Genome sequences of *Tropheus moorii* and *Petrochromis trewavasae*, two eco-morphologically divergent cichlid fishes endemic to Lake Tanganyika

C. Fischer¹², S. Koblmüller¹, C. Börger¹, G. Michelitsch³, S. Trajanoski³, C. Schlötterer⁴, C. Guelly³, G. G. Thallinger²⁵ & C. Sturmbauer¹⁵

With more than 1000 species, East African cichlid fishes represent the fastest and most species-rich vertebrate radiation known, providing an ideal model to tackle molecular mechanisms underlying recurrent adaptive diversification. We add high-quality genome reconstructions for two phylogenetic key species of a lineage that diverged about ~3–9 million years ago (mya), representing the earliest split of the so-called modern haplochromines that seeded additional radiations such as those in Lake Malawi and Victoria. Along with the annotated genomes we analysed discriminating genomic features of the study species, each representing an extreme trophic morphology, one being an algae browser and the other an algae grazer. The genomes of *Tropheus moorii* (TM) and *Petrochromis trewavasae* (PT) comprise 911 and 918 Mbp with 40,300 and 39,600 predicted genes, respectively. Our DNA sequence data are based on 5 and 6 individuals of TM and PT, and the transcriptomic sequences of one individual per species and sex, respectively. Concerning variation, on average we observed 1 variant per 220 bp (interspecific), and 1 variant per 2540 bp (PT vs PT)/1561 bp (TM vs TM) (intraspecific). GO enrichment analysis of gene regions affected by variants revealed several candidates which may influence phenotype modifications related to facial and jaw morphology, such as genes belonging to the Hedgehog pathway (*SHH*, *SMO*, *WNT9A*) and the BMP and GLI families.

With 1727 described species¹, cichlid fishes are among the most species-rich teleost fish families. Their hotspot of biodiversity lies in East Africa, and in particular the three Great Lakes, Malawi, Victoria and Tanganyika². Despite a large degree of similarity pointing to recurrent evolution of eco-morphologically equivalent species³, the three cichlid radiations show important differences with respect to species numbers, evolutionary age of lineages, diversity of parental care patterns and the degree of morphological divergence²⁴. This is likely due to different sets of colonizing species and most importantly due to their different evolutionary age.

With an age of 9–12 million years (myr)⁵⁻⁶, Lake Tanganyika is by far the oldest of these lakes. Due to its old age, the Lake Tanganyika species assemblage is at a mature stage, so that it comprises the largest genetic and phenotypic diversity among the East African cichlid radiations, but further diversification proceeds predominately without much eco-morphological innovation⁷. Upon colonization of the emerging lake, the cichlids took advantage of the window of ecological opportunity and rapidly diversified⁸. In fact, two colonizing lineages underwent hybridization at the very onset of the radiation, an event that might have triggered or boosted the start⁹. The Lake Tanganyika radiation holds a key position for the entire modern African cichlid fauna, in that three of the newly emerging lacustrine lineages managed to colonize surrounding rivers, so that the radiation repeatedly swept over the boundaries of the maturing lake⁸⁻¹⁰. Three of the emerging lineages, the non-mouthbrooding

¹Institute of Biology, University of Graz, Graz, Austria. ²Institute of Biomedical Informatics, Graz University of Technology, Graz, Austria. ³Center for Medical Research, Medical University of Graz, Graz, Austria. ⁴Institut für Populationsgenetik, Vetmeduni Vienna, Vienna, Austria. ⁵BioTechMed-Graz, Graz, Austria. ⁶email: gerhard.thallinger@tugraz.at; christian.sturmbauer@uni-graz.at
Lamprologini, the mouthbrooding Orthochromini and some early Haplochromini such as the ancestors of the genera *Pseudocrenilabrus* and *Serranochromis*, left the lake at various stages of lake maturation to colonize particular surrounding water bodies. One group of early haplochromines continued to evolve in the lake-swamp-river interface towards more elaborate maternal mouthbrooders, demarcated by increased sexual dimorphism and eggspots on the anal fin, the so-called modern haplochromines. These modern haplochromines not only colonized most river systems all over southern and eastern Africa but re-entered the—at this time already much deeper and mature—Lake Tanganyika ecosystem, to evolve into the endemic Lake Tanganyika tribe Tropheini. Thus, the Tropheini managed to break into an ongoing and already complex lacustrine radiation, while its non-lacustrine sisters spread across several river systems to seed radiations in emerging lakes along their routes of riverine dispersal.

The Lake Tanganyika-endemic tribe Tropheini represents the sister group of all modern haplochromines outside the lake and diverged from these ~ 3–9 mya. That five out of the 29 species of the Tropheini both occur in the lake itself and upstream in tributary rivers and/or parts of the Lukuga River, the lake's only outflow, might be owed to their swamp-river origin. This is why we decided to sequence and compare the genomes of two ecologically divergent species of the endemic Lake Tanganyika tribe Tropheini. In terms of genetics, the modern haplochromines, including the Tropheini, are iconic as their generalist riverine-adapted genomes repeatedly underwent recurrent adaptive modifications upon ecological opportunity—provided by newly emerging lakes. It has been suggested that ecologically and phenotypically flexible species adapted to seasonally unstable river habitats can outcompete other colonizers in seeding lacustrine radiations, as they can rapidly accommodate empty niche space via phenotypic plasticity. According to the flexible stem hypothesis, a phenotypically plastic population is subdivided into alternative adaptive phenotypes and subsequently adaptive genetic factors are sorted during speciation to proceed further via genetic accommodation and genetic assimilation. In the course of adaptive divergence during repeated adaptive radiations, genic evolutionary events such as gene duplications appear to be recurrently shaped by ecological opportunity, in combination with geographic fragmentation events, episodes of bottlenecks and population expansions, as well as repeated admixes or fusions in hybridization events caused by climate-induced lake level fluctuations. Along with divergence and incidental gene flow, gene duplication and selection events apparently reshaped the genotypes. On the phenotype level, the evolutionary success of East African cichlids has been attributed to particular key innovations including (1) the functional decoupling of oral and pharyngeal jaws facilitating the exploitation of diverse trophic niches, (2) the adaptation of the visual system to different water turbidities, and (3) parental care and male mating coloration driven by sexual selection facilitating reproductive isolation. At this stage, the suite of genetic mechanisms modifying the genomic substrate underlying the enormous phenotypic eco-morphospace covered by cichlids remains largely unknown (see for a recent review).

The first major steps towards understanding the molecular mechanisms behind those divergent morphologies were taken by elucidating the genomes and transcriptomes of five cichlid species: *Oreochromis niloticus* representing an outgroup lineage, *Neolamprologus pulcher* representing a Tanganyikan substrate brooder lineage, and three modern haplochromines, namely *Astatotilapia burtoni* representing a riverine lineage, *Maylandia zebra* representing Lake Malawi and *Pundamilia nyererei* representing Lake Victoria. This study found evidence for an excess of gene duplications in the East African lineage compared to *Oreochromis* and other teleosts, an abundance of non-coding element divergence, accelerated coding sequence evolution, expression divergence along with transposable element insertions, and regulation by novel microRNAs. The study also revealed genome-wide diversifying selection on coding and regulatory variants, some of which recruited from ancient polymorphisms. High quality (HQ) genome drafts based on Pacific Biosciences (PacBio) data became available especially in the last two years. HQ drafts of *Simochromis diagramma* (East Africa, Lake Tanganyika) and *Astatotilapia calliptera* (East Africa, Lake Malawi) were generated by the Sanger Institute (2018) and a HQ draft of *Archocentrus centrarchus* (Central America) was generated by the G10K-VGP group (2019); assemblies of the South American cichlids *Amphilophus citrinellus* (2014, University of Konstanz) and *Arindoacara coerleocpunctatus* (2015, Sanger Institute) are also available. The *O. niloticus* (ON) and *M. zebra* (MZ) genomes have recently (2019) been newly assembled and anchored with a high-coverage PacBio + genetic map approach; the genomes of *A. calliptera*, *A. centrarchus* and *S. diagramma* (not anchored) were reconstructed similarly. *Oreochromis niloticus*, *M. zebra*, *A. calliptera* and *A. centrarchus* are the only reconstructions on chromosome level (linkage groups). Seven HQ drafts received annotations from the NCBI Annotation Pipeline (S. diagramma not yet); *O. niloticus*, *A. calliptera* and *A. citrinellus* received annotations from Ensembl as well. These genomes cover species from the Great Lakes and rivers in Africa and from crater lakes in Central and South America (Fig. 1).

We present reconstructions of the two genomes of *Tropheus moorii* (TM) and *Petrochromis trewavasae* (PT), as well as sets of structural and functional annotations. The two species belong to two sublineages within the Tropheini that diverged ~ 2–6.5 mya and represent the deepest split within the tribe Tropheini. Aside from generating the genomic and transcriptomic data basis for two key species representing the modern haplochromines in Lake Tanganyika, the main interest in this study was in the genetic origin of the divergent facial and jaw morphologies of these morphologically diverse species. To this end we provide first insights on genetic variants potentially being involved in the morphological differentiation of the two study species.

**Results**

**Assemblies.** Based on the estimated genome sizes of ~ 900 Mbp (Supplementary Table S1), our sequencing efforts yielded sequence data with an average base coverage of ~ 1.5×, ~ 88×, ~ 34× and ~ 10.5× (PT) and ~ 1.2×, ~ 38×, ~ 29× and ~ 9.1× (TM) for Roche 454, Illumina PE, Illumina MP and PacBio, respectively (see Supplementary Table S23). The filtered sequence data was used to generate primary assemblies derived from different reconstruction algorithms (assemblers) and data combinations (see Methods). The final genome reconstructions
of the two species are based on meta-assemblies of these sets of primary assemblies. The meta-assemblies with the best scores based on misassemblies, contiguity and gene predictions were used in subsequent analyses. 

**Petrochromis trewavasae.** The primary assemblies exhibit assembly sizes from ~779 Mbp to ~966 Mbp (907 Mbp PacBio only; see Supplementary Table S11). The final assembly consists of 7261 scaffolds with a N50 of 1.84 Mbp, 1.44% of nucleotides are undetermined (N) and 90% of the assembled genome is contained in 885 fragments longer than 70 kbp. The total assembly size is 917.57 Mbp (Table 1).

**Tropheus moorii.** The primary assemblies exhibit assembly sizes from ~754 Mbp to ~952 Mbp (879 Mbp PacBio only; see Supplementary Table S11). The final assembly consists of 7662 scaffolds with a N50 of 1.64 Mbp, 1.29% of nucleotides are undetermined (N) and 90% of the assembled genome is contained in 657 fragments longer than 192 kbp. The total assembly size is 911.13 Mbp (Table 1). Both assembly sizes are in the expected range; k-mer spectra-based predictions hint to genome sizes of close to 900 Mbp (see Supplementary Table S11) and 900–1000 Mbp have also been reported for other cichlid genomes21,30.

Table 1. Assembly contiguity and size statistics: The assembled genomes consist of 917.57 and 911.13 Mbp for *P. trewavasae* and *T. moorii*, respectively. Count and number of bases for scaffolds and contigs are reported. Scaffolds were broken to contigs at stretches of Ns of length ≥ 10. Statistics on *O. niloticus* were obtained from NCBI and extended as necessary (in blue); technology-wise version 2 is comparable, version 4 is based on high-coverage PacBio and optical mapping data.

---

**Figure 1.** Time calibrated phylogeny of cichlid species with publicly available complete genomes. The phylogeny is based on 519 anchored genome loci either sequenced or extracted from the genome sequence6. The study species are highlighted in bold, species with annotated high quality genomes are in black and species with high quality assemblies (but not yet annotated genomes) are indicated in grey (Figure created with Inkscape https://inkscape.org).
In the following, we compare our results to published genomes and annotations of several cichlid fish with emphasis on *O. niloticus* and *M. zebra* due to their well-developed state. The latest versions (v4) of *O. niloticus* (44 × PacBio, newly anchored) and *M. zebra* (now 65 × PacBio and anchored) were published by Conte et al.\(^2^2\); the tendency with respect to earlier versions is clear, qualities of sequences and annotations are improved and the numbers of annotated structures were further increased. With respect to the gene length distributions (Supplementary Table S1), the contiguity measures achieved for PT and TM are satisfying and fall in the typical range, given the applied sequencing technologies and coverage (Table 1) for a comparison with *O. niloticus* versions see Supplementary Table S2, and for a general comparison with published fish genomes see Supplementary Table S23 of Vij et al.\(^2^9\).

**Annotations.** Structural annotation yielded ~40,300 (PT) and 39,600 (TM) genes and ~54,200 (PT) and 56,800 (TM) transcripts, respectively (Table 2); this is in line with the results of different annotation versions of ON (~30,200 to 42,600 genes). As to annotated features, PT and TM show similar numbers which often lie between those of version 2 and 3 of the respective ON annotations. For comparison, statistics for ON v2–v4 (the latest) are added, as ON received the most community effort and data for genome assembly and annotation of all cichlids (Supplementary Table S2). Prediction of long non-coding RNAs yielded 2782 and 2112 IncRNAs for PT and TM, respectively. With 57.7% and 63.2% a slight preference for the sense strand could be observed (Supplementary Table S3). Homology based functional annotation could be made for 41,970 (PT) and 43,918 (TM) of them, respectively (Table 3). Pfam domain mapping yielded 78,900 (PT) and 84,158 (TM) hits, respectively. RepeatMasker\(^2^7\) identified 31.1% (PT) and 30.0% (TM) of the genomes as repetitive, respectively; the largest proportions of classified repeat types were held by DNA transposons, LINEs and LTR transposons with ~13%, ~7% and ~2% (Table 4).

**Data availability and visualization.** The genome and transcriptome assemblies (FASTA), the structural and functional annotations (GFF3), read mappings (BAM) and additional Integrative Genomics Viewer (IGV)\(^3^3\) track files (short and long non-coding RNAs, repeats, ORFs, CpG islands, microsatellites, IPR and eggNOG domains, variant calls, read mappings, alternative splicing, and REAPR error calls; Fig. 2) are available at https://cichlidgenomes.tugraz.at.

**Quality evaluation.** Assembly quality was assessed with BUSCO\(^3^4\) and CEGMA\(^3^5\). BUSCO identified 98.3% and 98% of the 4584 proteins in the Actinopterygii database in complete form for PT and TM, respectively; 1.7% and 2% of the benchmarking universal single-copy orthologs (BUSCOs) were either fragmented or missing. These results compare well with those of published genomes and are generally on a par with those of the later versions of the *O. niloticus* genome drafts (Table 5). CEGMA identified all of the 248 core eukaryotic genes (CEGs) for both PT and TM (Table 6); CEGMA results for PT and TM transcriptome assemblies can be found in Supplementary Table S6. However, REAPR reports 17,166/11,992 (PT/TM) likely assembly errors (Supplementary Table S10); there are IGV tracks highlighting questionable regions to guide caution when analyzing in the vicinity (see Fig. 2). Completeness of conserved protein domains was assessed with DOGMA\(^3^6\). DOGMA found 91.8% and 90.5% of the 1051 expected conserved domains at a conserved domain arrangement size of 1 for PT and TM, respectively (Table 7).

**Comparative analysis.** We compared the genomes of PT and TM by mapping the raw reads of one species to the genome of the other species. This yielded 4,105,604 and 4,178,777 small variants (SMV; SNPs and InDels) in introns; (2) SV are (slightly) underrepresented in promoter, 5′UTR, coding, splice site, 3′UTR and intergenic regions; they are overrepresented in introns and splice sites (via overlap) (Fig. 3A).

SNPeff\(^3^7\) categorises variant effects on the gene into four groups based on the location and nature of the variant: ‘HIGH’, ‘MODERATE’, ‘LOW’, or ‘MODIFIER’, where the latter denotes non-coding variants or variants affecting non-coding genes, where predictions are difficult or there is no evidence of impact. In our analysis, more than 97% of the identified variants are classified as ‘MODIFIER’ (Table 9).

Genes of interest, which are possibly affected by mutations, are highlighted in Table 10 (see Supplementary Information for details on the gene selection and GO enrichment analysis, respectively). Here, we started to analyze genes which are related to the development of the viscerocranium (untargeted gene selection) and the pharyngeal system (targeted, i.e., biased gene selection based on literature—see Table S17); however, there are other GO terms of interest which are consistently enriched over different analysis approaches such as BMP signaling, for instance. A condensed GO analysis result for an untargeted approach (A2, see Table S14b) is shown in Table 11; here, gene categories are based on variant comparison groups (within and between species groups) combined with quantile ranking and thresholding ($p = 0.5$, i.e., median), and variant counts (‘mutation loads’) were used as criterion. The term ‘embryonic viscerocranium morphogenesis’ is enriched in the within and the
Table 2. Structural annotation statistic of PT and TM in comparison with ON: Structural annotation yielded ~ 40,300 and 39,600 genes, respectively. This is in line with the results of different annotation versions of ON (~ 30,200 to 42,600).
between species gene sets over all approaches (see Supplementary Table S16; genes belonging to this term were combined with genes from the targeted approach and used for further downstream analyses (see Supplementary Table S14a, Table S14b, Table S15, Table S16, Table S18 and Table S21). In the comparative analysis, biological species are coded as A (PT) and B (TM) (Table 11). The categories (AA, AB, BA and BB) refer to the within and between group comparisons. That is, there are mutations at the same genomic locations (nucleotides) which are either identical within and between species (referred to as, e.g., identical (AA) and redundantly identical (AB)) or nonidentical (referred to as, e.g., nonidentical (BB) and nonidentical (BA)); moreover, there are mutations which are unique to a group (referred to as, e.g., unique (AA) and unique (AB), i.e., at the genomic location there is only a variant in species A (unique (AA)) or there is only a variant between species A and B (unique (AB)), respectively. In the shown example, for SMV the calls for viscerocranium morphogenesis are symmetric except for the AB category (which fell below the threshold), i.e., the GO term is consistently enriched within and between species. Further analyses on the genes belonging to the term clearly verify the presence of shared and species-specific mutations in these genes (see example in Supplementary section Identification of genes putatively related to facial and jaw morphology). Hence, there is substantial variation in these genes which may drive changes in the manifestation of morphology. However, we cannot yet delineate possible effects from shared and non-shared variants.

Besides variants in the DNA structure, alternative splicing (AS) was analyzed. There are ~6200 AS events in ~2600 genes between sexes of each species and ~39,000 AS events in ~9400 genes between the two species (see Supplementary Table S13).

| Annotation database                        | P. trewavasae v1 | T. moorii v1 |
|--------------------------------------------|------------------|--------------|
| UNIPROT/NR (FUNCTIONAL ANNOTATION, BLAST)  | 41,970           | 43,918       |
| MEROPS PROTEASE DB                         | 1,920            | 2,115        |
| CAZYMES (DBCAN)                            | 572              | 551          |
| EGGNOG (FINOG)                             | 127,730          | 125,974      |
| BUSCO VERTEBRATA MODELS                    | 3,196            | 3,462        |
| SIGNALP (SECRETOME)                        | 5,899            | 6,016        |
| INTERPROSCAN 5                             | **640,654**      | **672,128**  |
| CCD                                        | 28,531           | 29,546       |
| COILS                                      | 22,201           | 23,355       |
| GEN3D                                      | 93,347           | 98,675       |
| HAMAP                                      | 477              | 572          |
| MOBIDBLITE                                 | 58,498           | 59,402       |
| PANTHER                                    | 36,336           | 38,455       |
| PFAM                                       | **78,900**       | **84,158**   |
| PIRSF                                      | 2,804            | 3,278        |
| PRINTS                                     | 50,894           | 53,148       |
| PRODOM                                     | 519              | 536          |
| PROSITEPATTERNS                            | 22,972           | 22,897       |
| PROSITEPROFILES                            | 60,769           | 64,001       |
| SFLD                                       | 181              | 237          |
| SMART                                      | 73,882           | 77,359       |
| SUPERFAMILY                                 | 69,102           | 72,987       |
| TIGRFAM                                    | 1,888            | 2,181        |
| TMHMM                                      | 39,353           | 41,341       |

Table 3. Functional annotation statistics: The number of proteins found in UniProt and NR are given. Furthermore, the table contains the number of proteins with putative protease (Merops) and carbohydrate activity (CAZymes), the number of orthologs in finOG, the number of proteins matching the BUSCO vertebrate models and the number of proteins with putative secretory signals (SignalP). Finally, the number of hits of the protein sequences for the various InterPro domain databases are presented.
Discussion

Assembly and annotation. Meta-assembly of a set of primary assemblies yielded high quality genome drafts with 918 and 911 Mbp for *Petrochromis trewavasae* and *Tropheus moori*, respectively. This is in line with the sizes between 900 and 1000 Mbp reported for other cichlid genomes and with the ~940 Mbp estimated by our assembly validation with REAPR. The latest *Oreochromis niloticus* assembly spans ~1 Gbp;—this variation in genome size may be due to biological differences but could also indicate that some portions of the repetitive DNA in the respective genome reconstruction (PT/TM) are not included in the assembly as a consequence of sequence collapse (e.g., collapsed repeats).

Typically, assemblies are based on genomic data from a single individual, which ideally stems from an inbred line. In this project, we assembled from 6 (PT) and 5 (TM) non-inbred individuals, respectively; this called for a more complex assembly approach. Furthermore, we performed de novo sequencing without any linkage or optical map data (as seen in the latest genome drafts of *O. niloticus* and *A. calliptera*) and the PacBio coverage was (with ~9–10 ×) considerably lower than that used for the assemblies of *O. niloticus* (44 × for v330 and v427) and *M. zebra* (16.5 × for v340, added on top of already high Illumina PE and MP coverages, and 65 × for v427). Still, both assemblies compare well with published genome drafts of comparable species with respect to typical metrics regarding gene content. BUSCO results (Table 5) show a low rate of ~4–8% (depending on database) of duplicated BUSCOs in PT and TM. This is slightly higher than the ~2–4% reported for other cichlid genomes, which may be a consequence of incorrect assembly of haplotypes from the non-inbred individuals. With respect to total BUSCOs identified, fragmented and missing BUSCOs, both the PT and TM genome reconstruction perform very well.

For both species, the annotations proved valid for first sensible downstream analyses, but certainly there are some gene models which may need further improvement, e.g., by repeated training of gene predictors; however, for AUGUSTUS, the central predictor, model evaluations already show good training states (see Supplementary Table S12). The most relevant sources of insufficient gene models might include gene fusions, splits and especially truncations, which are obvious under closer inspection—this is typical for early annotations, especially when the annotation pipeline is still under development. We observe a relatively low mean and median length of protein sequences (see Table 2) in both assemblies/annotations. This may reflect a systematic error in the generation process, e.g., InDels leading to frame shifts and, hence, wrong translations and premature stop codons. Investigation of this phenomenon showed non-triplet InDels; however, these are also found between, e.g., *O. niloticus* and *M. zebra* transcript models. Moreover, the rate of identified nonsense mutations in PT and TM is low (Table 9).

The NCBI and Ensembl annotation pipelines are state-of-the-art; additionally, the amount and diversity of RNA-seq data used for the annotation of, e.g., *O. niloticus* was much larger than was the case for either species in this project. Hence, the larger number of identified transcript isoforms (as well as the higher average numbers of exons per transcript) may be seen as straightforward consequences. However, the total number of exons in both species is on a par with the ON annotations. Interestingly, the number of gene models in PT and TM are also on a par with ON v3. As there is no well-established method to score the correctness of gene models (perhaps by a general structure check and a database-based similarity majority scoring), this is merely a comparison of

| Species | *P. trewavasae v1* | *T. moori v1* |
|---------|-----------------|--------------|
| GC percentage | 41 % | 41 % |
| **Total bases masked** | 285,037,142 | 31.06 | 273,442,180 | 30.01 |
| SINEs | 39,940 | 6,581,639 | 0.72 | 39,267 | 6,438,991 | 0.71 |
| LINEs | 191,893 | 70,153,888 | 7.65 | 188,268 | 65,197,769 | 7.16 |
| LTR transposons | 45,962 | 18,449,565 | 2.01 | 43,936 | 16,701,190 | 1.83 |
| DNA transposons | 382,758 | 117,387,485 | 12.79 | 379,174 | 114,770,288 | 12.60 |
| Unclassified | 220,179 | 56,236,845 | 6.13 | 216,706 | 54,110,764 | 5.94 |
| **Total interspersed repeats** | 268,809,422 | 29.30 | 257,219,002 | 28.23 |
| Small RNA | 2,665 | 386,061 | 0.04 | 2,672 | 388,465 | 0.04 |
| Satellites | 4,348 | 1,341,142 | 0.15 | 4,327 | 1,330,850 | 0.15 |
| Simple repeats | 299,777 | 13,269,318 | 1.45 | 290,876 | 13,375,298 | 1.47 |
| Low complexity | 42,780 | 2,083,961 | 0.23 | 42,064 | 2,013,702 | 0.22 |

Table 4. Repeat annotation statistics as determined by RepeatMasker.
Figure 2. IGV tracks and extended annotation example view. (a) The sequences, along with structural and functional information (mouse-over), are provided via IGV tracks. We added an extensive set of data tracks and annotations (not all shown) to facilitate quick downstream analyses. (b) Protein sequence-based data set with annotations of identified functional domains (Figures represent screenshots from https://cichlidgenomes.tugraz.at).
### Table 5. BUSCO results:

Identified genes are classified as ‘complete’ when their lengths are within two standard deviations of the BUSCO group mean length (i.e., within ~95% expectation). ‘Complete’ genes found with more than one copy are classified as ‘duplicated’; BUSCOs are expected to evolve under single-copy control, hence recovery of many duplicates may indicate erroneous assembly of haplotypes. Genes only partially recovered are classified as ‘fragmented’, and genes not recovered are classified as ‘missing’.

The latest versions of assemblies were used in all cases (i.e., V4 of *O. niloticus* and *M. zebra*). See BUSCO results for PT and TM transcriptome assemblies in Supplementary Table S5. Values are color coded according to the rank: Dark green, best; dark red, worst. BUSCO stands for benchmarking universal single-copy ortholog.

| Database: Actinopterygii | *P. trewavasae* | *T. moorii* | *O. niloticus* | *M. zebra* | *A. citrinellus* | *A. calliptera* | *H. burtoni* | *P. nyererei* | *N. bichirandri* |
|--------------------------|----------------|-------------|----------------|-----------|-----------------|---------------|-------------|-------------|-----------------|
| Complete BUSCOs (C)      | 4,306          | 4,492       | 4,396          | 4,483     | 4,472           | 4,428         | 4,467       | 4,455       | 4,584           |
| Complete and single-copy (S) | 4,319          | 4,266       | 4,319          | 4,271     | 4,302           | 4,349         | 4,349       | 4,248       |                 |
| Complete and duplicated (D) | 398           | 224         | 107            | 100       | 126             | 101           | 104         | 105         |                 |
| Fragmented (F)           | 189            | 224         | 107            | 100       | 126             | 101           | 104         | 105         |                 |
| Missing (M)              | 183            | 148         | 44             | 53        | 53              | 54            | 72          | 114         |                 |
| Total BUSCO groups searched | 4,584          | 4,584       | 4,584          | 4,584     | 4,584           | 4,584         | 4,584       | 4,584       |                 |

| Database: Vertebrata | *P. trewavasae* | *T. moorii* | *O. niloticus* | *M. zebra* | *A. citrinellus* | *A. calliptera* | *H. burtoni* | *P. nyererei* | *N. bichirandri* |
|---------------------|----------------|-------------|----------------|-----------|-----------------|---------------|-------------|-------------|-----------------|
| Complete BUSCOs (C) | 2,447          | 2,422       | 2,462          | 2,499     | 2,493           | 2,454         | 2,472       | 2,408       | 2,366           |
| Complete and single-copy (S) | 2,453          | 2,426       | 2,462          | 2,499     | 2,493           | 2,454         | 2,472       | 2,408       | 2,366           |
| Complete and duplicated (D) | 15            | 16          | 17             | 16        | 15              | 14            | 18          | 29          |                 |
| Fragmented (F)       | 14             | 13          | 14             | 13        | 14              | 14            | 11          | 8           |                 |
| Missing (M)          | 14             | 13          | 14             | 13        | 14              | 14            | 11          | 8           |                 |
| Total BUSCO groups searched | 2,586          | 2,586       | 2,586          | 2,586     | 2,586           | 2,586         | 2,586       | 2,586       |                 |

| Database: Metazoa | *P. trewavasae* | *T. moorii* | *O. niloticus* | *M. zebra* | *A. citrinellus* | *A. calliptera* | *H. burtoni* | *P. nyererei* | *N. bichirandri* |
|-------------------|----------------|-------------|----------------|-----------|-----------------|---------------|-------------|-------------|-----------------|
| Complete BUSCOs (C) | 956            | 955         | 955            | 955       | 955             | 954           | 954         | 954         | 954             |
| Complete and single-copy (S) | 888            | 880         | 858            | 894       | 900             | 896           | 896         | 896         | 896             |
| Complete and duplicated (D) | 68             | 75          | 42             | 42        | 37              | 37            | 37          | 37          | 37              |
| Fragmented (F)     | 4              | 1           | 4              | 3         | 3               | 3             | 3           | 3           | 3               |
| Missing (M)        | 3              | 1           | 3              | 3         | 3               | 3             | 3           | 3           | 3               |
| Total BUSCO groups searched | 978            | 978         | 978            | 978       | 978             | 978           | 978         | 978         | 978             |

### Table 6. CEGMA results:

Shown are the latest versions in all cases; for ON and MZ additionally to v4 (PacBio-based) v2 (Illumina PE + MP-based) is listed for comparison (as PT and TM were primarily constructed using the same technologies). Values are color coded according to the rank: Dark green, best; dark red, worst. CEG stands for core eukaryotic gene.

| Species | *P. trewavasae* | *T. moorii* | *O. niloticus* | *M. zebra* | *A. calliptera* | *H. burtoni* | *P. nyererei* | *N. bichirandri* |
|---------|----------------|-------------|----------------|-----------|-----------------|-------------|-------------|-----------------|
| v1      | 100           | 100         | 100            | 99.6      | 100             | 99.19       | 99.6        | 98.79          | 98.79           |
| v2      | 99.6          | 98.79       | 98.79          | 98.39     | 99.19           | 99.64       | 98.79       | 98.79          | 98.79           |
| v3      | 99.6          | 98.79       | 98.79          | 98.39     | 99.19           | 99.64       | 98.79       | 98.79          | 98.79           |
| v4      | 99.6          | 98.79       | 98.79          | 98.39     | 99.19           | 99.64       | 98.79       | 98.79          | 98.79           |

### Table 7. DOGMA results:

DOGMA samples a score transcriptome/proteome regarding its completeness of conserved protein domains provided as percentage of a defined core set (conserved domains are structural and functional building blocks of proteins). The analysis supports the notion (see mean and median protein lengths in Table 2) that gene models of protein-coding genes need improvement. Values are color coded according to the rank: Dark green, best; dark red, worst. CDA stands for conserved domain arrangement.

| Species | *P. trewavasae* | *T. moorii* | *O. niloticus* | *M. zebra* | *D. rerio* |
|---------|----------------|-------------|----------------|-----------|-----------|
| CDA size | #expected | #found | [%] | #found | [%] | #found | [%] | #found | [%] | #found | [%] |
| 1       | 1,051      | 965        | 91.82 | 953     | 90.49    | 1,049    | 99.81 | 1,044    | 99.33 | 1,043    | 98.67 |
| 2       | 441        | 322        | 73.02 | 329     | 74.60    | 410      | 90.93 | 308      | 90.25 | 309      | 89.21 |
| 3       | 167        | 101        | 60.48 | 94      | 56.29    | 133      | 79.64 | 133      | 79.64 | 132      | 79.04 |
| Total   | 1,659      | 1,388      | 83.66 | 1,374   | 82.82    | 1,583    | 95.42 | 1,575    | 94.94 | 1,558    | 93.91 |

Comparative analysis. We picked the two study species for the following reasons. *Tropheus moorii* is a highly successful algae browser found in large numbers in all types of rocky shore, while *Petrochromis trewavasae* is an algae grazer distributed at rocky shores on the western side of the lake, living in sympathy with *Tropheus*. The Tropheini comprise 3 predatory species, one omnivore, 10 algae browsers and 15 algae grazers. Algae grazers have chisel-like teeth to bite off filamentous algae from the rocky substrate, while algae grazers have comb-
subdivided into alternative adaptive phenotypes and subsequently adaptive genetic factors are sorted during speciation sampled in the natural environment, but better reflects intra-population diversity and ultimately the older Lake Tanganyika radiation. Two species and even a considerable amount within each species. The large amount of variation at the intraspecific level may in fact be owed to our approach of using several non-inbred F1 individuals of a single population. This degree of eco-morphological divergence is greater than in the much younger eco-morphological equivalents in Lake Victoria, but comparable with the eco-morphospace covered by the entire Lake Malawi flock. Interestingly, the genus *Tropheus* comprises about 120 mostly allopatric and in terms of color distinct populations and sister species that are morphologically similar. They all remained in the same trophic niche at all rocky shorelines throughout the lake. *Petrochilus trewavasae* does not show much color variation, has a restricted distribution at the southwestern shoreline of the lake and is a member of a complex and morphologically distinct grazer lineage including the much more diverse *P. polyodon* species complex. When considering the entire lineage, it underwent a similar evolutionary trajectory as *Tropheus*. It should be noted here that the generally much lower species number in Lake Tanganyika when compared to Lakes Malawi and Victoria also results from the different species concepts employed, in that several allopatric entities are treated as species in Lakes Victoria and Malawi, whereas as geographical varieties in the older Lake Tanganyika radiation. The comparative analysis presented here yielded, as expected, a large number of variant regions between the two species and even a considerable amount within each species. The large amount of variation at the intraspecific level may in fact be owed to our approach of using several non-inbred F1 individuals of a single population sampled in the natural environment, but better reflects intra-population diversity and ultimately the old evolutionary age of the lineage. We used GATK\(^4\) and DELLY\(^\delta\), two well established tools, for variant calling; however, the calling of variants is still not a well solved problem with often little overlap between results of different algorithmic routes (e.g., see\(^44,45\)). As to the reported statistics on variant effects, it is known that the state of the structural annotation and the used variant effect annotator strongly influence the results\(^46\). The analysis results presented here reflect the state of the genome reconstructions (v1).

| Basic statistics | PT vs TM | TM vs PT | PT vs PT | TM vs TM |
|------------------|---------|---------|---------|---------|
| Genome           | TM draft v1.0 | PT draft v1.0 | PT draft v1.0 | TM draft v1.0 |
| Data             | PT filtered reads | TM filtered reads | PT filtered reads | TM filtered reads |
| Number of variants processed | 4,105,604 | 4,178,777 | 356,428 | 577,124 |
| Number of multi-allelic vcf entries | 17,885 | 26,099 | 2,607 | 3,940 |
| Number of variants with effects\(^1\) | 9,046,617 | 8,610,805 | 742,872 | 1,305,064 |
| Genome total length | 911,126,605 | 917,573,940 | 917,573,940 | 911,126,605 |
| Genome effective length | 906,396,323 | 913,305,292 | 905,543,956 | 901,211,269 |
| Variant rate      | 1 variant in 220 bases | 1 variant in 218 bases | 1 variant in 2540 bases | 1 variant in 1561 bases |

| Small-scale local variants (SMV) |
|---------------------------------|
| Snp                | 3,081,328 | 3,159,251 | 241,245 | 416,002 |
| Ins                | 511,111  | 487,934  | 54,489  | 75,775  |
| Del                | 513,165  | 531,592  | 60,694  | 85,347  |
| Total              | 4,105,604 | 4,178,777 | 356,428 | 577,124 |

| Large-scale structural variants (SV) |
|-------------------------------------|
| Duplication                        | 3559     | 2247     | 2870     | 3628     |
| Deletion                           | 56,396   | 58,692   | 11,989   | 21,047   |
| Inversion                          | 1853     | 1218     | 1343     | 1692     |
| Translocation                      | 18,829   | 12,351   | 11,566   | 15,022   |
| Insertions                         | 13,205   | 14,981   | 1316     | 2147     |
| Total                              | 93,842   | 89,489   | 29,084   | 43,536   |

Table 8. Overview on inter- and intraspecies variant analysis result: The numbers represent heterogeneity between species (for PT vs TM and TM vs PT) and heterogeneity within species (for PT vs PT and TM vs TM). These numbers may include the net effect of technical issues (e.g., with assembly, annotation, mapping and calling algorithms). \(^{1\text{as determined by SNPeff}}\).
Figure 3. Variant location distribution and proportion of genes exhibiting a variant in a particular structural/functional region. (A) With respect to structural/functional regions the distribution of called small and structural variants is typical—i.e., it largely follows the proportion of the respective genomic region with respect to the entire genome. Around 60% of variants are located in gene introns, followed by ~25% in intergenic regions and ~8.5% in gene promoters. (B) Under the applied parameters for calling and filtering almost all genes (~96%) are affected by some small variant and ~50% by some structural variant. Interestingly, the proportions of genes exhibiting a SMV in the promoter (~92%; defined as 2 kbp upstream and 200 bp downstream of the TSS) or coding regions (~75%) are very high. Gene regions are defined as the gene body plus 5 kbp up- and downstream. SMV, small variant; SV, structural variant; GCOV, genome coverage of specific region; TSS, transcription start site.
of an ancestral population exploit distinct ecological niches in a new habitat, such as different food types. Within a single generation, plasticity in anatomy may lead to a fitness increase, e.g., more efficient food capture or processing, in each niche. Newly exposed phenotypic variation will be targeted by selection, and if the new environment is stable, the plastic phenotypes may be canalized through genetic assimilation. The assumption is that the molecular mechanisms for the plastic response also underlie the evolution of key phenotypes, i.e., genetic variation in the same molecules/signaling pathways, which enable plasticity, is targeted by selection and fixed in order to canalize the phenotype. In a recent study, the role of hedgehog (Hh) signaling in the craniofacial plasticity in teleosts has been highlighted, demonstrating that Hh levels tune the sensitivity to mechanical signals related to foraging conditions—where adaptive morphological changes in immediately affected structures, e.g., the pharyngeal bones, may propagate morphological changes to other craniofacial structures.48

Variants have been called in virtually all gene regions. About 99% have at least one—under the applied parameter settings—possible variant in the gene body or 5 kb up/downstream (Fig. 3B). Genes with at least one mutation were subjected to Gene Ontology (GO) analysis to get hints on possible interesting functional groups affected by more variants—i.e., the number of variants (or ‘mutation load’) was used as pointer for the probability of effective changes. The rationale behind this approach was the assumption of correctness of the infinitesimal model or the omnigenic model, respectively. One may expect that the observed phenotype shifts are not due to few high impact (usually coding region) variants but rather due to several ‘lower impact’ variants (in the used categories probably the ‘modifier variants’ which typically represent >90% of the mutation load). Even if at this stage the relevance of the variation in the selected genes is not clear, all listed genes have multiple calls regarding SMV and SV (Fig. 3B) which may increase chances of effective influences on phenotypes. Given their assigned functions reported in other organisms (Table 10), however, these genes are also worth being probed. For instance, five genes being related to nose and chin shape definition (DCHS2, RUNX2, GLI3, PAX1, and EDAR) have recently been identified in a human GWAS study,49 several variants in all these genes have also been found between the two species. Additionally, PAX3, KCTD15 and TBX family members (TBX1 and TBX10, but not TBX15) as previously reported) are in the result set; these genes have been related to facial morphology in humans in two other recent GWAS studies (Table 10). Particular focus of future downstream analyses should be on genes with stable differences in gene expression among the study species. As stated earlier, our focus lies on the differences in facial and pharyngeal shapes (see Supplementary Fig. S1). It is interesting that this simple method of unbiased variant counting (‘mutation load’) output the GO terms related to the morphogenesis of the viscerocranium reproducibly (see Supplementary Information), without giving a rather unspecific long list of GO terms. From the GO result follows the highlighting of several important signaling pathways: BMP signaling (e.g., bmp2, bmp4), Hedgehog (Hh) signaling (e.g., Shh, Gli family, Sec family, smo, med12, plc3), endothelin signaling (e.g., edn1, furin, dlx family), retinoic acid (RA) signaling (e.g., rere, rer2), and fibroblast growth factor (FGF) signaling (e.g., fgf8, fgf20b) (see Table 10 and Supplementary Table S21). All of these signaling networks are known to play roles in the regulation of vertebrate facial morphogenesis, and they interact. There are, for instance, strong co-operative and functional interactions between Shh and retinoic acid.41,42 A more in-depth comparative analysis of the observed gene variant distribution across the two species and its respective phenotypes was not carried out at this stage; this will be an important task for follow-up studies.

To summarize, the two new draft genomes add two monophyletic and eco-morphologically divergent key species that fill an important phylogenetic gap. Moreover, they represent the earliest offshoot of the so-called modern haplochromine cichlids, the most species-rich lineage of East African cichlids. While the Trophæini radiated within the confines of Lake Tanganyika, their allies spread over several rivers to seed additional radiations such as those in Lake Malawi and Victoria, where those reached comparable eco-morphological diversity.

Methods

Study species. The sampled specimens of P. moorii are F2 offspring of wild caught individuals from the Zambian section of the southwestern shore of Lake Tanganyika (08°38’ S 30°52’ E) near the village Nakumu, which were brought to the University of Graz in 2005. The P. trewavasae specimens used in this study are F1

| Expected impact category | PT vs TM | TM vs PT | PT vs PT | TM vs TM |
|--------------------------|----------|----------|----------|----------|
| [Count] [%]              | [Count] [%] | [Count] [%] | [Count] [%] |
| High                     | 13,761   | 0.15     | 12,131   | 0.14     |
| Moderate                 | 84,185   | 0.93     | 82,142   | 0.95     |
| Low                      | 149,561  | 1.65     | 142,956  | 1.66     |
| Modifier                 | 8,799,110| 97.26    | 8,373,576| 97.25    |

Table 9. Overview on putative effects of intra- and interspecies variants: Shown are variant effect annotations as determined by SNPEff37. The numbers represent heterogeneity between species (for PT vs TM and TM vs PT) and heterogeneity within species (for PT vs PT and TM vs TM). These numbers may include the net effect of technical issues (e.g., with assembly, annotation, mapping and calling algorithms).
| Gene | Description | Variant type | Variant location |
|------|-------------|--------------|------------------|
| Predicted: Insulin-Like Growth Factor-Binding Protein 3 (igfbp-3) | IGFBP-3 plays a role in regulating pharyngeal cartilage and inner ear development and growth in zebralis<sup>143</sup> | SMV | 3’ UTR, intron, exon, downstream, upstream |
| Predicted: Fibroblast Growth Factor 8-Like (fgf-8) ** | FGF-8 is active in mouse and rat bone cells in vitro, stimulating osteoblast proliferation in a MAPK-independent pathway and inhibiting osteoclastogenesis via a RANKL/OPG-independent mechanism<sup>144</sup>. Plays an important role in the regulation of embryonic development, cell proliferation, cell differentiation and cell migration. Required for normal brain, eye, ear and limb development during embryogenesis [UniProt] | SMV | 3’ UTR, intron, exon, downstream, upstream, splice |
| Predicted: Barx Homebox 1 (barx1) | BARX1 represses joints and promotes cartilage formation in the craniofacial skeleton in zebralis<sup>150</sup> | SMV | 5’ UTR, 3’ UTR, intron, exon, downstream, upstream, splice |
| Predicted: Distal-Less Homeobox 2 (dlx2) ** | Also related to craniofacial structures in humans<sup>150</sup> | SMV | 3’ UTR, intron, exon, downstream, upstream, splice |
| Predicted: Potassium Channel Tetramerization Domain Containing 15 (kctd15) * | KCTD15 regulates neural crest formation and, when mutated, results shown to regulate TFAPl2A, which has a critical role in neural crest formation and, when mutated, results in reduced snout length in mice, among other defects. Perhaps KCTD15 affects nasal tip shape in humans by influencing chondrocyte proliferation in the nasal septum | SMV | 3’ UTR, 3’ UTR, intron, downstream, upstream, splice |
| Predicted: Secreted Protein Acylated with Cysteine-rich (Osteonectin) (sparc) | SPARC is a cysteine-rich acidic matrix-associated protein which is required for the collagen in bone to become calcified, and it is also involved in extracellular matrix synthesis and promotion of changes to cell shape. SPARC is required for normal growth of zebralis otoliths<sup>141,142</sup> | SMV | 5’ UTR, 3’ UTR, intron, downstream, upstream, splice |
| Predicted: T-box 10 (tbx10) ** | Tbx10 has been shown to be under positive selection in East African cichlid fishes<sup>147</sup>. There are HOXD-DLX3-DLX4 clusters in vertebrates, respectively<sup>152</sup> | SMV | 3’ UTR, intron, downstream, upstream, splice |
| Predicted: T-box 1 (tbx1) ** | Tbx1 has been related to abnormal pharyngeal arch and facial development in mouse<sup>143</sup>. T-box genes make major contributions to craniofacial development (Tbx1, Tbx10, Tbx15, Tbx22) and to development of the brain (Tbx1, Eomes), mammary gland (Tbx2, Tbx3), pituitary gland (Tbx3, Tbx18), thymus (Tbx1), liver (Tbx3), lung (Tbx2, Tbx4, Tbx5), pigmentation (Tbx15) and the immune system (Tbx21), among others<sup>145</sup> | SMV | 5’ UTR, 3’ UTR, intron, downstream, upstream, splice |
| Predicted: Potassium Channel Tetramerization Domain Containing 15 (kctd15) * | KCTD15 regulates neural crest formation by affecting Wnt signaling and the activity of transcription factor AP-2 in zebrafish embryos and human cells<sup>146</sup>. In humans, expression of KCTD15 showed a highly focal effect limited to the nasal tip<sup>147</sup>. KCTD15 has been shown to regulate TFAPl2A, which has a critical role in neural crest formation and, when mutated, results in reduced snout length in mice, among other defects. Perhaps KCTD15 affects nasal tip shape in humans by influencing chondrocyte proliferation in the nasal septum | SMV | 3’ UTR, 3’ UTR, intron, downstream, upstream, splice |
| Predicted: FAS-Associated Factor 1-Like (FAF1) | FAF1 is disrupted in clef palate and has conserved function in zebrafish. Knockdown of zebralis FAF1 leads to pharyngeal cartilage defects and jaw abnormality<sup>148</sup> | SMV | 5’ UTR, 3’ UTR, intron, downstream, upstream, splice |
| Predicted: PR Domain Containing 1 With ZNF domain (preg1) | In zebrafish, misexpression of pregn1 inhibits the formation of dermoeipithelial structures and reduces expression of chordin, which encodes a BMP antagonist. Later in development pregn1/blimp1 is expressed in many tissues, including the pharyngeal arches<sup>149</sup> | SMV | 3’ UTR, intron, downstream, upstream, splice |
| Predicted: Arginine-Glutamic Acid Dipeptide (RE) repeats (rere) | RERE/Atrophin-2 is thought to function as a transcriptional co-repressor during embryonic development in Drosophila<sup>148</sup>. This transcriptional regulator is required for the normal patterning of the early vertebrate embryo, including the central nervous system, pharyngeal arches, and limbs. Consistent with a role as a transcriptional corepressor, RERE binds histone deacetylase 1 and 2 (HDAC1/2), and orphan nuclear receptors such as TRs in zebrafish<sup>150</sup>. It plays a role in bilateral symmetry in mice<sup>151</sup> and variants are also related to craniofacial structures in humans<sup>149</sup> | SMV | 3’ UTR, 3’ UTR, intron, downstream, upstream, splice |
| Predicted: Distal-Less Homeobox 2 (dlx2) ** | The DLX proteins are postulated to play a role in forebrain and craniofacial development. The gene family has been shown to be under positive selection in East African cichlid fishes<sup>152</sup>. There are DLX1-DLX2, DLX3-DLX4, DLX5-DLX6 clusters in vertebrates, linked to Hox gene clusters HOXD, HOXB, and HOXA respectively<sup>152</sup> | SMV | 3’ UTR, 3’ UTR, intron, downstream, upstream, splice |

Continued
| Gene | Description | Variant type | Variant location |
|------|-------------|--------------|------------------|
| Predicted: Retinoic Acid Receptor Gamma-A-Like (RARGA) | RARs are involved in embryonic patterning and organogenesis—with craniofacial skeletal deficiencies affecting Rarα/β-null mutants. In zebrafish, putative roles for retinoic acid receptors in the hindbrain, limbs and pharyngeal arches have been identified. | SMV, SV | 5' UTR, 3' UTR, intron, exon, downstream, upstream |
| Predicted: Wingless-Type Mmtv Integration Site Family Member 9A (wnt9a) | Shown to play a role in zebrafish palate morphogenesis. Ligand for members of the frizzled family of seven transmembrane receptors. Functions in the canonical Wnt/beta-catenin signaling pathway. Required for normal timing of Ihh expression during embryonic bone development, normal chondrocyte maturation and for normal bone mineralization during embryonic bone development. Plays a redundant role in maintaining joint integrity. | SMV, SV | 3' UTR, intron, downstream, upstream |
| Predicted: Transforming Growth Factor Beta 2 (tgfb2) | Shown to play a role in zebrafish palate morphogenesis. The majority of osteoblasts and chondrocytes in the craniofacial region are derived from cranial neural crest cells (CNCC), which produce the facial skeleton. TGF-β signaling plays a crucial role in craniofacial development, and loss of TGF-β signaling in CNCC results in craniofacial skeletal malformations. | SMV, SV | Downstream, upstream |
| Predicted: Mothers Against Decapentaplegic Homolog 5 (SMAD5) | Shown to play a role in zebrafish palate morphogenesis. Transcriptional modulator activated by BMP (bone morphogenetic proteins) type 1 receptor kinase. SMAD5 is a receptor-regulated SMAD [UniProt]. | SMV, SV | 5' UTR, 3' UTR, intron, exon, downstream, upstream |
| Predicted: Paired Box 9 (pax9) | Shown to play a role in zebrafish palate morphogenesis. Transcription factor required for normal development of thymus, parathyroid glands, ultimobranchial bodies, teeth, skeletal elements of skull and larynx as well as distal limbs [UniProt] | SMV, SV | 5' UTR, 3' UTR, intron, exon, downstream, upstream |
| Predicted: Fibroblast Growth Factor 10-Like (FGF10) | Shown to play a role in zebrafish palate morphogenesis. Plays an important role in the regulation of embryonic development, cell proliferation and cell differentiation. Required for normal branching morphogenesis [UniProt]. FGF10 is largely expressed in mesenchymal tissues and is essential for postnatal life because of its critical role in development of the craniofacial complex. Genetic mouse models have demonstrated that the dysregulation or absence of FGF10 function affects the process of palate closure, the development of salivary and lacrimal glands, the inner ear, eye lids, tongue taste papillae, teeth, and skull bones. Mutations within the FGF10 locus have been described in connection with craniofacial malformations in humans. | SMV, SV | 5' UTR, 3' UTR, intron, exon, downstream, upstream |
| Predicted: Ectodysplasin a Receptor (edar) * | EDAR, which has previously been linked to ear and tooth shape and hair texture, affects chin protrusion in humans. | SMV, SV | 5' UTR, 3' UTR, intron, exon, downstream, upstream |
| Predicted: Dachous Cadherin-Related 2 (dchs2) * | DCHS2, also related to cartilage, controls nose pointiness in humans. | SMV, SV | Intron, exon, downstream, upstream, splice |
| Predicted: GLI Family Zinc Finger 1 (gli1) | GLI1 (2 and 3) are involved in craniofacial development in mice. | SMV, SV | Exon, downstream, upstream |
| Predicted: GLI Family Zinc Finger 3 (gli3) * | GLI3 known to be involved in cartilage growth is linked to the breadth of a person's nostrils in humans. | SMV, SV | 5' UTR, 3' UTR, intron, exon, downstream, upstream, splice |
| Predicted: Runt-Related Transcription Factor 2 (runx2) * | RUNX2, which drives bone development, is associated with the width of the nose bridge, the upper area of the nose in humans. | SMV, SV | 5' UTR, 3' UTR, intron, exon, downstream, upstream |
| Predicted: Paired box 1 (PAX1) * | PAX1 known to be involved in cartilage growth is linked to the breadth of the nostrils in humans. | SMV, SV | 5' UTR, 3' UTR, intron, downstream, upstream |
| Predicted: Paired box 3 (PAX3) * | PAX3 influences the position of the nasion in humans. | SMV, SV | 5' UTR, intron, exon, upstream |

Table 10. Selected genes affected by variants. To narrow down the list of genes carrying variants, a targeted approach and GO enrichment analysis were performed; this table lists genes related to facial and jaw morphology. Shown results are filtered and simplified: (1) variant types and locations have been unified for transcript isoforms and (annotated) gene duplicates, and (2) they have been intersected between species comparisons. SMV, small variant(s); SV, structural variant(s). * Also identified in human to play a role in craniofacial morphology. ** Other family member identified in human to play a role in craniofacial morphology.
| Type               | GO ID        | Description                                                      | p value  |
|--------------------|--------------|------------------------------------------------------------------|----------|
| SMV:identical (AA) | GO:0030513   | positive regulation of BMP signaling pathway                     | 2.80E-08 |
| SMV:identical (AA) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 3.00E-06 |
| SMV:identical (AB) | GO:0030509   | BMP signaling pathway                                             | 3.20E-09 |
| SMV:identical (AB) | GO:0060972   | left/right pattern formation                                      | 1.40E-04 |
| SMV:identical (BA) | GO:0000536   | cartilage morphogenesis                                           | 3.80E-05 |
| SMV:identical (BA) | GO:0015102   | cartilage condensation                                            | 2.10E-04 |
| SMV:identical (BA) | GO:0048747   | muscle fiber development                                          | 1.30E-08 |
| SMV:identical (BA) | GO:0007517   | muscle organ development                                          | 5.70E-04 |
| SMV:identical (BA) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 1.30E-08 |
| SMV:identical (BB) | GO:0048048   | embryonic eye morphogenesis                                       | 6.00E-07 |
| SMV:identical (BB) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 8.20E-10 |
| SMV:identical (BB) | GO:0048048   | embryonic eye morphogenesis                                       | 1.20E-09 |
| SMV:nonidentical (AA) | GO:0030513   | positive regulation of BMP signaling pathway                     | 2.30E-08 |
| SMV:nonidentical (AA) | GO:0001501   | skeletal system development                                       | 7.50E-04 |
| SMV:nonidentical (AA) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 2.30E-06 |
| SMV:nonidentical (AB) | GO:0030509   | BMP signaling pathway                                             | 6.20E-11 |
| SMV:nonidentical (AB) | GO:0030513   | positive regulation of BMP signaling pathway                     | 6.60E-09 |
| SMV:nonidentical (AB) | GO:0043010   | camera-type eye development                                       | 1.60E-04 |
| SMV:nonidentical (AB) | GO:0006972   | left/right pattern formation                                      | 5.40E-06 |
| SMV:nonidentical (BA) | GO:0015102   | cartilage condensation                                            | 3.40E-09 |
| SMV:nonidentical (BA) | GO:0000536   | cartilage morphogenesis                                           | 2.40E-04 |
| SMV:nonidentical (BA) | GO:0043010   | camera-type eye development                                       | 1.40E-04 |
| SMV:nonidentical (BA) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 5.90E-13 |
| SMV:nonidentical (BA) | GO:0048048   | embryonic eye morphogenesis                                       | 1.90E-09 |
| SMV:nonidentical (BB) | GO:0030500   | regulation of bone mineralization                                 | 5.70E-06 |
| SMV:nonidentical (BB) | GO:0048747   | muscle fiber development                                          | 8.10E-05 |
| SMV:nonidentical (BB) | GO:0048048   | embryonic eye morphogenesis                                       | 2.20E-10 |
| SMV:nonidentical (BB) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 1.20E-09 |
| SMV:unique (AA)    | GO:0030513   | positive regulation of BMP signaling pathway                     | 1.60E-07 |
| SMV:unique (AA)    | GO:0048703   | embryonic viscerocranium morphogenesis                            | 3.50E-05 |
| SMV:unique (AB)    | GO:0030513   | positive regulation of BMP signaling pathway                     | 2.40E-08 |
| SMV:unique (AB)    | GO:0030509   | BMP signaling pathway                                             | 1.50E-07 |
| SMV:unique (AB)    | GO:0048747   | muscle fiber development                                          | 2.60E-05 |
| SMV:unique (BA)    | GO:0000536   | cartilage morphogenesis                                           | 1.70E-04 |
| SMV:unique (BA)    | GO:0048048   | embryonic eye morphogenesis                                       | 4.00E-07 |
| SMV:unique (BB)    | GO:0048703   | embryonic viscerocranium morphogenesis                            | 1.00E-06 |
| SMV:unique (BB)    | GO:0048747   | muscle fiber development                                          | 2.40E-05 |
| SMV:unique (BB)    | GO:0048703   | embryonic viscerocranium morphogenesis                            | 1.70E-09 |
| SMV:unique (BB)    | GO:0048048   | embryonic eye morphogenesis                                       | 1.80E-09 |
| SMV:identical (AA) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 4.40E-11 |
| SMV:identical (AB) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 4.00E-13 |
| SMV:identical (BA) | GO:0030510   | regulation of BMP signaling pathway                               | 2.30E-06 |
| SMV:identical (BA) | GO:0048048   | embryonic eye morphogenesis                                       | 8.30E-07 |
| SMV:identical (BA) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 2.60E-05 |
| SMV:identical (BA) | GO:00009953  | dorsal/ventral pattern formation                                  | 2.00E-05 |
| SMV:identical (BB) | GO:0030510   | regulation of BMP signaling pathway                               | 7.30E-07 |
| SMV:identical (BB) | GO:0048703   | embryonic viscerocranium morphogenesis                            | 7.90E-06 |
| SMV:identical (BB) | GO:00009953  | dorsal/ventral pattern formation                                  | 6.10E-06 |
| SMV:unique (AA)    | GO:0048703   | embryonic viscerocranium morphogenesis                            | 5.40E-06 |
| SMV:unique (AB)    | GO:0048747   | muscle fiber development                                          | 5.90E-05 |
| SMV:unique (AB)    | GO:0048703   | embryonic viscerocranium morphogenesis                            | 8.80E-06 |
| SMV:unique (BA)    | GO:0030510   | regulation of BMP signaling pathway                               | 9.20E-07 |
| SMV:unique (BA)    | GO:0030500   | regulation of bone mineralization                                 | 4.40E-06 |
| SMV:unique (BB)    | GO:0048747   | muscle fiber development                                          | 4.50E-09 |

Table 11. GO enrichment analysis result—biological process terms (condensed). This table shows results from approach A2 (see Supplementary Table S14b). Enrichment was assessed via a Fisher's exact test with a cutoff of p ≤ 0.001 and GO topology was accounted for (R package topGO, method weight). In the Type column biological species are coded as A (PT) and B (TTM); identical and nonidentical variants at same nucleotide positions, and unique variants are indicated. The categories (AA, AB, BA and BB) refer to the within and between comparison: identical (AA) means that the intraspecific variant(s) (SMV and SV) in this group have also been called in the related interspecific (AB) comparison at the same location, with nonidentical (AA) a different variant has been called at the same location (e.g., A → T within and A → G between species), with unique (AA) only within species A and with unique (AB) only between species A and B a variant was called at that position; the same holds for species B and the BB and BA categories. Comparisons have been conducted two-way, i.e., A vs B and B vs A; the groups were tested against a gene universe containing all genes with GO information (the dataset contains 7905 (PT) and 7688 (TM) GO terms in total). SMV: small variant(s) (SNPs and InDels); SV: structural variant(s) (insertions, deletions, duplications, inversions and translocations). See Supplementary Table S15 for detailed lists.
offspring of wild fish also from the southwestern shore, but further northeast near the village Katete (08°20'30"S 30°30'E) and were obtained from an ornamental fish importer. Collection of the parental generation of fish was carried out in the framework of a Memorandum of Understanding between the Department of Fisheries, Ministry of Agriculture and Cooperatives, Zambia, the Department of Biological Sciences at the University of Zambia in Lusaka, the Department of Zoology at the University of Graz, Austria, the Department of Behavioural Ecology at the University of Bern, Switzerland, and the Department of Zoology at the University of Basel, Switzerland, under the research permit issued to CSt by the Zambian Ministry of Home Affairs (permit number: SP006515). Sequence data presented here are based on DNA extractions of 6 P. trewa wasae and 5 T. moorii individuals; the specimens included both sexes and were about one year old.

Sequencing and laboratory procedures. We sequenced the genomic DNA extracted from the specimens above using several sequencing technologies: Illumina HiSeq paired-end 2 × 101 bp (300 bp and 600 bp fragment size), Illumina Nextera mate-pair 2 × 100 bp (1–6 kbp fragment size), 454 Life Sciences (~ 350 bp average read length; 8 and 20 kbps fragment size) and single-molecule real-time (SMRT) sequencing technology from Pacific Biosciences (PacBio) (~ 8000–9000 bp average read length after correction).

Laboratory-related methods (DNA extraction, library preparation and sequencing) have, in part, been previously described in the accompanying paper on the mitochondrial genomes of Platanus. In addition, we carried out two sequencing runs using second-generation Pacific Biosciences sequencing technology based upon one individual per species. DNA extraction was carried out in Graz, library preparation and sequencing at the Lausanne Genomic Technologies Facility: DNA was sheared in a Covaris g-TUBE (Covaris, Woburn, MA, USA) to obtain 20 kbp fragments. After shearing the DNA size distribution was checked on a Fragment Analyzer (Advanced Analytical Technologies, Ames, IA, USA). 5 μg of the sheared DNA was used to prepare one SMRTbell library with the PacBio SMRTbell Template Prep Kit 1 (Pacific Biosciences, Menlo Park, CA, USA) according to the manufacturer's recommendations. The resulting library was size selected on a BluePippin system (Sage Science, Inc., Beverly, MA, USA) for molecules larger than 11 kbp. The recovered library was sequenced on thirteen/sixteen (TM/PT) SMRT cells with P6/C4 chemistry and MagBeads on a PacBio RSII system (Pacific Biosciences, Menlo Park, CA, USA) at 240 min movie length.

For RNA-seq, total RNA from one male and female individual per species (pooled from the following tissues: liver, spleen, brain, heart and skeletal muscle) was extracted with Trizol as follows: tissue was homogenized with MagnaLyser and incubated with Trizol-tube 5 min at room temperature (RT); 200 μl Chloroform (/ml of Trizol) was added and shaken vigorously for 15 s, incubated for 2–3 min/RT and centrifuged at 12,200 rpm/4 °C/15 min; supernatant was transferred to a new 1.5 ml tube and 500 μl isopropanol (/ml of Trizol) were added; after vortexing, incubaion for 10 min/RT, centrifugation at 12,200 rpm/4 °C/10 min supernant was discarded and the pellet placed on ice immediately. The pellets were washed 2 times: add 1 ml EtOH 80% (~ 20 °C), centrifuge: full speed/4 °C/5 min discard supernatant and finally dried at 37 °C. Dried pellets were resuspended in 20 μl distilled water. RNA-seq libraries were derived from total RNA which was rRNA-depleted, normalized and sequenced on a single Illumina HiSeq 2500 lane per species.

General data (pre)processing. All pipelining and higher-level processing was done with R/Bioconductor, some minor pipelining in Bash and some workhorse functionality was written in C++ (called from R). For details on parameter settings for important steps/tools see Supplementary Table S22.

FastQC v0.10.1 was used for basic read quality evaluation. A custom k-mer spectrum-based approach using JELLYFISH v2.0.9 in (conjunction with a database of known technical sequences) and a De Bruijn-based approach (implemented in Minion from the Kraken v1.27–27 package) were used for the automatic identification of technical contaminants and suspicious sequences (based on expected frequencies). In addition, FastQScreen v0.4.4.2 was utilized for the species-specific identification of biological contamination and DeconSeq v0.4.3.65 for the removal of technical contaminants, Scythe v0.99466 for additional adapter trimming, CLC quality trim v4.2.0 for quality-score-based read trimming and Reaper v13.274 for further quality and complexity-based filtering. BBmerge v33.40.48 was used for overlapping paired-end read merging and FastUniq v1.1.99 for duplicate removal. Nextclip v1.2.80 was used for Nextera mate-pair read filtering and classification. 454 datasets were additionally filtered with sffToCA (Celera Assembler utility). BAMtools v2.4.0.71, SAMtools/BCFtools/HTSlib v1.4.72 and Picard tools v1.1.1973 were used for mapping and sequence file manipulations such as indexing, merging, sorting, and generation of subsets, removal of duplicate reads, and removal of PE contamination from MP libraries in sequence files. Proovread v2.13.10.74 was used for PacBio read correction utilizing all available Illumina PE data and the unigts created by MaSuRCA v2.3.2.75. SEECER v0.1.3.76 and Rcor corrector v1.0.2.77 were used for RNA-seq and Musket v1.1.78 for DNA-seq base-call correction. DNA-seq and RNA-seq datasets were preprocessed using the same pipeline (with different parameter settings); in general, two filter regimes were applied to each data set (‘stringent’/‘standard’ and ‘relaxed’) in preparation for different downstream use cases (see Supplementary Table S22). Genome sizes were estimated by a k-mer spectrum-based approach implemented in gce v1.0.2.78.

Genome assembly. From the perspective of the conducted meta-assembly, the algorithm implemented in MaSuRCA v2.3.2.75 (utilizes Celera Assembler v6.5.06) served as the core assembly procedure; all at this time available data sets (i.e., Illumina PE and MP, Illumina Nextera MP and 454 MP and SE) were used. Celera Assembler v8.3rc2 (CA)68 was used for the ‘PacBio only’ assemblies. As several individuals per species (all non-inbred diploids) have been sequenced in this project, heterozygosity was a concern. Hence, assembly algorithms specifically designed to better handle divergence were incorporated into the reconstruction approach: Platanus v1.2.11 is a recent assembler tailored to more sensibly deal with heterozygosity issues in genomic data (5 iterations; all
Illumina data sets were used); Redundans v0.12c42 (utilizes SSPACE353, GapCloser44, bwa45 and last65) also aims at providing more accurate and contiguous assemblies of highly heterozygous genomes (5 iterations; all Illumina data sets were used). The PBJelly Suite v15.8.243 (utilizes BLASR88) was used to incorporate the long sequence reads (PacBio) in a reference-guided process into the established assembly drafts (5 iterations). The diverse set of generated genome drafts was subjected to MeTRambler46 in an attempt to generate high quality consensus sequences. A custom algorithm, which takes into account several measures on probable misassemblies, contiguity and gene predictions (drawing information from QUAST90 and REAPR38), was applied to determine the best order of successive meta-assemblies.

**Genome finishing.** For another round of inter-scaffold gap closing, GMcloser91 (utilizes Nucmer92 / BLAST93 and Bowtie294) was applied on the meta-assemblies with PacBio and Illumina PE data. Finally, Sealer95 (utilizes Konnector, a part of the ABySS assembler pipeline96) was used with the Illumina PE (liberal) libraries for final gap filling and a custom GATK-based42 genome finishing (via Illumina PE back mapping and consensus recalling) was applied.

**Genome validation.** REAPR v1.0.1898 (utilizing SMAL v0.7.0.197) was used with the Illumina Nextera mate-pair (6 kbp) and Illumina PE (600 bp) libraries to evaluate the correctness of assemblies and QUAST v4.190 was applied for contiguity and gene prediction statistics. Completeness of the assemblies was assessed using CEGMA v2.535 (utilizing GeneWise v2.4.198, HMMER v3.099 and NCBI BLAST + v2.2.29 + 93) with parameter optimization for vertebrate genomes (~vrt) and BUSCO v3.0.224 (utilizing NCBI BLAST + v2.2.29 +, HMMER v3.199 and AUGUSTUS v3.2.100).

**Transcriptome assembly and RNA-seq read mapping.** The transcriptome assemblies were conducted with Trinity v2.3.2101,102 and the PASA2 v0.2.0 pipeline103 (utilizing GMAP v2014-12–06104, BLAT v36.115 and MySQL v5.7.12106); also Transdecoder v3.0.1102 was applied to identify candidate coding regions (used with MAKER3107). RNA-seq read alignments for other analyses were generally conducted with STAR v2.4.2a108 using default parameters.

**Genome annotation.** Structural annotations were performed based on experimental data from mRNA-Seq datasets. Additionally, information was drawn from transcript and protein models from selected publicly available datasets (Danio rerio, H. burtoni, M. zebra, N. brichardi, O. niloticus, and P. nyererei) and from further models in UniProtSwiss-Prot, nr/nt and UniRef90teleost. Functional annotation was primarily conducted via BLAST-based comparisons against mentioned databases and via a host of databases coordinated by InterProScan (see Table 2).

**Transcriptome assembly and RNA-seq read mapping.** The transcriptome assemblies were conducted with Trinity v2.3.2101,102 and the PASA2 v0.2.0 pipeline103 (utilizing GMAP v2014-12–06104, BLAT v36.115 and MySQL v5.7.12106); also Transdecoder v3.0.1102 was applied to identify candidate coding regions (used with MAKER3107). RNA-seq read alignments for other analyses were generally conducted with STAR v2.4.2a108 using default parameters.

**Functional annotation.** Structural annotations of coding genes and tRNAs was generated using the pipelines MAKER v3.0107 (utilising the gene finders GeneMark-ES v4.32109, AUGUSTUS v3.2.1100, SNAP v2013–11–29110 and tRNAcan v1.3.1111), Funannotate v0.5.5–v0.7.0.12 (FA) and BRAKER1 v1.8112 (utilising GeneMark-ET v4.3.2114 and AUGUSTUS v3.2.119); BRAKER1 was also used for AUGUSTUS training. In addition, gene models were created with String-Tie v1.3.22110 and Cufflinks v2.2.1110. All models were combined by EvidenceModeler v1.1111 (EVM) under the control of MAKER3. For non-coding RNAs, Infernal v1.1.2114, Rfam v12.1119 and FEELnc v0.1.0120 were utilized. The mRNA training set for FEELnc was derived from the FA/MAKER annotation data, where presumed ‘good’ gene models with similar structure to previously published models were selected; the IncRNA training set was generated by shuffling of the mRNA sequences. Microsatellites were called with MISA v1.021, CpG islands with EMBoss v6.6.0122, CpGplot and ORFs with EMBoss v6.6.0 getorf (and R post-processing). Repeats were determined using RepeatMasker v4.0.6123 (with RepBase v20160321123 and species-specific libraries generated with RepeatModeler v1.0.8124), RepeatScout v1.0.5125 and TRF v4.06126.

**Functional annotation.** Functional annotation was conducted using InterProScan v5.24–63.0127 (utilizing the databases CDD-3.14, Coils-2.2.1, Gene3D-3.5.0, Hamap-201605.11, MobiDBLite-1.0, PANTHER-11.1, Pfam-30.0, PIRSF-3.01, PRINTS-42.0, ProDom-2006.1, ProSitePatterns-20.119, ProSiteProteins-20.119, SPLD-2, SMART-7.1, SUPERFAMILY-1.75, TIGRFAM-15.0 and TMHMM-2.0c). Furthermore, under the control of FA the databases eggNOG v4.5.1128 (bNOG), MEROPS v12.0129, dbCAN v5.0130 and BUSCO vertebrata v3131 were used for similarity searches and SignalP v4.132 for identification of target location signal sequences.

**Final integration of all annotations was done with R 3.4.3/Bioconductor 3.6 using the packages data.**}

**DNA-seq read mapping.** Preprocessed reads were aligned in paired-end mode with BWA mem45 using the default parameters with –M and –R flags. Aligned reads were coordinate sorted with Picard SortSam v1.119 and indexed with SAMtools index v1.4.122. Duplicates were removed with Picard MarkDuplicates v1.119. The quality of the mappings was assessed with QualiMap v2.046.

**Comparative analysis—small (SMV) and structural variant (SV) calling—variant effect prediction.** The Genome Analysis Toolkit (GATK) v3.7 was used for local realignment of reads and the detection and filtering of SNP/InDel variants (referred to as small variant/s, SMV)49 as recommended by the GATK documentation; the HaplotypeCaller was applied—with a minimum score for variant emission of 10, for calling of 30, and a minimum pruning of 10. SMV with a quality score ≥ 30 were included in further analyses. DELLY v0.7.45 was applied to call structural variant/s (SV, insertions, deletions, duplications, inversions and translocations) with an insert size cut-off of 3 (for deletions) and a minimum paired-end mapping quality of 20. All variants
with a minimum of 5 broken read pairs supporting the variant as well as with a minimum length of 300 bp (for deletions, inversions, and duplications) were included in further analyses, as recommended by the DELLY documentation. Presumed variant effects were called with SNPeff v4.3r37. Whippet v0.11.133 was used for the calling of alternative splicing events. The comparative analyses were conducted in R 3.4.3/Bioconductor 3.6 using the packages data.table 1.12.2, GenomicFeatures 1.30.3, VariantAnnotation 1.24.5 and their dependencies.

GO analysis. To narrow down the candidate gene list, GO enrichment analysis was performed on the gene regions carrying variants using the R package topGO v2.30.134; the custom GO annotations were generated based on the InterProScan mappings. GO topology was accounted for (method weight) and enrichment was assessed via a Fisher’s exact test with a cutoff of \( p \leq 0.001 \). See details on the GO analysis in the Supplementary Information.

Ethics approval and consent to participate. Animal treatment reported in this paper complies with the standards of the Animal Welfare Act in Austria and the European Community Directive 86/609. Fish were kept in our certified aquarium facility at the Institute of Biology, University of Graz. Individuals were sampled by CSt and SK, euthanized using an overdose of clove oil and decapitated conforming to the Austrian Animal Welfare legislation. According to the Austrian Animal Experiments Acts (TVG, BGBI. Nr. 501/1989, last changed by BGBI. Nr. 162/2005), approval was not required because no experimental treatment was performed.

Consent for publication. Not applicable.

Data availability
The genome drafts were uploaded to EBI, TM: [GCA_902810505], PT: [GCA_902810495]; the genome and transcriptome assemblies (FASTA), the structural and functional annotations (GFF3), read mappings (BAM) and additional IGV33 track files are available at https://cichlidgenomes.tugraz.at.

Received: 5 May 2020; Accepted: 28 December 2020
Published online: 22 February 2021

References
1. Van der Laan, R. & Fricke, R. Eschmeyer’s Catalog of Fishes Family Group Names. http://www.calacademy.org/scientists/catalog-of-fishes-family-group-names (2020).
2. Greenwood, P. H. African cichlids and evolutionary theories. In Evolution of Fish Species Flock (eds Echelle, A. A. & Kornfeld, I.) 141–154 (University of Maine at Orono Press, Orono, 1984).
3. Muschick, M., Indermaur, A. & Salzburger, W. Convergent evolution within an adaptive radiation of cichlid fishes. Curr. Biol. 22, 2362–2368 (2012).
4. Wagner, C. E., Harmon, L. J. & Seehausen, O. Ecological opportunity and sexual selection together predict adaptive radiation. Nature 487, 366–369 (2012).
5. Tiercelin, J.-J. & Mondeguer, A. The geology of the Tanganyika trough. In Lake Tanganyika and its Life (ed. Coulter, G. W.) 7–48 (Oxford University Press, Oxford, 1991).
6. Irarissi, I. et al. Phylogenomics uncovers early hybridization and adaptive loki shaping the radiation of Lake Tanganyika cichlid fishes. Nat. Commun. 9, 3139 (2018).
7. Salzburger, W., Meyer, A., Baric, S., Verheyen, E. & Sturmbauer, C. Phylogeny of the Lake Tanganyika Cichlid species flock and its relationship to the Central and East African Haplochromine Cichlid Fish Faunas. Syst. Biol. 51, 113–135 (2002).
8. Salzburger, W., Mack, T., Verheyen, E. & Meyer, A. Out of Tanganyika: genesis, explosive speciation, key-innovations and phylogeny of the haplochromine cichlid fishes. BMC Evol. Biol. 5, 17 (2005).
9. Koblmüller, S. et al. Age and spread of the haplochromine cichlid fishes in Africa. Mol. Phylogenet. Evol. 49, 153–169 (2008).
10. Sturmbauer, C., Salzburger, W., Duftner, N., Schelly, R. & Koblmüller, S. Evolutionary history of the Lake Tanganyika cichlid tribe Lamprologini (Teleostei: Perciformes) derived from mitochondrial and nuclear DNA data. Mol. Phylogenet. Evol. 57, 266–284 (2010).
11. Sturmbauer, C., Levinton, J. S. & Christy, J. Molecular phylogeny analysis of fiddler crabs: test of the hypothesis of increasing behavioral complexity in evolution. Proc. Natl. Acad. Sci. U. S. A. 93, 10855–10857 (1996).
12. Joyce, D. A. et al. An extant cichlid fish radiation emerged in an extinct Pleistocene lake. Nature 435, 90–95 (2005).
13. Katongo, C., Koblmüller, S., Duftner, N., Mumba, L. & Sturmbauer, C. Evolutionary history and biogeographic affinities of the serranochromine cichlids in Zambian rivers. Mol. Phylogenet. Evol. 45, 326–338 (2007).
14. Sturmbauer, C., Koblmüller, S., Seef, K. M. & Duftner, N. Phylogeographic history of the genus Trophius, a lineage of rock-dwelling cichlid fishes endemic to Lake Tanganyika. Hydrobiologia 542, 335–366 (2005).
15. Meier, J. I. et al. Ancient hybridization fuels rapid cichlid fish adaptive radiation. Nat. Commum. 8, 14363 (2017).
16. Svardal, H. et al. Ancestral hybridization facilitated species diversification in the Lake Malawi Cichlid fish adaptive radiation. Mol. Biol. Evol. 37, 1100–1113 (2020).
17. Kullander, S. O. & Roberts, T. R. Out of Tanganyika: endemic lake fishes inhabit rapids of the Lukuga River. Ichthyol. Explor. Freshw. 22, 355–376 (2011).
18. West-Eberhard, M. J. Developmental Plasticity and Evolution (Oxford University Press, Oxford, 2003).
19. Rossiter, A. The Cichlid fish assemblages of Lake Tanganyika: ecology, behaviour and evolution of its species flocks. In Advances in Ecological Research (eds Begon, M. & Fitter, A. H.) 187–252 (Academic Press Ltd., London, 1995).
20. Malinsky, M. et al. Whole-genome sequences of Malawi cichlids reveal multiple radiations interconnected by gene flow. Nat. Ecol. Evol. 2, 1940–1955 (2018).
21. Brawand, D. et al. The genomic substrate for adaptive radiation in African cichlid fish. Nature 513, 375–381 (2014).
22. Liem, K. F. Evolutionary strategies and morphological innovations: Cichlid Pharyngeal Jaws. Syst. Biol. 22, 425–441 (1973).
23. Carleton, K. L., Dalton, B. E., Escobar-Camacho, D. & Nandamuri, S. P. Proximate and ultimate causes of variable visual sensitivity: Insights from cichlid fish radiations. Genesis 54, 299–325 (2016).
24. Maan, M. E. & Sefc, K. M. Colour variation in cichlid fish: Developmental mechanisms, selective pressures and evolutionary consequences. Semin. Cell Dev. Biol. 24, 516–528 (2013).
25. Salzburger, W. Understanding explosive diversification through cichlid fish genomics. Nat. Rev. Genet. 19, 705–717 (2018).
28. Thibaud-Nissen, F. et al. P8008 the NCBI eukaryotic genome annotation pipeline. *J. Anim. Sci.* 94, 184 (2016).
29. Zerbino, D. R. et al. Ensembl 2018. *Nucleic Acids Res.* 46, D754–D761 (2018).
30. Conte, M. A., Gammesinger, T. W.J., Bartie, K. L., Penman, D. J. & Kocher, T. D. A high quality assembly of the Nile Tilapia (*Oreochromis niloticus*) genome reveals the structure of two sex determination regions. *bioRxiv* https://doi.org/10.1101/099564 (2017).
31. Vij, S. et al. Chromosomal-level assembly of the Asian Seabass genome using long sequence reads and multi-layered scaffolding. *PLoS Genet.* 12, e1005954 (2016).
32. Smit, A. F. A., Hubley, R. & Green, P. RepeatMasker Open-4.0. http://www.repeatmasker.org (2015).
33. Simão, F. A., Waterhouse, R. M., Ioannidis, P., Kriventseva, E. V. & Zdobnov, E. M. BUSCO: Assessing genome assembly and annotation completeness with single-copy orthologs. *Bioinformatics* 31, 3210–3212 (2015).
34. Parra, G., Bradnam, K. & Korf, I. CEGMA: A pipeline to accurately annotate core genes in eukaryotic genomes. *Bioinformatics* 23, 1061–1067 (2007).
35. Dohmen, E., Kremer, L. P. M., Bornberg-Bauer, E. & Kemen, A. Domain-based transcriptome and proteome quality assessment. *Bioinformatics* 32, 2577–2581 (2016).
36. Cingolani, P. et al. A program for annotating and predicting the effects of single nucleotide polymorphisms, SnpEff. *Fly* 6, 80–92 (2012).
37. Hunt, M. et al. REAPR: a universal tool for genome assembly evaluation. *Genome Biol.* 14, R47 (2013).
38. Asalone, K. C. & de Andrade Souza, D. W. B. Genomic analysis of Cichlidae fish (African cichlids) using new generation sequencing technologies. *Mol. Ecol. Res.* 11, 159–174 (2011).
39. Finn, R. D. et al. Pfam: The protein families database. *Nucleic Acids Res.* 38, D217–D221 (2010).
40. McKenna, A. et al. The genome analysis toolkit: A MapReduce framework for analyzing next-generation DNA sequencing data. *Genome Res.* 20, 1297–1303 (2010).
41. Rausch, T. et al. DELLY: Structural variant discovery by integrated paired-end and split-read analysis. *Bioinformatics* 28, i333–i339 (2012).
42. Liu, Y. et al. Comparison of multiple algorithms to reliably detect structural variants in pears. *BMC Genomics* 21, 61 (2020).
43. Supernat, A., Vidarsson, O. V., Steen, Y. M. & Stokowy, T. Comparison of three variant callers for human whole genome sequencing. *Sci. Rep.* 8, 17851 (2018).
44. McCarthy, D. J. et al. Choice of transcripts and software has a large effect on variant annotation. *Genome Med.* 6, 26 (2014).
45. Gunter, H. M., Schneider, R. F., Karner, I., Sturmbauer, C. & Meyer, A. Molecular investigation of genetic assimilation during the rapid adaptive radiations of East African cichlid fishes. *Mol. Cell.* 26, 6634–6653 (2017).
46. Navon, D. et al. Hedgehog signaling is necessary and sufficient to mediate craniofacial plasticity in tetosods. *Proc. Natl. Acad. Sci. U. S. A.* 117, 19321–19327 (2020).
47. Boyle, E. A., Li, Y. I. & Pritchard, J. K. An expanded view of complex traits: From polygenic to omnigenic. *Nat. Rev. Genet.* 19, 455–471 (2018).
48. Navon, D. et al. Hedgehog signaling is necessary and sufficient to mediate craniofacial plasticity in tetosods. *Proc. Natl. Acad. Sci. U. S. A.* 117, 19321–19327 (2020).
49. Boyle, E. A., Li, Y. I. & Pritchard, J. K. An expanded view of complex traits: From polygenic to omnigenic. *Cell* 169, 1177–1186 (2017).
50. Adhikari, K. et al. A genome-wide association scan implicates DCHS2, RUNX2, GLI3, PAX1 and EDAR in human facial variation. *Nat. Commun.* 7, 11616 (2016).
51. Liu, E. et al. A genome-wide association study identifies five loci influencing facial morphology in Europeans. *PLoS Genet.* 8, e1002932 (2012).
52. Claes, P. et al. Genome-wide mapping of global-to-local genetic effects on human facial shape. *Nat. Genet.* 50, 414–423 (2018).
53. Lupu, G., Harris, W. A. & Lewis, K. E. Mechanisms of ventral patterning in the vertebrate nervous system. *Nat. Rev. Neurosci.* 7, 103–114 (2006).
54. Dworkin, S., Boglev, Y., Owens, H. & Goldie, S. J. The role of sonic hedgehog in craniofacial patterning, morphogenesis and cranial neural crest survival. *J. Dev. Biol.* 4, 24 (2016).
55. Szabo-Rogers, H. L., Smithers, L. E., Yakob, W. & Liu, K. J. New directions in craniofacial morphogenesis. *Dev. Biol.* 341, 84–94 (2010).
56. Zhou, H., Kim, S., Ishii, S. & Boyer, T. G. Mediator modulates GlI3-dependent Sonic hedgehog signaling. *Mol. Cell. Biol.* 26, 8667–8682 (2006).
57. Vihlais-Neto, G. et al. Bore controls retinoic acid signalling and somite bilateral symmetry. *Nature* 463, 953–957 (2010).
58. Clouthier, D. E., Garcia, E. & Schilling, T. F. Regulation of facial morphogenesis by endothelin signaling: Insights from mice and fish. *Am. J. Med. Genet. A* 152A, 2962–2973 (2010).
59. Fischer, C. et al. Complete mitochondrial DNA sequences of the Threadfin Cichlid (*Petrochromis tewaasa*) and the Blunthead Cichlid (*Tropheus moorii*) and patterns of mitochondrial genome evolution in cichlid fishes. *PLoS ONE* 8, e67048 (2013).
60. Andrews, S. FastQC A Quality Control tool for High Throughput Sequence Data. http://www.bioinformatics.babraham.ac.uk/projects/fastqc/ (2016).
61. Marçais, G. & Kingsford, C. A fast, lock-free approach for efficient parallel counting of occurrences of k-mers. *Bioinformatics* 27, 764–770 (2011).
62. Davis, M. P. A., van Dongen, S., Abreu-Goodger, C., Bartonicek, N. & Enright, A. J. Kraken: A set of tools for quality control and analysis of high-throughput sequence data. *Methods* 63, 41–49 (2013).
63. Wingett, S. W. & Andrews, S. FastQ Screen: A tool for multi-genome mapping and quality control. *F1000Res.* 7, 1338 (2018).
64. Schmieder, R. & Edwards, R. Fast identification and removal of sequence contamination from genomic and metagenomic datasets. *PLoS ONE* 6, e12788 (2011).
65. Martin, M. Cutadapt removes adapter sequences from high-throughput sequencing reads. *EMBnet J.* 17, 10–12 (2011).
66. Buffalo, V. Scythe. https://github.com/vsbuffalo/scythe (2014).
67. CLCbio Assembly Cell. https://www.qiagenbioinformatics.com/products/clc-assembly-cell (2015).
68. Bushnell, B., Rood, J. & Singer, E. BBMerge—Accurate paired shotgun read merging via overlap. *PLoS ONE* 12, e0185056 (2017).
69. Xu, H. et al. FastUniq: A fast de novo duplicates removal tool for paired short reads. *PLoS ONE* 7, e52249 (2012).
70. Leggatt, R. M., Clavijo, B. J., Clissold, L., Clark, M. D. & Caccamo, M. NextClip: An analysis and read preparation tool for Nextera Long Mate Pair libraries. *Bioinformatics* 30, 566–568 (2014).
71. Barnett, D. W., Garrison, E. K., Quinlan, A. R., Stromberg, M. P. & Marr, G. T. BamiTools: A C++ API and toolkit for analyzing and managing BAM files. *Bioinformatics* 27, 1691–1692 (2011).
72. Li, H. et al. The sequence Alignment/Map format and SAMtools. *Bioinformatics* 25, 2078–2079 (2009).
73. Broad Institute Picard Tools. https://github.com/broadinstitute/picard (2016).
74. Hackl, T., Hedrich, R., Schultz, J. & Förster, F. Proovread: large-scale high-accuracy PacBio correction through iterative short read consensus. *Bioinformatics* 30, 3004–3011 (2014).
75. Zimin, A. V. et al. The MaSuRCA genome assembler. *Bioinformatics* 29, 2669–2677 (2013).
76. Le, H. S., Schule, M. H., McCaulley, B. M., Himman, V. F. & Bar-Joseph, Z. Probabilistic error correction for RNA sequencing. *Nucleic Acids Res.* **41**, e109 (2013).
77. Song, L. & Florea, L. Rcorrector: efficient and accurate error correction for Illumina RNA-seq reads. *GigaScience* **4**, 48 (2015).
78. Liu, Y., Schröder, J. & Schmidt, B. Musket: A multistage k-mer spectrum-based error corrector for Illumina sequence data. *Bioinformatics* **29**, 308–315 (2013).
79. Liu, B. et al. Estimation of genomic characteristics by analyzing k-mer frequency in de novo genome projects. *arXiv:1308.2012* (2013).
80. Denisov, G. et al. Consensus generation and variant detection by Celera Assembler. *Bioinformatics* **24**, 1035–1040 (2008).
81. Kajitani, R. et al. Efficient de novo assembly of highly heterozygous genomes from whole-genome shotgun short reads. *Genome Res.* **24**, 1384–1395 (2014).
82. Prysacza, L. P. & Gabaldón, T. Redundans: An assembly pipeline for highly heterozygous genomes. *Nucleic Acids Res.* **44**, e113 (2016).
83. Boetzer, M., Henkel, C. V., Jansen, H. J., Butler, D. & Pirovano, W. Scaffolding pre-assembled contigs using SSPACE. *Bioinformatics* **27**, 578–579 (2011).
84. Luo, R. et al. SOAPdenovo2: an empirically improved memory-efficient short-read de novo assembler. *GigaScience* **1**, 18 (2012).
85. Li, H. & Durbin, R. Fast and accurate short read alignment with Burrows–Wheeler transform. *Bioinformatics* **25**, 1754–1760 (2009).
86. Frith, M. C., Wan, R. & Horton, P. Incorporating sequence quality data into alignment improves DNA read mapping. *Nucleic Acids Res.* **38**, e100 (2010).
87. English, A. C. et al. Mind the Gap: Upgrading genomes with Pacific biosciences RS long-read sequencing technology. *PLoS ONE* **7**, e47768 (2012).
88. Chaisson, M. J. & Tesler, G. Mapping single molecule sequencing reads using basic local alignment with successive refinement (BLASR): application and theory. *BMC Bioinform.* **13**, 238 (2012).
89. Wences, A. H. & Schatz, M. C. Metassembler: Merging and optimizing de novo genome assemblies. *Genome Biol.* **16**, 207 (2015).
90. Gurevich, A., Saveliev, V., Vyahhi, N. & Tesler, G. QUAST: Quality assessment tool for genome assemblies. *Bioinformatics* **29**, 1072–1075 (2013).
91. Kosugi, S., Hirakawa, H. & Tabata, S. GMcloser: closing gaps in assemblies accurately with a likelihood-based selection of contig or long-read alignments. *Bioinformatics* **31**, 3733–3741 (2015).
92. Kurt, S. et al. Versatile and open software for comparing large genomes. *Genome Biol.* **5**, R12 (2004).
93. Camacho, C. et al. BLAST+: Architecture and applications. *BMC Bioinformatics* **10**, 421 (2009).
94. Langmead, B. & Salzberg, S. L. Fast gapped-read alignment with Bowtie 2. *Nat. Meth.* **9**, 357–359 (2012).
95. Paulino, D. et al. Sealer: A scalable gap-closing application for finishing draft genomes. *BMC Bioinf.* **16**, 230 (2015).
96. Simpson, J. T. et al. ABySS: A parallel assembler for short read sequence data. *Genome Res.* **19**, 1117–1123 (2009).
97. Ponstingl, H. & Ning, Z. SMALT. https://www.sanger.ac.uk/science/tools/smalt-0 (2018).
98. Birney, E., Clamp, M. & Durbin, R. GeneWise and genomewise. *Genome Res.* **14**, 988–995 (2004).
99. Finn, R. D., Clements, J. & Eddy, S. R. HMMER web server: Interactive sequence similarity searching. *Nucleic Acids Res.* **39**, W29–W37 (2011).
100. Stanke, M. & Morgenstern, B. Augustus: A web server for gene prediction in eukaryotes that allows user-defined constraints. *Nucleic Acids Res.* **33**, W465–W467 (2005).
101. Grabherr, M. G. et al. Trinity: reconstructing a full-length transcriptome without a genome from RNA-Seq data. *Nat. Biotechnol.* **29**, 644–652 (2011).
102. Haas, B. J. et al. De novo transcript sequence reconstruction from RNA-Seq: reference generation and analysis with Trinity. *Nat. Protoc.* **8**, 1494–1512 (2013).
103. Haas, B. J. et al. Improving the Arabidopsis genome annotation using maximal transcript alignment assemblies. *Nucleic Acids Res.* **31**, 5654–5666 (2003).
104. Wu, T. D. & Watanabe, C. K. GMAP: A genomic mapping and alignment program for mRNA and EST sequences. *Bioinformatics* **21**, 1859–1875 (2005).
105. Kent, W. J. BLAT—The BLAST-like alignment tool. *Genome Res.* **12**, 656–664 (2002).
106. Oracle Inc. MySQL. https://www.mysql.com (2016).
107. Cantarel, B. L. et al. MAKER: An easy-to-use annotation pipeline designed for emerging model organism genomes. *Genome Res.* **18**, 188–196 (2008).
108. Dobin, A. et al. STAR: ultrafast universal RNA-Seq aligner. *Bioinformatics* **29**, 15–21 (2013).
109. Lomsadze, A., Ter-Hovhannisyan, V., Chernoff, Y. O. & Borodovsky, M. Gene identification in novel eukaryotic genomes by self-training algorithm. *Nucleic Acids Res.* **33**, 6494–6506 (2005).
110. Korf, I. Gene finding in novel genomes. *BMC Bioinf.* **5**, 59 (2004).
111. Schattner, P., Brooks, A. N. & Lowe, T. M. The tRNAscan-SE, snoscan and snoGPS web servers for the detection of tRNAs and snoRNAs. *Nucleic Acids Res.* **33**, W666–W689 (2005).
112. Palmer, J. M. Funanotate: A fungal genome annotation and comparative genomics pipeline. https://github.com/nextgenusf/funanotate (2016).
113. Hoff, K. J., Lange, S., Lomsadze, A., Borodovsky, M. & Stanke, M. BRAKER1: Unsupervised RNA-Seq-based genome annotation with GeneMark-ET and AUGUSTUS. *Bioinformatics* **32**, 767–769 (2016).
114. Lomsadze, A., Burns, F. D. & Borodovsky, M. Integration of mapped RNA-Seq reads into automatic training of eukaryotic gene finding algorithm. *Nucleic Acids Res.* **42**, e119 (2014).
115. Pertea, M. et al. StringTie enables improved reconstruction of a transcriptome from RNA-seq reads. *Nat. Biotechnol.* **33**, 290–295 (2015).
116. Trapnell, C. et al. Transcript assembly and quantification by RNA-Seq reveals unannotated transcripts and isoform switching during cell differentiation. *Nat. Biotechnol.* **28**, 511–515 (2010).
117. Haas, B. J. et al. Automated eukaryotic gene structure annotation using EVidenceModeler and the program to assemble spliced alignments. *Genome Biol.* **9**, R7 (2008).
118. Nawrocki, E. P., Kolbe, D. L. & Eddy, S. R. Infernal 1.0: inference of RNA alignments. *Bioinformatics* **25**, 1335–1337 (2009).
119. Griffiths-Jones, S., Bateman, A., Marshall, M., Khanna, A. & Eddy, S. R. Rfam: an RNA family database. *Nucleic Acids Res.* **31**, 439–441 (2003).
120. Wucher, N. et al. FEELnc: A tool for Long non-coding RNAs annotation and its application to the dog transcriptome. bioRxiv https://doi.org/10.1101/064436 (2016).
121. Thiel, T., Michealek, W., Varshney, R. K. & Graner, A. Exploiting EST databases for the development and characterization of gene-derived SSR-markers in barley (*Hordeum vulgare* L.). *Theor. Appl. Genet.* **106**, 411–422 (2003).
122. Rice, P., Longden, I. & Bleasby, A. EMBOSS: The European molecular biology open software suite. *Trends. Genet.* **16**, 276–277 (2000).
123. Jurka, J. W. RepBase. https://www.girinst.org/server/RepBase (2016).
124. Smit, A. F. A. & Hubley, R. RepeatModeler Open-1.0. http://www.repeatmasker.org (2014).
125. Price, A. L., Jones, N. C. & Pevzner, P. A. D. novo identification of repeat families in large genomes. *Bioinformatics* **21**, i351–i358 (2005).

126. Benson, G. Tandem repeats finder: A program to analyze DNA sequences. *Nucleic Acids Res.* **27**, 573–580 (1999).

127. Jones, P. et al. InterProScan 5: genome-scale protein function classification. *Bioinformatics* **30**, 1236–1240 (2014).

128. Huerta-Cepas, J. et al. eggNOG 4.5: a hierarchical orthology framework with improved functional annotations for eukaryotic, prokaryotic and viral sequences. *Nucleic Acids Res.* **44**, D286–D293 (2016).

129. Rawlings, N. D., Barrett, A. J. & Finn, R. Twenty years of the MEROPS database of proteolytic enzymes, their substrates and inhibitors. *Nucleic Acids Res.* **44**, D343–D350 (2016).

130. Yin, Y. et al. dbCAN: A web resource for automated carbohydrate-active enzyme annotation. *Nucleic Acids Res.* **40**, W445–W451 (2012).

131. Petersen, T. N., Brunak, S., von Heijne, G. & Nielsen, H. SignalP 4.0: discriminating signal peptides from transmembrane regions. *Nat. Methods* **8**, 785–786 (2011).

132. Oktomirkochnikov, K., Conesa, A. & García-Alcalde, F. Qualimap 2: advanced multi-sample quality control for high-throughput sequencing data. *Bioinformatics* **32**, 292–294 (2016).

133. Sterne-Weiler, T., Peclet-Peltier, A. J., Best, A. J., Ka. C. H. & Blencowe, B. J. Efficient and accurate quantitative profiling of alternative splicing patterns of any complexity on a laptop. *Mol. Cell. Biol.* **72**, 187–190 (2018).

134. Aletra, A., Rahnenführer, J. & Lengauer, T. Improved scoring of functional groups from gene expression data by decorrelating GO graph structure. *Bioinformatics* **22**, 1600–1607 (2006).

135. Li, Y., Xiang, J. & Duan, C. Insulin-like growth factor-binding protein-3 plays an important role in regulating pharyngeal skeleton and inner ear formation and differentiation. *J. Biol. Chem.* **280**, 3613–3620 (2005).

136. Lin, J. M. et al. Actions of fibroblast growth factor-8 in bone cells in vitro. *Am. J. Physiol. Endocrinol. Metab.* **297**, E142–E150 (2009).

137. Nichols, J. T., Pan, L., Moens, C. B. & Kimmel, C. B. barx1 represses joints and promotes cartilage in the craniofacial skeleton. *Development* **140**, 2765–2775 (2013).

138. Bush, J. O., Lan, Y. & Jiang, R. The cleft lip and palate defects in Dancer mutant mice result from gain of function of the Tbx1 gene. *Proc. Natl. Acad. Sci. U. S. A.* **101**, 7022–7027 (2004).

139. Vieira, A. R. et al. Medical sequencing of candidate genes for nonsyndromic cleft lip and palate. *PLoS Genet.* **1**, e64 (2005).

140. Pappago, V. E. The T-box gene family: Emerging roles in development, stem cells and cancer. *Development* **141**, 3819–3833 (2014).

141. Kang, Y. J., Stevenson, A. K., Yau, P. M. & Kollmar, R. Sparc protein is required for normal growth of zebrafish otoliths. *Hum. Mutat.* **30**, 525–530 (2009).

142. Zhang, Z., Huynh, T. & Baldini, A. Mesodermal expression of Tbx1 is necessary and sufficient for pharyngeal arch and cardiac outflow tract development. *Development* **133**, 3587–3595 (2006).

143. Zarelli, V. E. & Dawid, I. B. Inhibition of neural crest formation by Kctd15 involves regulation of transcription factor AP-2. *Hum. Mutat.* **30**, 666–675 (2009).

144. Sterne-Weiler, T., Weatheritt, R. J., Best, A. J., Ha, K. C. H. & Blencowe, B. J. Efficient and accurate quantitative profiling of alternative splicing patterns of any complexity on a laptop. *Mol. Cell. Biol.* **72**, 187–190 (2018).

145. Lok, Y. T., Pan, L., Moens, C. B. & Kimmel, C. B. Translating growth factor-beta regulates basal transcriptional regulatory machinery to control cell proliferation and differentiation in cranial neural crest-derived osteoprogenitor cells. *J. Biol. Chem.* **285**, 4975–4982 (2010).

146. Zarelli, V. E. & Dawid, I. B. Inhibition of neural crest formation by Kctd15 involves regulation of transcription factor AP-2. *Hum. Mutat.* **30**, 525–530 (2009).

147. Thomason, M. & Bradshaw, A. D. SPARC osteoectenin in mineralized tissue. *Matrix Biol.* **52–54**, 78–87 (2016).

148. Rossel, E. M. & Bradshaw, A. D. SPARC osteoectenin in mineralized tissue. *Matrix Biol.* **52–54**, 78–87 (2016).

149. Selmar, J. E. & Dawid, I. B. Inhibition of neural crest formation by Kctd15 involves regulation of transcription factor AP-2. *Proc. Natl. Acad. Sci. U. S. A.* **110**, 2870–2875 (2013).

150. Zhang, Z., Huynh, T. & Baldini, A. Mesodermal expression of Tbx1 is necessary and sufficient for pharyngeal arch and cardiac outflow tract development. *Development* **133**, 3587–3595 (2006).

151. Zarelli, V. E. & Dawid, I. B. Inhibition of neural crest formation by Kctd15 involves regulation of transcription factor AP-2. *Hum. Mutat.* **30**, 525–530 (2009).

152. Plaster, N., Sonntag, C., Schilling, T. F. & Hammerschmidt, M. R. E. A. T. B. 2 interacts with histone deacetylase and FGF8 signaling to regulate multiple processes of zebrafish development. *Dev. Dyn.* **236**, 1891–1904 (2007).

153. Jordan, V. K. et al. Genotype–phenotype correlations in individuals with pathogenic RER variants. *Hum. Mutat.* **39**, 666–675 (2018).

154. Diegeseen, E. T., Kim, F. D. & Salzburger, W. Sequence analyses of the distal-less homeobox gene family in East African cichlids fishes reveal signatures of positive selection. * BMC Evol. Biol.* **13**, 153 (2013).

155. Stock, D. W. et al. The evolution of the vertebrate Dlx gene family. *Proc. Natl. Acad. Sci. USA* **93**, 10858–10863 (1996).

156. Mark, M., Ghyselincx, N. B. & Chambon, P. Function of retinoic acid receptors during embryonic development. *Nucl. Recept. Signal.* **7**, e002 (2009).

157. Linville, A., Radlke, K., Waxman, J. S., Yelon, D. & Schilling, T. F. Combinatorial roles for zebrafish retinoic acid receptors in the hindbrain, limbs and pharyngeal arches. *Dev. Biol.* **325**, 60–70 (2009).

158. Swartz, M. E., Sheehan-Rooney, K., Dixon, M. J. & Eberhart, J. K. Examination of a patogenic gene program in Zebrafish. *Dev. Dyn.* **240**, 2204–2220 (2011).

159. Iwata, J. et al. Transforming growth factor-beta regulates basal transcriptional regulatory machinery to control cell proliferation and differentiation in cranial neural crest-derived osteoprogenitor cells. *J. Biol. Chem.* **285**, 4975–4982 (2010).

160. Prochazkova, M., Prochazka, J., Marrangoni, P. & Klein, O. D. Bones, Glands, Ears and More: The Multiple Roles of FGF10 in Craniofacial Development. *Front. Genet.* **9**, 542 (2018).

161. Du, J. et al. Different expression patterns of Gli1–3 in mouse embryonic maxillofacial development. *Acta Histochem.* **114**, 620–625 (2012).

Acknowledgements
We thank Viola Nolte for expert technical support during sequencing, and Wolfgang Gessl for fish keeping and photographs.

Author contributions
Conceived the study: C.S. Designed the experiments: C.S., G.G.T., C.G. and C.Sch. Performed the experiments: S.K., C.B., G.M. and S.T. Analyzed the data: C.F. Wrote the paper: C.F., G.G.T., C.S. Contributed to the manuscript: S.K., C.G., C.Sch. Approved the final manuscript: all authors.

Funding
This work was supported by the Austrian Science Fund projects [FWF Grants P22737 and P29838]. The funders had no role in study design, data collection and analysis, decision to publish, or preparation of the manuscript. The purchase of the Pacific Biosciences RS II instrument at the University of Lausanne was financed in part by the Loterie Romande through the Fondation pour la Recherche en Médecine Génétique.
Competing interests
The authors declare no competing interests.

Additional information
Supplementary Information The online version contains supplementary material available at https://doi.org/10.1038/s41598-021-81030-z.

Correspondence and requests for materials should be addressed to G.G.T. or C.S.

Reprints and permissions information is available at www.nature.com/reprints.

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2021