2020). For this, we extracted the flanking sequences of each SNP (+/-10 bp) within candidate region LM (Fig. 3 and Fig. S2) and screened for TFBSs above a conservative threshold of 0.85 (Kwon et al. 2012) of the relative matrix score in at least one of the two species. The difference between relative scores (delta Pdem - Pcy) of Ps. demasoni and Ps. cyaneorhabdos serves as an indicator of differential regulation of agrp2 between the non-striped and striped species. Table S2 gives a summary of all results as well as location of gene expression which we collected from ZFIN (https://zfin.org/).

We performed the same analysis for region LV 1 using the non-striped species P. nyererei, Pnye, and striped H. sauvagei, Hsau. We focused on LV 1, as it had the strongest association, also when we included closely related riverine and Lake Kivu lineages (see overlapping region LVRS in Fig. 4).

Since we find the association of non-coding regions within agrp2 to stripe phenotypes in all striped species from Lake Victoria in our data set, we traced the evolutionary origin of those haplotypes. First, we included striped and non-striped ancestral lineages of Lake Victoria and repeated the analysis of mean relative genetic differentiation (F_{ST}) between striped and non-striped species. The resulting region (labeled LVRS for Lake Victoria Region Superflock) of highest differentiation overlaps with the previous identified region LV1, however, is slightly shorter (538 bp instead of 687 bp). We used the R package pegas (Paradis 2010) to plot a haplotype network from this 538 bp region and included the Lake Malawi mbuna to show that the stripe haplotype is not shared between radiations.

Lastly, we used ChromoPainter (Lawson, et al. 2012) to elucidate haplotype relationships within the agrp2 locus in Lake Victoria. ChromoPainter models each recipient haplotype as a mosaic of the donor haplotypes while capturing which donors are essential to explain the recipient. There, we used different striped and non-striped donors (ancestral species which are the sources of admixture) and three striped, and three non-striped recipient
species, which represent the recipients of admixture. For visualization, we used the *pheatmap* function (Kolde 2019) in R. Scripts are available at the GitHub repository: https://github.com/sabineurban.

**Data Availability**

Fastq raw reads of all samples sequenced for this study (Table S1) are deposited at the NCBI Sequence Read Archive under the bioproject accession number PRJNA649899.

**Author Contributions**

S.U. wrote the paper with contributions from all authors; S.U. and C.F.K. conducted sample collection; S.U. did bench work; S.U. and A.N. conducted analyses; C.F.K. designed the study. C.F.K. and A.M. supervised the study.

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Fig. 1. Recurrent evolution of horizontal stripes in East African cichlids. (A) Phylogeny of East African cichlid lineages based on 6,545 genome-wide random loci of 3 kb. Node color indicates lake of origin (Malawi in blue, Tanganyika in green, Victoria in orange, and the Lake Victoria outgroups in purple) and if the species shows horizontal stripes (light color) or not (dark color). (B) Map of the African Great Lakes and surrounding lakes with dotted lines connecting species in the phylogeny to their lake of origin. Horizontal stripes are present in about one third of the species in all radiations. Photographs from top to bottom: Pundamilia nyererei, Haplochromis sauvagei, Astatotilapia stappersii, Pseudotropheus demasoni, Melanochromis kaskazini. Photograph credits: Jan Gerwin, Claudius Kratochwil, Adrian Indermauer, Sabine Urban.
Fig. 2. The same gene but different lake-specific regulatory regions are associated with the repeated evolution of stripe patterns in the adaptive radiations of haplochromine cichlids of Lakes Malawi and Victoria. (A) Association of stripes with genomic regions.
Black dots represent midpoints of every associated region ($F_{ST}$ value) as identified by Saguaro and black lines are smoothened local regressions between striped and non-striped species from Lake Victoria (top) and Lake Malawi mbuna species (bottom). This is plotted together with topology weights for topologies in which striped and non-striped species are reciprocally monophyletic (orange bars Lake Victoria, blue bars Lake Malawi mbuna). Each value gives the proportionate contribution of a particular taxon tree to the full tree with values ranging from 0 to 1. An example for such a topology in which striped species are reciprocally monophyletic is provided for both radiations. (B) Gene structure of $agrp2$ with regions of elevated $F_{ST}$ ($F_{ST} > 0.75$). Grey boxes indicate two isoforms of the 5'UTR that harbors variants associated with stripe divergence in mbuna Lake Malawi cichlids. (C) Unrooted gene tree from the region of highest differentiation in Lake Malawi (LM, 90 bp). The two monophyletic groups of non-striped Malawi cichlids differ in two variants at position 438,598 and 438,657 (Fig. 3 and Fig. S2). Numbers represent posterior probabilities. (D) Unrooted gene tree inferred from the region of highest differentiation in Lake Victoria (LV 1, 687 bp). (E) Unrooted gene tree from the region of second highest differentiation in Lake Victoria (LV 2, 411 bp).
Fig. 3. Different lake-specific substitutions are associated with stripe pattern evolution in Lakes Malawi and Victoria. (A) Grey represents the “non-striped” allele of the *P. nyererei* reference genome. All non-reference alleles within candidate region LM are indicated in blue. Substitutions in LM are not shared between Lake Victoria and Lake Malawi cichlids suggesting that they originated *de novo* in Lake Malawi mbuna. Orange indicates substitutions that were found in candidate region LV 1 that are associated with stripe patterns in Lake Victoria cichlids. While those haplotypes are divergent between striped and non-striped cichlids in Lake Victoria this is not the case in Lake Malawi mbuna.

Black arrows highlight substitutions that are highly divergent between striped and non-striped species (>90% frequency difference since some striped species are heterozygous, see also Fig. 4A and Fig. S3). (B) A screen for potential transcription factor binding sites in a non-striped (*Ps. Demasoni, Pdem, in Malawi and P. nyererei, Pnye, in Victoria*) and striped (*Ps. cyaneorhabdos, Pcy, in Malawi and H. sauvagei, Hsau, in Victoria*) representative species revealed that associated transcription factors in Lake Malawi cichlids are distinct from those in Lake Victoria cichlids. Shown are those TFs with the highest delta relative score. Variable positions in the motif sequence are written in bold.
Fig. 4. The evolutionary origin of the major effect haplotype of Lake Victoria stripe divergence predates the adaptive radiation of Lake Victoria. (A) Haplotype network of the highest $F_{ST}$ region (LVRS, 538 bp) between striped and non-striped species from the Lake Victoria outgroup.
Victoria superflock shows that there are both a stripe and a non-stripe haplotype present in the endemic radiation of haplochromine cichlids in Lake Victoria. Mismatches represent heterozygous individuals, i.e. not all species that were assigned a stripe phenotype are homozygous for the "stripe haplotype". Interestingly, these species do not show a complete stripe pattern consisting of a dorsolateral and a midlateral stripe but either show only one stripe (*H. paropius*) or represent a striped individual of a species (*H. phythophagus*) that displays a polymorphism by presence/absence of horizontal stripes. After $F_{ST}$ calculation we included the Lake Malawi mbuna which do not share the Lake Victoria stripe haplotype. However, the riverine *A. stappersii*, the striped species from Lake Kivu (*H. vittatus*), and all striped Lake Victoria cichlids share the stripe haplotype. (B) ChromoPainter analysis visualizes the probability of haplotypic segments to be shared among lineages and thereby the ancestral relationships at the *agrp2* locus of striped and non-striped recipient species. Every horizontal line in the heatmap represents one haplotype of a recipient species. The structure of *agrp2* is given at the bottom and grey dashed lines connect a SNP in the ChromoPainter ancestry matrix to its relative region within *agrp2*. This heatmap visualization of the results illustrates a high probability of ancestry in darker colors. Candidate regions LV 1, LV 2 and the 538 bp region from panel A (region LVRS, indicated by dark orange dashed lines) of striped recipient species are more closely related to the riverine species (*A. stappersii*). In contrast, in non-striped recipient species these regions are more closely related to the non-stripe donor species from Lake Kivu (*H. gracilior*). Candidate region LM (indicated at the top in blue) does not show a clear donor-recipient signal since there is also a high probability of relatedness of non-stripe Lake Victoria recipients to our control of a striped Lake Victoria donor species (*H. sauvagei*).