

Differential use of multiple genetic sex determination systems in divergent ecomorphs of an African crater lake cichlid

Munby, Hannah; Linderoth, Tyler; Fischer, Bettina; Du, Mingliu; Vernaz, Grégoire; Tyers, Alexandra M.; Ngatunga, Benjamin P.; Shechonge, Asilatu; Denise, Hubert; McCarthy, Shane A.; Bista, Iliana; Miska, Eric A.; Emilia Santos, M.; Genner, Martin J.; Turner, George F.; Durbin, Richard

10.1101/2021.08.05.455235

Published: 06/08/2021

Publisher's PDF, also known as Version of record

Cyswllt i'r cyhoeddiad / Link to publication

Dyfyniad o'r fersiwn a gyhoeddwyd / Citation for published version (APA): Munby, H., Linderoth, T., Fischer, B., Du, M., Vernaz, G., Tyers, A. M., Ngatunga, B. P., Shechonge, A., Denise, H., McCarthy, S. A., Bista, I., Miska, E. A., Emilia Santos, M., Genner, M. J., Turner, G. F., & Durbin, R. (2021). Differential use of multiple genetic sex determination systems in divergent ecomorphs of an African crater lake cichlid. (bioRxiv). BioRxiv. https://doi.org/10.1101/2021.08.05.455235

Hawliau Cyffredinol / General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- · Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
 - You may not further distribute the material or use it for any profit-making activity or commercial gain
 You may freely distribute the URL identifying the publication in the public portal?

Take down policy
If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

1 Differential use of multiple genetic sex determination systems in divergent ecomorphs of

2 an African crater lake cichlid

- 3 Hannah Munby^{1,†}, Tyler Linderoth^{1,†,*}, Bettina Fischer¹, Mingliu Du^{1,2,5}, Grégoire Vernaz^{1,2,5}, Alexandra M.
- 4 Tyers³, Benjamin P. Ngatunga⁴, Asilatu Shechonge⁴, Hubert Denise¹, Shane A. McCarthy^{1,5}, Iliana
- 5 Bista^{1,2,5}, Eric A. Miska^{1,2,5}, M. Emília Santos⁶, Martin J. Genner⁷, George F. Turner³, Richard Durbin^{1,5,*}
- 6 ¹Department of Genetics, University of Cambridge, Cambridge, UK
- 7 ²Wellcome/CRUK Gurdon Institute, University of Cambridge, Cambridge, UK
- 8 ³School of Natural Sciences, Bangor University, Bangor, UK
- ⁹ ⁴Tanzania Fisheries Research Institute, Dar es Salaam, Tanzania
- ¹⁰ Wellcome Sanger Institute, Hinxton, Cambridge, UK
- 11 ⁶Department of Zoology, University of Cambridge, Cambridge, UK
- 12 ⁷School of Biological Sciences, University of Bristol, Bristol, UK
- 13 [†]Authors contributed equally to the work.
- 14 *Authors for correspondence: tl483@cam.ac.uk, rd109@cam.ac.uk

15 Abstract

- 16 African cichlid fishes not only exhibit remarkably high rates of speciation but also have some of
- 17 the fastest evolving sex determination systems in vertebrates. However, little is known
- 18 empirically in cichlids about the genetic mechanisms generating new sex-determining variants,
- 19 what forces dictate their fate, the demographic scales at which they evolve, and whether they
- are related to speciation. To address these questions, we looked for sex-associated loci in full
- 21 genome data from 647 individuals of Astatotilapia calliptera from Lake Masoko, a small isolated
- 22 crater lake in Tanzania, which contains two distinct ecomorphs of the species. We identified
- 23 three separate XY systems on recombining chromosomes. Two Y alleles derive from mutations
- 24 that increase expression of the gonadal soma-derived factor gene (qsdf) on chromosome 7; the
- 25 first is a tandem duplication of the entire gene observed throughout much of the Lake Malawi
- 26 haplochromine cichlid radiation to which A. calliptera belongs, and the second is a 5 kb insertion
- 27 directly upstream of *gsdf*. Both the latter variant and another 700 bp insertion on chromosome
- 28 19 responsible for the third Y allele arose from transposable element insertions. Males
- 29 belonging to the Masoko deep-water benthic ecomorph are determined exclusively by the gsdf

- 30 duplication, whereas all three Y alleles are used in the Masoko littoral ecomorph, in which they
- 31 appear to act antagonistically among males with different amounts of benthic admixture. This
- 32 antagonism in the face of ongoing admixture may be important for sustaining multifactorial sex
- 33 determination in Lake Masoko. In addition to identifying the molecular basis of three coexisting
- 34 sex determining alleles, these results demonstrate that genetic interactions between Y alleles
- 35 and genetic background can potentially affect fitness and adaptive evolution.

Introduction

- 37 Sex, as a means of generating beneficial combinations of alleles, is one of the most effective
- 38 evolutionary innovations used among eukaryotes to surmount fitness challenges. Many different
- 39 means of establishing separate sexes have arisen across the tree of life, operating through a
- 40 combination of genetic and environmental mechanisms (Bachtrog et al., 2014; Pennell et al.,
- 41 2018). The continued evolution of new sex determination systems can provide a means to
- 42 improve fitness via altering sex ratios (Kocher, 2004), resolving sexually antagonistic mutations
- 43 (van Doorn & Kirkpatrick, 2007; 2010), and avoiding the negative consequences of sex
- 44 chromosome degeneration (Blaser et al., 2013). Given this adaptive role of sex determination,
- 45 this begs the question of whether it is any coincidence that the fastest reported rates of sex
- 46 chromosome and heterogamety transitions among vertebrates (El Taher et al., 2020) have
- 47 occurred in East African cichlid fishes, renowned also for their extremely high speciation rates
- 48 (Brawand et al., 2014; Ronco et al., 2020). In support of such an association, population genetic
- 49 models have demonstrated how heterogamety switches arising from a new sex-determining
- 50 locus coupled with sexual and sex-ratio selection can help generate reproductive isolation in
- 51 sympatry (Lande *et al.*, 2001).
- 52 Sex-determination across African cichlid species is largely governed genetically in either a
- 53 single-locus or polygenic fashion (Ser et al., 2010). The loci controlling sex are known to exist
- 54 both on homomorphic sex chromosomes, for which there is little if any evidence for long range
- 55 suppression of recombination around the sex-determining alleles (Parnell & Streelman, 2013),
- 56 and on supernumerary B chromosomes (Clark et al., 2017; Clark & Kocher, 2019). Within the
- 57 Lake Malawi haplochromine cichlid radiation, the characterized sex determining loci are the
- 58 orange blotch associated ZW locus and an XY locus on chr5 (Roberts et al., 2009; Ser et al.,
- 59 2010), two XY loci on chr7 (Albertson, 2002; Parnell & Streelman, 2013; Roberts et al., 2009),
- 60 an XY locus on chr3, and a ZW locus on chr20 (Parnell & Streelman, 2013), using the

- chromosome numbering established for the *Metriaclima zebra* genome (Conte & Kocher, 2015).

 In most of these cases, multiple sex determination systems have been observed to act within a
- 63 single species. Most studies to date have identified sex-associated loci through
- 64 captive-breeding experiments (e.g. Parnell & Streelman, 2013; Ser et al., 2010), which provide
- 65 only broad genomic resolution, or through GWAS on relatively small sample sizes in wild
- 66 populations with limited power to detect intraspecific associations (El Taher et al., 2020). While
- 67 these studies point to cichlid sex determination as being highly fluid on the timescale of
- 68 hundreds of thousands to millions of years, studies on the dynamics within populations would
- 69 provide the context for examining how recombination, selection, and drift interact with molecular
- 70 mechanisms to shape the evolution of nascent sex chromosomes (Furman et al., 2020). To this
- 71 end, we sought to understand how sex determination acts in a single population of the eastern
- 72 happy cichlid Astatotilapia calliptera.
- 73 Astatotilapia calliptera is found both in the shallow margins of Lake Malawi as well as in the
- 74 surrounding rivers and smaller lakes. Peterson et al. (2017) found that the major chr7 XY locus
- 75 previously identified in Malawi Mbuna cichlids determined sex in a population of *A. calliptera*
- 76 from Lake Malawi. Despite only mapping the effect to megabase-scale resolution, they
- 77 postulated that a variant in the gonadal soma-derived factor (gsdf) gene on chromosome 7 was
- 78 responsible for dictating sex given its repeated role in sex determination in other fish species
- 79 (Einfeldt et al., 2021; Jiang et al., 2016; Kaneko et al., 2015; Myosho et al., 2012).
- 80 In particular, we studied A. calliptera in crater Lake Masoko to the north of Lake Malawi, which is
- 81 estimated to have formed ~50,000 years ago (Williamson et al., 1999). Lake Masoko is only 700
- metres in diameter with a shallow littoral margin and walls steeply descending to around 36 m at
- 83 its deepest point (Turner et al., 2019). It is currently a closed system, without surface
- 84 connections to any other water bodies (Turner et al., 2019). With the only other fish being two
- 85 cichlid species distantly related to *A. calliptera* and one clariid catfish species, the lake provides
- 86 a relatively simple context for studying the evolutionary genetics of sex determination,
- 87 speciation and their potential interaction. Genomic evidence suggests that A. calliptera
- 88 colonised the shallow littoral habitat from nearby river systems ~10,000 years ago, and
- 89 subsequently extended its range into the deeper benthic habitat ~1,000 years ago (Malinsky et
- 90 al., 2015). These shallow littoral and deep benthic populations are phenotypically distinct
- 91 ecomorphs, with the differences in habitat use coinciding with differences in body shape and jaw
- 92 morphology. Moreover, the ecomorphs can be distinguished by differences in male breeding

- 93 colouration, with reproductively active littoral males being typically yellow, and benthic males
- 94 dark blue. Both ecomorphs are sexually dimorphic, with males generally larger and more
- 95 brightly coloured than the females, which tend to have a duller, silvery brown colouration.

96 Results

- 97 We collected whole genome shotgun sequencing data for 548 Astatotilapia calliptera from Lake
- 98 Masoko at a median coverage of 14.5x (range 4.5x 22x, mean of 12.2x), and combined this
- 99 with data from 99 previously published samples (Malinsky et al., 2015), resulting in whole
- 100 genome sequence data for 596 male and 51 female fish (Supplementary Table 1). Reads were
- 101 mapped to the high-quality fAstCal1.2 A. calliptera reference genome and variants called at
- 102 3,328,052 quality-screened single nucleotide polymorphism (SNP) sites (see Methods for
- 103 details).
- 104 Multiple Y alleles determine sex in Lake Masoko
- 105 We carried out a genome wide association study (GWAS) for sex using a linear mixed model
- 106 framework (Figure 1a). The most strongly associated SNP is very highly significant (log₁₀
- p-value = 2.02e-22), and located at position 18,098,212 on chromosome 7 approximately 8 kb
- 108 downstream of the gene gsdf. By considering read depth summed over all fish heterozygous for
- this SNP, we established that it, and the entire gsdf gene, are contained in a 20 kb-long region
- 110 that exhibits 50% inflated relative coverage in the heterozygotes, suggesting that the associated
- 111 variant chromosome contains a duplication of this region (Figure 1b). We examined paired end
- 112 Illumina reads from Masoko A. calliptera samples homozygous for the apparent duplication
- (Supplementary Figure 1a), and long Pacific Biosciences reads from a male fish from a related
- 114 species (*Tropheops* sp. 'mauve') which also shows the inflated coverage pattern
- 115 (Supplementary Figure 1b), and in both cases confirmed the presence of a tandem duplication
- spanning coordinates 18,079,155 to 18,100,834 of chr7. We also confirmed the presence of this
- duplication junction by PCR (Supplementary Figure 1c). Copy number of the duplication is a
- 118 stronger predictor of sex than the best associated SNP from the GWAS scan (Table 1),
- 119 suggesting that the duplication itself operates as a Y allele in an XY sex determination system.

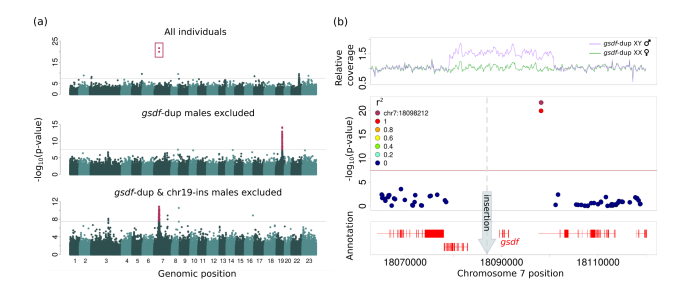


Figure 1: Genome-wide association study for sex. (a) P-values for the likelihood ratio test of an association between sex of Astatotliapia calliptera from Lake Masoko and their posterior mean genotypes at SNPs across the genome. The panels in order from top to bottom show results from the serial GWAS in which we looked for sex associations using all females and a subset of males not possessing the alternate allele of the single most highly-ranked SNP (or asdf-dup specifically for iterations two and three) from any of the previous GWAS. The grey. horizontal line in each of the Manhattan plots indicates the 0.05 Bonferroni-adjusted significance threshold, correcting for the number of tested SNPs. Significant SNPs tagging sex-determining loci are shown in maroon. (b) A zoomed-in view of the region harboring the SNPs most strongly associated with sex on chromosome 7. SNPs are coloured based on their degree of linkage disequilibrium with the most strongly sex-associated SNP tagging the qsdf duplication. The top panel shows the average sequencing depth in 100 bp bins of males heterozygous for the gsdf duplication compared to females. The sequencing depth of each individual was normalized with respect to their average depth in the non-duplicated flanking regions such that an increase of 0.5x in males compared to females indicates the presence of an extra copy of this locus. The duplication spans the region containing the entire qsdf gene and SNPs just downstream of qsdf were highly associated with sex in the GWAS run on all males and females. A 5 kb insertion upstream of gsdf indicated by the grey arrow characterizes the chr7-ins Y allele, which was in high linkage with the strongly sex-associated chromosome 7 SNPs in the bottom panel of (a).

122

123

124

125

127

128

129

130131

132

133134

135

136137

138

Table 1: Frequency of sex-determining genotypes in Lake Masoko *Astatotilapia calliptera* Multilocus genotypes for the sex determining loci are based on the number *qsdf* gene copies an

individual carries and their combination of reference (0) and insertion (1) alleles at the loci characterized by the chr19-ins and chr7-ins alleles. Among the 51 females in our sample, 46 were classified as low PC1 and five were middle PC1, none of which carried the *gsdf* duplication nor any of the insertion alleles.

gsdf copies	chr19-ins genotype	chr7-ins genotype	All males	Low PC1 males	Middle PC1 males	High PC1 males	Females
2	0/0	0/0	5	5	0	0	51
3	0/0	0/0	481	177	127	177	0
4	0/0	0/0	20	4	6	10	0
2	0/1	0/0	59	38	21	0	0
2	1/1	0/0	2	2	0	0	0
2	0/0	0/1	23	14	9	0	0
3	0/1	0/0	3	1	2	0	0
2	0/1	0/1	1	1	0	0	0
3	missing	0/0	2	1	0	1	0

The duplicated *gsdf* Y allele, which we call *gsdf*-dup, does not determine sex in all males: 90 of the 596 males (15%) are homozygous unduplicated, while 20 (3%) are apparently homozygous duplicated (2x relative sequence depth). To establish whether another locus might control sex in the males lacking *gsdf*-dup, we carried out a second sex GWAS with the 51 females and 90 males without the duplication. This revealed a region on chromosome 19 with multiple SNPs that were highly significant, the highest of which (position 21,581,905, log₁₀ p-value = 6.327883e-15) is located 77 bp upstream of the *e2f2* gene (Figure 1a). The inferred ancestral allele at this SNP was found exclusively among males across 59 heterozygotes and 3 homozygotes, suggesting a second XY system (Supplemental Table 2). We inspected the genomic region harboring variants in high linkage disequilibrium (LD) with the SNP to determine

155 whether it was tagging any other variants having an even stronger sex association not detected

by the GWAS, which was limited to biallelic SNPs. We discovered one such variant, a 700 bp insertion at position 21,572,413, which is located 1.7 kb upstream of the id3 gene (Supplementary Figure 2). This male-exclusive insertion, hereafter called chr19-ins, is found in 158 159 62 of the 90 males without *gsdf*-dup, of which 60 are heterozygotes and two are homozygotes. There are also three males with *qsdf*-dup that are heterozygous for chr19-ins. The additional sequence inserted in chr19-ins occurs in 37 places across 17 chromosomes and two unplaced scaffolds of the reference genome (blastn evalue = 0, > 96% identity, 100% coverage), and matches an LTR/Unknown family transposable element (blastn evalue = 0, 97% identity, 99% coverage) identified by repeatModeler2. At a more relaxed level of identity this transposable element is found in 126 places spread across all chromosomes and eight scaffolds of the reference genome (blastn evalue = 0, > 92% identity, 100% coverage). Since there remain 28 males carrying neither qsdf-dup nor chr19-ins, we repeated the GWAS procedure a third time, yielding another highly significant region of association on chromosome 7 around *gsdf* (Figure 1a). The most significant individual SNP in this case is approximately 371 kb upstream of gsdf (position 17,718,711, log_{10} p-value = 1.386670e-11), with a derived allele exclusively in males; 19 of the 28 males are heterozygous and one is homozygous (Supplemental Table 2). This pattern is consistent with a third Y allele that affects the asdf gene independently of the *qsdf* duplication. Further investigation in the window of elevated LD with this top GWAS SNP revealed a 5 kb insertion at position 18,086,980, hereafter called chr7-ins, located just 2.5 kb upstream of qsdf. This insertion is again exclusive to males including all with the chr7:17718711 derived allele as well as three additional males without any previously identified Y allele. Two subregions of the chr7-ins sequence, one 638 bp and the other 510 bp, are respectively found at 19 and 18 places throughout 15 chromosomes and three unplaced scaffolds of the A. calliptera reference genome (blastn evalue = 0, >90% identity, 100% coverage). RepeatModeler2 assigns them both to the ends of an unknown repeat family, 181 indicating that the chr7-ins insertion was also introduced by a transposable element. There remain 5 males (0.8% of 596) not carrying any of the three putative Y alleles (*qsdf*-dup, chr19-ins, chr7-ins). These results showing all genotypes are summarized in Table 1. It has been reported that B chromosomes can act dominantly to determine female sex in some rock-dwelling Mbuna Lake Malawi cichlids (Clark et al., 2017; 2018; 2019). We therefore 186 examined whether any of our Lake Masoko samples contained excess sequence indicative of B chromosomes, as defined in Clark *et al.* (2018). None of our samples showed any such excess, indicating that B chromosomes do not contribute to sex determination in this system.

9 Gsdf is expressed at higher levels in individuals carrying gsdf-affected Y alleles

Comparison of gene expression in the gonads of two adult male and two adult female *A*.

calliptera shows seven-fold higher *gsdf* expression in males than in females (Figure 2a),

consistent with observations in other fish species of higher levels of *gsdf* in testis than ovary

(Zhu et al., 2018). Furthermore, male carriers of *gsdf*-dup and chr7-ins, the latter which could

plausibly be in a promoter region of *gsdf* given its upstream proximity, express *gsdf* in

non-gonadal tissues (liver, eye, gill and anal fin) at substantially higher levels than males lacking

these alleles (Figure 2b & Supplementary Figure 3). Thus, we infer that higher *gsdf* expression

resulting from more copies of the actual gene itself or changes to a regulatory element triggers

masculinization in Masoko *A. calliptera*. In contrast, the inserted chr19-ins sequence upstream

of *id3*, the nearest gene to this insertion, did not show any associated changes in expression. It

remains unclear how this variant results in masculinization.

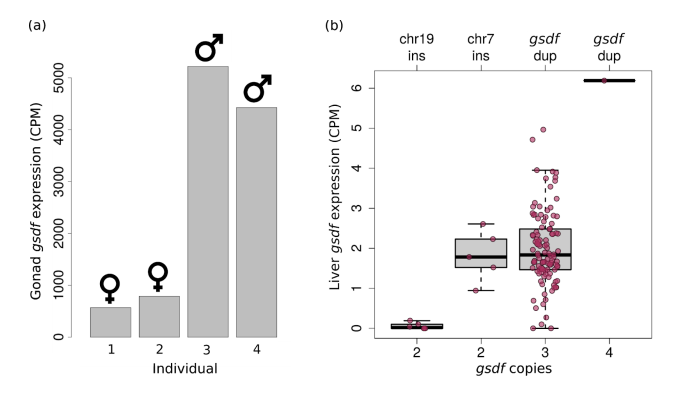


Figure 2: Expression of *gsdf*. (a) Expression levels of *gsdf* in the gonads of two male and two female *A. calliptera* reveals approximately seven times higher *gsdf* expression in males. (b)

203 Comparison of *gsdf* expression levels in the livers of Masoko male *A. calliptera* heterozygous 204 (three copies) and homozygous (four copies) for the *gsdf* duplication and males lacking the 205 duplication (two copies) but who carry Y alleles generated through insertions on chromosomes 206 7 and 19. The chromosome 7 insertion (chr7-ins) is directly upstream of *gsdf*, potentially in a 207 regulatory element of this gene. Thus, all males carrying Y alleles resulting from mutations 208 thought to affect *gsdf* express this gene more than other males on average. Gene expression 209 was quantified as counts per million reads (CPM).

210 Differential use of Y alleles in Lake Masoko

A principal component analysis (PCA) of the SNP data for the Lake Masoko samples reveals a 212 primary axis of genetic variation distinguishing the benthic from littoral ecomorph (Figure 3a). and this axis is strongly correlated with catch depth (Supplementary Figure 4). There is a tight 213 214 cluster of samples at high principal component 1 (PC1) corresponding to the benthic ecomorph. For the purposes of this paper we denote fish with PC1 > 0.4 as genetically benthic, and those 215 216 with PC1 < 0.4 as genetically littoral. The genetically littoral fish are more broadly distributed in the PCA plot, consistent with varying degrees of benthic admixture (Supplementary Figure 5), 217 218 and for some analyses below we partition them into a "low PC1" subgroup with PC1 < -0.02, 219 and a "middle PC1" group with -0.02 < PC1 < 0.4.

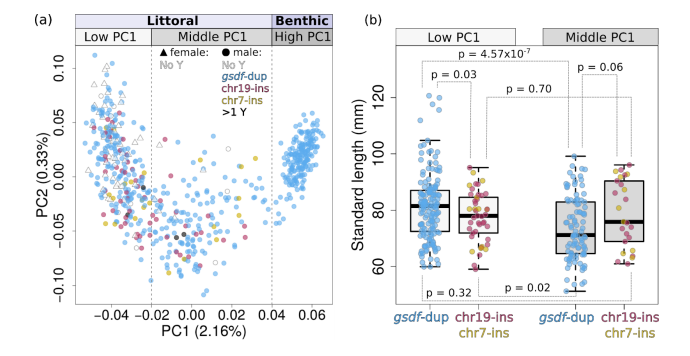


Figure 3: Genetic characterization of Masoko A. calliptera. (a) The first two components from a principal component analysis of the genome-wide variation among A. calliptera from Lake Masoko shows different Y allele usage between fish belonging to distinct genetic clusters. 223 The points represent individuals and their colours denote which of the sex determining alleles 224 identified from the GWAS individuals carry. PC1 separates fish adhering to the benthic ecomorph from littoral morph fish. The dashed grey lines show the demarcations that were used to classify fish as low, middle, and high PC1, which corresponds to their level of benthic 226 ancestry across the genome. (b) Comparisons between the standard lengths of littoral males heterozygous for *qsdf*-dup versus males heterozygous for chr19-ins or chr7-ins shows an 228 interaction between Y allele type and benthic admixture levels on body size. Males carrying 229 more than one type of Y allele were excluded. Two-tailed t-tests were used to test for significant 230 231 differences between the lengths of males characterized by different genetic PC1 background and Y allele combinations (p-values shown). 233 The genetically benthic fish were almost exclusively found in deep waters (> 20 metres), with just three of 188 individuals at intermediate depth (5-20 metres). The genetically littoral fish 234 were found predominantly at shallow (< 5 metres) and intermediate depths, though there were 235 236 some littoral fish caught in deep water, with a strong bias for these to be amongst fish with 237 higher PC1 values: in particular, amongst the 289 low PC1 subgroup individuals 138 were 238 caught shallow, 114 at intermediate depth, and 6 deep, while out of the 170 middle PC1 239 subgroup individuals 25 were caught shallow, 63 at intermediate depth, and 46 deep. Interestingly, all 188 genetically benthic males carried the gsdf duplication compared to 318/408 240 (78%) of the remaining males (Figure 3a); this deviates significantly from a null hypothesis in 242 which the frequency of males using *gsdf*-dup is independent of PC1 ($\chi^2 = 7.35$, p = 0.007). Correspondingly, the chr19-ins and chr7-ins alleles are only present in the genetically littoral males, at respective frequencies of 8.2% and 2.9%. Antagonism between Y alleles and admixture Fish grow throughout life, and there is evidence that physical size is a correlate of resource holding potential and reproductive success in males of African mouthbrooding cichlids (Hermann et al., 2015; Nelson, 1995; Sefc, 2011) where even a 1 mm size difference can 248 severely impact an individual's chances of winning bouts of male-male aggression (Turner & 249

250 Huntingford, 1986). In Lake Malawi haplochromine cichlids specifically, body size is a key predictor of the ability to successfully hold essential breeding territory from which to court females (Markert & Arnegard 2007). Even in the absence of male-male competition, at least in the case of South American convict cichlids, females prefer to mate with larger males 253 254 (Dechaume-Moncharmont et al., 2011), thus there is substantial evidence to suggest that male 255 cichlids may commonly benefit from being larger. In Lake Masoko, the genetically littoral male fish tend to be smaller as their amount of benthic 256 ancestry increases (Supplementary Figure 6, Supplementary Table 3). This decrease in size 257 with greater benthic admixture is significantly influenced by the type of Y allele that a male 258 carries (ANOVA F = 3.66, p = 0.027, comparing a linear model with interaction between genetic 259 PC1 and Y allele to a model with no interaction term). Chr19-ins males and chr7-ins males are the same size in both low and middle PC1 subgroups (low PC1 two-tailed t = -0.40, p = 0.70, middle PC1 two-tailed t = -0.24, p = 0.81), and together their size remains stable regardless of the level of benthic ancestry (two-tailed t = 0.38, p = 0.7, Figure 3b). In contrast, gsdf-dup males 263 with middle PC1 genetic ancestry are significantly smaller than those with low PC1 ancestry (two-tailed t = 5.21, p = $4.57*10^{-7}$). This size difference for *qsdf*-dup males is so pronounced that 265 266 while they are significantly larger than males using the other two Y alleles on the low PC1 267 background (two-tailed t = 2.24, p = 0.03) they tend to be smaller in an intermediate PC1 268 background. In contrast, the qsdf-dup genetically benthic (high PC1) males do not suffer from the size deficit seen in qsdf-dup middle PC1 males (Supplementary Figure 7a). Males 269 270 homozygous for *qsdf*-dup are on average 81 mm long, which is no different than heterozygotes (two-tailed t = -0.48, p = 0.64), and so by this proxy are equally fit. 271 Because PC1, which reflects benthic genetic content, is correlated with fish capture depth, we examined whether there could be an interaction between environment and genotype contributing to these size differences. Interestingly, while the gsdf-dup males with middle PC1 274 275 ancestry are smaller at all catch depths, chr19-ins and chr7-ins males with middle PC1 backgrounds are noticeably larger at depths greater than five metres (Supplementary Figure 276 277 7a). This larger size of the deeper-caught chr19-ins and chr7-ins middle PC1 males is counteracted by their shallow-caught counterparts tending to be the overall smallest, contributing to these males appearing similar in size across genetic backgrounds when not 279 280 accounting for depth. Despite numbers of some categories being low, this three-way interaction 281 between the depth at which fish are caught, Y allele type, and level of benthic ancestry, is

borderline significant in its ability to predict fish length (ANOVA F = 3.02, p = 0.05), suggesting that depth is relevant in contextualizing how different genetic combinations relate to body size, 284 and therefore fitness. 285 If the low PC1 and middle PC1 fish were sufficiently separated from each other genetically, these differences in size would be expected to lead to differences in the fraction of littoral males 287 carrying the rarer insertion alleles at greater depth or PC1 values. However, a three-way interaction between PC1 (restricted to low and middle PC1), catch-depth, and Y allele type is not significant in modeling the frequency of males ($\chi^2_2 = 0.08$, p = 0.96), nor are interactions 289 between Y allele type and depth or PC1 (Wald test z = -0.85 to 1.16, all p-values > 0.25 in the homogeneous association model of male frequency, which includes all pairwise interactions 291 between depth, Y allele and PC1) (Supplementary Figure 7b). Indeed, pooled across depths, gsdf-dup males are 3.5x more common than males carrying either of the other two Y alleles 293 among fish with low PC1 genetic backgrounds and 3.9x more common among middle PC1 294 males (difference not significant, Fisher's exact test p = 0.45). 295 296 Although the results of the last paragraph fail to provide direct evidence of a selective benefit for the Y insertion alleles at deeper depths or highly admixed genetic backgrounds in terms of allele frequency differences, it is noteworthy that elevated linkage disequilibrium (LD) extends for hundreds to thousands of kilobases from the strongest sex-associated GWAS SNPs tagging chr19-ins and chr7-ins (Supplementary Figure 2). To quantify this extent of LD we measured the mean squared physical distance between the chr19-ins and chr7-ins tagging SNPs and other 301 SNPs that were within a megabase and in strong LD ($r^2 > 0.5$) with these focal SNPs; these 302 values are in the 81st and 87th percentiles respectively compared to other randomly-sampled focal SNPs across the genome with the same allele frequencies. This is consistent with 305 long-range LD generated by recent positive selection, suggesting that either the sex-determining variants or another locus that they are physically linked to could be the target of selection. 307 308 Distribution of sex-determining alleles across the Lake Malawi cichlid radiation 309 We next investigated the presence of these Y alleles in other species from the Malawi radiation 310 for which we have sequenced samples. The *gsdf* duplication is seen in 100 additional species, suggesting that it is old and may correspond to the major male-determining allele in the chr7 XY

system observed to act previously in multiple Lake Malawi cichlid species (Parnell & Streelman, 2013; Ser *et al.*, 2010) (Supplementary Table 5). However, its use in sex determination appears to be quite dynamic; for example, it was not seen in the entire sample of 32 *A. calliptera* males from crater lake Itamba near to Lake Masoko (Figure 4a), and it has been lost or gained multiple times within the *Maylandia* genus (Figure 4b).

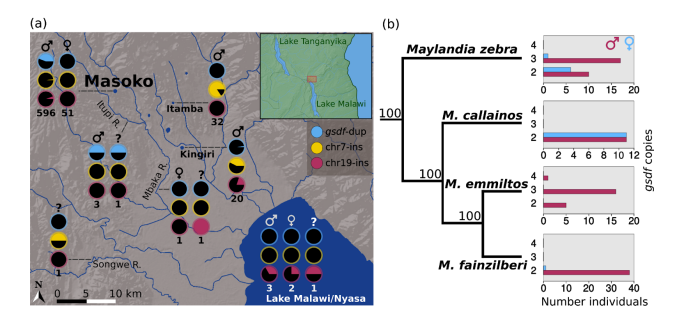


Figure 4: Geographic and taxonomic distribution of Y alleles. (a) The frequency of the gsdf-dup, chr7-ins, and chr19-ins alleles among A. calliptera males, females, and individuals of unknown sex sampled from lakes and rivers throughout Tanzania and Malawi suggests varied 320 usage of these alleles as sex determiners. The sample sizes for each sex and locality are indicated under pie charts of allele frequencies. (b) The frequency of male (blue) and female 321 322 (maroon) individuals from four *Maylandia* species that are either heterozygous (three copy), homozygous (four copy), or lacking (two copy) the duplicated qsdf allele exemplifies the 323 dynamic role of qsdf-dup in sex determination across the Malawi cichlid radiation. The presence 325 of the gsdf duplication in relation to the neighbor-joining species tree, rooted using the distantly-related outgroup Rhamphochromis longiceps, suggests that the qsdf duplication has 326 been lost or gained at least twice during the diversification of the Maylandia lineage. 327 Additionally, the *qsdf* duplication is found in both sexes of *M. zebra*, although at significantly 328 329 different frequencies (Fisher's exact test p = 0.035), consistent with it playing a role in sex determination in this population.

331 Among our specimens, the chr19-ins allele is exclusive to A. calliptera, and is geographically widespread, occurring in populations from another Tanzanian crater lake, Kingiri (Figure 4a), as well three other lakes, and five rivers (Supplementary Table 5) that span an area extending south and north of Lake Malawi. Among the 20 non-Masoko chr19-ins carriers for which we 334 335 have sex information, 18 were chr19-ins heterozygote males from the Bua River and lakes Kingiri, Malombe, Chilwa, and Malawi, and two were heterozygote females from the Salima 336 337 population of Lake Malawi and the Ruvuma River. 338 The chr7-ins allele occurs in other lake and riverine populations of A. calliptera mostly from the regions surrounding northern Lake Malawi except for one southern Lake Malawi population 339 (Southwest Arm). Among 20 Lake Kingiri males 55% are heterozygous for chr7-ins and 15% are 340 homozygous, while in 32 Lake Itamba males 31% are heterozygous and 69% are homozygous 341 (Figure 4a and Supplementary Table 5). The high frequency of chr7-ins homozygotes, 342 343 particularly in Itamba, suggests that this variant is either not sex determining or is being epistatically masked by a feminizing allele in these populations. We also detected the chr7-ins 344 variant in nine species from the genus *Tropheops* and two *Pseudotropheus* species 346 (Supplementary Table 6). Both genera are endemic to Lake Malawi and belong to the Mbuna 347 clade that is phylogenetically close to A. calliptera (Malinsky et al., 2018). Small sample sizes of 348 both males and females for these species and the coincidence of both the asdf-duplication and 349 chr7-ins make it difficult to confidently discern whether chr7-ins could be involved in sex determination, although there is an indication in some cases. For instance, in *Tropheops* sp. 350 351 'Chilumba' and *Tropheops* sp. 'mauve' there are males heterozygous for chr7-ins without a duplicated *qsdf*, however there are no females for comparison. Such a male is also found from 352 Tropheops sp. 'black' but in this species, and Tropheops sp. 'white dorsal', females occur that 354 carry both *gsdf*-dup and chr7-ins. While sexing errors could be responsible, a potentially more plausible explanation is the presence in *Tropheops* of a dominant female-determining variant at 355 another locus, given that females with either or both chr7-ins and *qsdf*-dup are observed 356 multiple times. Of the two *Pseudotropheus* species positive for chr7-ins, only one, 357 Pseudotropheus fuscus, had sexed individuals; 2/2 males are heterozygous for chr7-ins and have an unduplicated *qsdf*, while the only female lacks both *qsdf*-dup and chr7-ins, which is 359 consistent with chr7-ins being male-determining.

361 Discussion

362 Our genome-wide survey for genetic associations with sex revealed that there are three putative XY determination systems segregating within a single natural population of Astatotilapia calliptera from the crater lake Masoko. Among these, two are associated with qsdf on chromosome 7: the duplication present in 85% of males, which is the primary mechanism, and 365 366 an upstream insertion present in 4% of males. The third Y allele is characterized by an insertion on chromosome 19 in 11% of males. These systems are used differentially between the 367 divergent ecomorphs in the lake, with the deep-water benthic morph only using the duplication, 368 while littoral fish use all three systems. 369 370 Although use of multiple sex determination systems might seem likely to create sex-ratio biases, multiple Y alleles can coexist without problem in a population, with each male just carrying one of them, and females carrying none of them; Mendelian segregation in the offspring then gives 50% males with the paternal Y and 50% females. Indeed, we saw no females with any of the Y alleles. However in our larger set of males we did detect some that carried two Y alleles, including males homozygous for the *qsdf* duplication and others with two different Y alleles, 375 suggesting that there are some females carrying Y alleles present in the broader population. A 377 possible explanation for this is that a dominant ZW system may also be present at low 378 frequency, in which a dominant feminizing W allele acts epistatically to any of the Y alleles, as seen in some other Lake Malawi cichlid species (Parnell & Streelman, 2013; Ser et al., 2010). 380 We did not detect such a W allele in our association scans, possibly because the number of females in our data set did not give sufficient power to detect it at the frequency which would 381 382 explain our observations. Alternatively, there could be incomplete penetrance of the duplication allele, or genetically male fish could rarely undergo environmentally-induced sex reversal, which 383 has been documented in more taxonomically distant cichlids (Baroiller et al., 1995). 384 Complete genomic sequencing of many wild individuals enabled us to identify the likely causal 385 genetic mechanisms creating new Y alleles and corroborate the suspicion by Peterson et al. 386 387 (2017) that gsdf is a sex determination locus in A. calliptera. Our findings indicate that the tandem duplication of *qsdf* and the proximal upstream insertion both boost *qsdf* expression, 388 389 consistent with leading to masculinization as shown in *Oryzias* (Myosho et al., 2012). Upregulated *gsdf* expression appears to be generally important for testicular development in fish (Matsuda & Sakaizumi, 2016) and qsdf has been reported as a sex determiner in multiple fish 391 species (Einfeldt et al., 2021; Jiang et al., 2016; Kaneko et al., 2015; Myosho et al., 2012). 392 Recycling of this gene for sex determination through repeated distinct mutations is evidence for

394 evolutionary conservation of the genetic pathways controlling sex even as the specific sex determining alleles turn over (see Bachtrog et al. 2014 and Vicoso 2019 for discussion on this topic). The second gene we identified, id3, has not previously been directly associated with sex 397 determination, and while we believe we have identified the responsible mutation we cannot be 398 certain of the affected gene. 399 The genetic mechanisms generating the Masoko Y alleles parallel those involved in the origin of the dmy/dmrt1bY male determining gene in Oryzias latipes, which arose from a duplication of 400 dmrt1. Two transposable elements (TEs) introduced transcription factor binding sites upstream 401 of the *dmrt1b* paralog, which altered its expression leading to it becoming the master 402 sex-determining gene (Herpin et al., 2010; Schartl et al., 2018). Similarly, both the chr19-ins and 403 chr7-ins Y alleles were created by TE insertions directly upstream of the id3 and gsdf genes 404 respectively, offering support for the notion that TEs may play a potent role in rewiring the 405 406 expression of genes to function as sex determiners (Dechaud et al., 2019). Usage partitioning among three different Y alleles within a single, isolated population provides a 407 408 striking example of how dynamic sex determination is in African cichlids. This complements 409 recent work showing that across the Lake Tanganyika cichlid radiation sex systems turn over at a higher rate than previously established for vertebrates (El Taher et al., 2020). Previous studies 410 411 showed that multiple sex determination systems can segregate within captive families involving crosses between Lake Malawi species (Parnell & Streelman, 2013; Ser et al., 2010), but did not characterize their distributions within natural populations. Our results from Lake Masoko allow us to explore how multiple co-occurring sex systems segregate in the wild, and their relationship to subpopulation structure. 416 All of the variants that we identified for controlling sex also exist outside of Lake Masoko. The presence of gsdf-dup across all major clades of the Lake Malawi radiation, except for 417 Diplotaxodon and Rhamphochromis, suggests that it either predated the radiation or arose early 418 in it. Despite this, the *qsdf* duplication has not fixed, instead showing evidence of gains and loss at fine taxonomic scales within genera and even species. In contrast, chr19-ins and chr7-ins are 420 both far more taxonomically constrained, with chr19-ins exclusive to A. calliptera, despite being 421 widespread geographically. This suggests that these variants, although at low frequency, are 422 423 also old and in the case of chr7-ins could have been introduced into *Tropheops* and 424 Pseudotropheus through introgression. Another possibility is that chr7-ins, seen in 11/69 (~16%) 425 of the uniquely-classified Mbuna species (2/14 genera) in our dataset, could have arisen in a common ancestor of A. calliptera and Mbuna and remained as a minor sex-determining player in comparison to gsdf-dup, which we detected in ~72% of the Mbuna species (11/14 genera). This scenario would suggest that *qsdf*-dup may be selectively advantageous over chr7-ins in 428 429 most circumstances, while there are some conditions that favour chr7-ins. A common feature of all of the Y alleles we identified is that outside of Masoko they do not always appear to determine sex, suggesting that multifactorial sex determination is common and highly variable with respect to which alleles serve as the major sex determiners, even in closely related species. Having identified some of the precise variants influencing sex differentially across the 433 radiation enables future studies into the evolutionary factors supporting their turnover at a 434 variety of evolutionary scales. Our results raise the question of which eco-evolutionary contexts promote the invasion and eventual maintenance or loss of new sex determining variants. Theorized evolutionary mechanisms contributing to sex system turnover include resolving sexually antagonistic traits (van Doorn & Kirkpatrick, 2007), escape from deleterious mutational load (Blaser et al., 2013), 439 440 selection on sex ratios (Eshel, 1975), genetic drift (Saunders et al., 2018), and transmission distortion (Clark & Kocher, 2019; Werren & Beukeboom, 1998). In considering how our findings 441 align with such models it is important to recognize that we are only observing a snapshot of whatever dynamics may be occurring in Masoko, rather than seeing the evolutionary trajectories of Y allele usage. 445 Under the classic model of sexually antagonistic selection (van Doorn & Kirkpatrick, 2007), autosomal alleles with differential fitness effects between sexes gain an advantage if they become linked to a new sex determination locus, thus coupling the male-benefiting allele with males and vice versa. The resulting linkage disequilibrium can be reinforced in the long term through reduced recombination in the region containing the sex-determining and sexually antagonistic loci. When multiple sex loci co-occur in a population as in our case, the Y allele 450 conferring the greatest fitness advantage to males will spread. 451 452 We found evidence of an antagonistic relationship in terms of body size between the different Y alleles and genetic PC1 in littoral males. In cichlids, larger size confers higher fitness to males by providing them with an advantage in defending spawning sites and procuring access to 454 reproductively active females (Hermann et al., 2015). In the shallow waters where spawning

littoral fish have been observed, the frequencies of males characterized by different combinations of Y alleles and levels of benthic ancestry correlate well with their average size: gsdf-dup males with low benthic ancestry (low PC1) are largest and most common compared to 459 males that either carry the chr19-ins or chr7-ins Y alleles or have more benthic ancestry (middle 460 PC1). This suggests that in shallow water among males with low levels of benthic ancestry, gsdf-dup males have a fitness advantage over males that carry the rarer Y alleles. This size 461 advantage disappears however in fish with an increased benthic ancestry component, with 462 middle PC1 gsdf-dup males being smaller by nearly 8 mm on average. Furthermore, in waters 463 deeper than five metres, among the fish with middle PC1 ancestry, chr19 and chr7 insertion 464 males actually gain a size advantage over *qsdf*-dup males. These size differences are all 465 greater than the level known to be sufficient for preventing smaller males of another African 466 cichlid species from being able to effectively compete for territories (Turner & Huntingford, 467 1986). In A. calliptera specifically, body size has been shown to significantly influence 468 469 male-male aggression, presumably because it signals the resource holding potential of competing males (Theis et al., 2015). Therefore, we suggest that the insertion Y alleles may be maintained in the population by a relative advantage under these depth and genetic background 471 472 conditions, while there is sufficient genetic mixing between the low and middle PC1 subgroups 473 of littorals to prevent establishment of significant allele frequency differences. 474 We suggest two possible reasons, not mutually exclusive, for why the chr7-ins and chr19-ins Y alleles are not seen in the high PC1 benthic ecomorph. The first is that the PCA and admixture 476 plots (Figure 3a, Supplementary Figures 4, 5) are consistent with an asymmetry of gene flow between the benthic and littoral ecomorphs, with the benthic ecomorph that is adapted to the 477 cold, hypoxic environment at the bottom of the lake being genetically isolated with little if any gene flow from littorals into it, whereas there is gene flow from the benthics into littorals. This supports the cline of benthic admixture reflected in PC1 variation amongst the littorals. Second, 480 even if there is hybridisation leading to low levels of gene flow into benthics, there are reasons 481 to suggest it is sex-biased involving littoral females and benthic males. We never caught 482 genetically benthic fish in the shallow depths where littorals breed, but we do see occasional 483 484 genetic littorals in deep water. Benthic males appear to exclusively use the deep water mating territories that have been observed at the base of the crater wall, and we suggest that littoral 485 males may be unable to compete successfully in this forbidding environment to which they are 486 not adapted whereas littoral females may accept mating. In this scenario low frequency Y alleles 487 488 from the littorals would not invade the benthics at an appreciable rate, and any that were

present in the founders or entered through rare hybridization events could have been easily lost by drift.

In conclusion, our discovery that at least three different alleles control sex and segregate differentially within an isolated population of A. calliptera provides evidence that genetic sex determination in nature can be extremely fluid even at very small demographic scales. All of the 493 alleles we identified involved structural genetic variants, with two of the three generated by 494 transposable element insertions, highlighting a potentially important role for TEs in the rapidly evolving sex systems of African cichlids, similar to their role in adaptive variation in opsin 496 regulation (Carleton et al. 2020). Our results also indicate that genetic background differences 497 likely created by admixture can bring about antagonistic relationships among males carrying 498 different Y alleles, providing an evolutionary context that may favour multifactorial sex systems. 499 This has interesting implications for the incipient speciation between littoral and benthic Masoko 500 501 ecomorphs in that alternative Y alleles circumvent negative genetic interactions brought about by admixture, allowing for sustained back-crossing that reduces the level of divergence. It is 502 possible that this contributes to the low genome-wide F_{ST} (4%) between the ecomorphs, which 503 also lack fixed genetic differences, although there are tens of islands of high F_{ST} divergence 504 505 potentially associated with loci under differential selection (Malinsky et al., 2015). Admixture and relatively low divergence are hallmarks of the Malawi cichlid radiation, so it seems plausible that 507 similar processes could exist or have existed elsewhere. The fact that we and other studies 508 have found polygenic sex determination systems that differ markedly between closely related 509 species and populations across the radiation supports this possibility.

510 Methods

11 Samples and sequencing

Fish were primarily collected by professional aquarium fish catching teams. Fish at a target depth range (determined by diver depth gauges) were chased into block nets by SCUBA divers and transferred to a holding drum, then brought to the surface, where they were euthanized with clove oil. The right pectoral fin of sampled individuals was then removed and stored in ethanol, and the remainder of the specimen pinned, photographed, labelled and preserved in ethanol for later morphological analysis. Standard lengths were measured using calipers. Females were distinguished from juvenile males among the smaller fish by visual inspection of the gonads

519 after opening the abdominal cavity. Adult males were identified from secondary sexual traits of larger size, brighter colour and possession of elongate filaments on the pelvic, dorsal and anal 521 fins (confirmed to be reliable by visual inspection of the gonads in a number of specimens from 522 earlier collections). DNA was extracted from preserved fin clips using Qiasymphony DNA tissue extraction kits or 523 PureLink® Genomic DNA extraction kits and samples were sequenced on the Illumina HiSeg2000 as in Malinsky et al. (2015) or on the HiSegX in three batches: 1) 118 "ILBCDS" samples collected in 2011 sequenced at 3.9-19.2x coverage (median 7.5x), 2) 194 "CMASS" 526 samples collected in 2014-2016 sequenced to 4.3-9.0x coverage (median 5.7x), 3) 336 "cichl" 527 528 samples collected in 2014-2016 and 2018 sequenced to 12.0-23.2x coverage (median 15.8x). One sample that was initially part of the study was removed following conflicting data being 529 detected during the analysis. Further testing with our PCR assay of both the original tissue 530 sample obtained in the field, and a second sample from the supposed same ethanol-preserved, 531 532 whole specimen, produced one male and one female genotype respectively, indicating a labeling error (Supplementary Figure 1c). RNA was extracted from the gonads of two male and two female A. calliptera collected from the Itupi River in 2016. To ensure accurate quantification of transcripts, we used PolyA selection on 535 one male and one female sample and RNA depletion on the other male and female sample. The 536 gonads were then sequenced using 75 bp paired-end reads on three lanes of the Illumina HiSeq 537 2500 (SBS kit v4). Adapter sequences and bases with Phred quality below 20 were removed from the ends of gonad RNAseg reads using Trim Galore 0.6.2 539 (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) and read quality was checked 540 using FastQC 0.11.8 (https://www.bioinformatics.babraham.ac.uk/projects/fastqc/). We also extracted RNA from the anal fins, eyes, gills and livers of 151 A. calliptera collected from Lake Masoko in 2015, 2016 and 2018 (Supplementary Table 1), which was stored in RNALater, using 543 Direct-zol™ RNA MiniPrep Plus kits (Zymo, R2072) with an additional Chloroform step before loading the sample onto filtration columns. RNA samples were quantified with the Qubit™ RNA HS Assay Kit and quality assessed on the Agilent 4200 TapeStation. Libraries were prepared 547 using Illumina mRNA sequencing kits with polyA enrichment and sequenced using 100 or 150 548 bp paired-end reads on three lanes of the Illumina HiSeq4000 and five S4 lanes of the Illumina NovaSeg. Adapter sequences and bases with Phred quality below 20 were removed from the

550 ends of all resulting RNAseg reads using Trim Galore 0.6.4 and read quality was checked using FastQC 0.11.9. 552 Variant discovery Sequencing reads for all A. calliptera samples were mapped to a high-quality A. calliptera 553 reference genome (fAstCal1.2, accession GCA 900246225.3) (Rhie et al., 2021) using 554 bwa-mem 0.7.17 (Li, 2013). We used GATK 3.8 (McKenna et al., 2010) to identify individual-level variation with the HaplotypeCaller program followed by joint genotype calling 556 among all samples using GenotypeGVCFs (Poplin et al., 2017; Van der Auwera & O'Connor, 557 2020). Sites exhibiting any of the following indications of quality issues in the medium-coverage 558 (~15x) "cichl" subset of 336 individuals were masked from all analyses; total sequencing depth across individuals more extreme than the genome-wide median total site depth (DP) +/-25%, 560 fewer than 95% of individuals covered by at least five reads, root mean square mapping quality less than 40, an alternate allele assertion quality score below 30, excess heterozygosity (exact test p-value < 1e-4), biases between reference and alternate alleles in terms of strand (exact 563 564 test p-value < 1e-6), base quality (z-score > 6), mapping quality (z-score > 6), and read position 565 (z-score > 6). Sites spanning indels or having more than two alleles were also masked from 566 analyses. Quality control for sites was carried out using the program vcfCleaner 567 (https://github.com/tplinderoth/ngsQC/tree/master/vcfCleaner). 568 Population genetic characterization We used principal component analysis (PCA) based on genotype posterior probabilities at the quality-controlled SNPs to characterize the distribution of A. calliptera genetic variation throughout Lake Masoko. Specifically, we used ANGSD 0.929 (Korneliussen et al., 2014) to 571 estimate minor allele frequencies from genotype likelihoods (-GL 1 model) calculated using 572 reads with minimum base and map Phred qualities of at least 20. These minor allele frequency 573 (MAF) estimates and genotype likelihoods were used to obtain genotype posterior probabilities for all individuals under a Hardy-Weinberg genotype prior. We used ngsCovar 1.0.2 (Fumagalli et al., 2014) to estimate the genetic covariance matrix among individuals based on their genotype posteriors at SNPs with MAF greater than 5%, which we decomposed in R 3.6.3 (R 577 Core Team, 2020) with the eigen() function. In addition, we used the program ADMIXTURE

579 1.3.0 (Alexander et al., 2009) to infer the proportions of distinct genetic ancestry for individuals assuming two ancestral populations (K parameter). Genome-wide association tests for sex 581 For statistical association testing we relaxed the excess heterozygosity filter to accept biallelic 582 SNPs with exact test p-value > 1e-20, and queried all such SNPs across the genome with MAF of at least 5% for association with sex under the linear mixed model framework implemented in GEMMA 0.98.1 (Zhou & Stephens, 2012). Sex was treated as a binary response which we regressed against posterior mean genotypes calculated from the GATK genotype likelihoods 586 using vcf2bimbam (https://github.com/tplinderoth/ngsQC/tree/master/vcfCleaner) under a 587 Hardy-Weinberg genotype prior. We accounted for confounding effects of ancestry among 588 individuals through incorporating a centered pairwise kinship matrix calculated using GEMMA 589 as a random effect in the LMM. We identified significantly associated loci using the 590 likelihood-ratio test p-values from GEMMA run in the LMM mode at a 5% significance level after a Bonferroni correction for the number of tested SNPs. In order to identify as many sex-associated loci as possible, we iteratively tested conditional subsets of individuals who did not carry alleles significantly associated with sex from previous iterations, that is, subsets of 595 individuals whose sex was not accounted for by other candidates. 596 Characterizing sex-determining variants throughout Lake Masoko and the Malawi radiation We only used SNPs with GEMMA and so following the sex GWAS we checked for the presence of structural variants (SVs) that might have a stronger association with sex in 10 kb windows extending from the significantly associated SNPs. We extracted read mapping information directly from the BAM files to look for mapping signatures that would be consistent with 600 structural variation, considering both read pair and depth information, using IGV 2.8.0 (Robinson 601 et al., 2011). We initially screened at least five males and five females for structural variation in 602 IGV and then used a custom perl script to call SVs if at least 5% of read pairs among all individuals within 480 bp of any putative SV positions had mates which mapped to a different chromosome. We assembled the anomalously mapped read pairs across all individuals for each SV that we called using MEGAHIT 1.2.9 (Li et al., 2016) and performed a blastn (Altschul et al., 1990; Camacho et al., 2009) search of the resulting contigs against fAstCal1.2. This approach led to the discovery of the putative sex-determining insertions on chromosomes 7 and 19, which

609 blasted with at least 90% identity across their full length to multiple places across the genome. We used repeatModeler2 2.0.2 (Flynn et al., 2020) with default options but including the -LTRStruct option to identify transposable element sequences in the fAstCal1.2 genome. Then we compared the SV contigs to these transposable element sequences to further characterize the insertions. The chr19-ins allele matched a 700 bp transposable element (blastn evalue = 0, 97% identity, 99% coverage) identified by repeatModeler2 as belonging to an LTR/Unknown family. The two partial contigs of the chromosome 7 insertion matched with 94% identity (631/673 bp with 35/673 bp (5%) gaps) and 97% (496/509 bp with 11/509 bp (2%) gaps) to either end of a 3,947 bp unknown transposable element. 617 In order to characterize the presence or absence of the chromosome 7 and 19 insertions, we mapped sequencing reads from all Masoko A. calliptera to the assembled insertion sequences 619 including 1 kb of upstream and downstream flanking sequence using BWA. We considered any reads mapping within the flanking regions and which spanned the insertion as reference allele 621 reads (with respect to fAstCal1.2) and any reads which mapped within the insertion by a 623 minimum of three bp as alternate allele reads. An individual's genotype was called heterozygous (0/1) if they possessed reads from both alleles that were each at a minimum frequency of 10%, otherwise, with more than 90% of either the reference or insertion reads, individuals were called as homozygous for the reference allele (0/0) or homozygous for the 627 insertion allele (1/1), respectively. We also genotyped fish based on the copy number of the duplicated gsdf-containing locus which spans positions 18,079,155 to 18,100,834 of 628 629 chromosome 7 in the fAstCal1.2 reference. For each individual, we translated their average sequencing depth across this region relative to their average sequencing depth from 38,320 bp 630 flanking sequence (19,154 bp upstream and 19,166 bp downstream of the duplication breakpoints) into copy number in increments of 0.5x: Relative coverage of 1.25 or lower was recorded as a non-duplicated gsdf region, (1.25,1.75] as three gsdf copies, (1.75, 2.25] as four 633 copies, and so on. Individuals with three and four copies of the qsdf locus were called 634 heterozygous and homozygous for the duplication respectively. Though it is possible for a 635 four-copy individual to have one chromosome with three qsdf copies this would necessitate another duplication and so is less parsimonious than the assumption that they are homozygous 637 for a chromosome with two copies. We also developed a PCR assay for the *qsdf* duplication (Supplementary Table 7), which we 639 640 used to confirm its presence in a subset of A. calliptera and Maylandia zebra. Genomic DNA

641 was extracted from fin clips using PureLink Genomic DNA Mini Kits (ThermoFisher Scientific, K182001) following the manufacturer's protocols and eluted in 30-60 µL elution buffer. We carried out PCRs in 20 µL reaction volumes consisting of 1X Platinum™ II PCR Buffer, 0.2 mM of each dNTP (ThermoFisher Scientific, R0192), 0.2 µM of each primer (Merck Life Science, 644 desalted), less than 500 ng template DNA (1 µL genomic DNA at ~1-5 ng/µL), 0.04 U/µL Platinum™ II Tag Hot-Start DNA Polymerase (ThermoFisher Scientific, No 14966001) and nuclease-free water. We amplified the DNA using the following thermal profile: 94°C for two 647 minutes followed by 30-35 cycles of 94°C for 15 seconds, 60°C for 15 seconds, 68°C for 15 seconds, and a final 68°C extension for five minutes. The PCR products were separated using 649 electrophoresis run at 100 volts for 30 minutes on a 2% agarose gel. 650 We genotyped 1.552 additional individuals from all seven of the Lake Malawi radiation clades (A. calliptera, Mbuna, Benthic, Deep, Utaka, Diplotaxodon, and Rhamphochromis; see Malinsky et al. 2018) for the gsdf duplication as well as the chromosome 7 and 19 insertions in the same way as for Masoko A. calliptera described above. This set of Malawi radiation individuals 654 represents 270 species (some are not formally established but recognized as distinct taxa) from 655 656 48 genera, including A. calliptera from locations other than Lake Masoko. In order to 657 characterize how the gsdf duplication is acquired and lost as lineages diversify we mapped its 658 presence at different copy number in males and females to the species tree for four Mbuna 659 species from the Maylandia genus: M. zebra, M. callainos, M. emmiltos, and M. fainzilberi. We generated the species tree using 12,133,030 genome-wide segregating sites among the four 660 661 Maylandia species identified using GATK 3.8 in the same manner as for Masoko A. calliptera. These SNPs passed quality controls addressing abnormally low and high sequencing coverage 662 and low mapping quality for the ingroup samples as well as for samples from the distantly-related species Rhamphochromis longiceps, which served as an outgroup. We used 664 ngsDist 1.0.8 (Vieira et al., 2016) to calculate a pairwise genetic distance matrix based on 665 genotype likelihoods for all of the ingroup and outgroup samples, as well as to bootstrap sites in 666 order to generate 100 additional bootstrap distance matrices. For this *Maylandia* species tree, 667 we used fastME 2.1.6.1 (Lefort et al., 2015) to infer neighbor-joining trees from the genetic distance matrices using the BIONJ algorithm with SPR tree topology improvement. RAxML-NG 1.0.1 (Kozlov et al., 2019) was used to determine the bootstrap support for the genome-wide 671 tree.

672 B chromosome assay

In addition to autosomal sex loci, B chromosomes, which are supernumerary chromosomes not required for organismal function and variably present across taxa and individuals, have been 675 implicated as sex modifiers in Lake Malawi cichlids (Clark et al., 2017). Accordingly, we assayed for the presence of B chromosomes among Masoko A. calliptera to discern whether they may influence sex. B chromosome material initially derives from autosomes, so their presence can be detected through inflated read coverage in homologous regions of the reference genome where B reads mismap. Accordingly, we assayed for B chromosomes based on inflated coverage at regions containing sequence known to exist on B chromosomes from Lake Malawi 680 cichlids (Clark et al., 2018). Regions identified as core B block sequence according to Clark et 681 al. (2018) were translated into fAstCal1.2 coordinates and the mean coverage across each of these segments for each Masoko A. calliptera individual was calculated directly from the BAM files. We used a minimum coverage ratio for the core B region compared to the genome-wide average of 2x to call B positive individuals. None of the Lake Masoko A. calliptera passed this threshold although this process did identify individuals carrying B chromosomes from other 687 species.

688 Expression of sex-associated genes

We mapped the quality-controlled liver, eye, gill, and anal fin RNAseq reads to the fAstCal1.2 genome with STAR 2.7.3a (Dobin & Gingeras, 2015) and counted reads derived from sex-associated genes with featureCounts 2.0.1 (Liao *et al.*, 2014). These read counts were normalized to counts per million (CPM) reads using edgeR 3.30.3 (Robinson et al., 2010). We mapped the quality-controlled gonad reads to the fAstCal1.2 reference using bwa-mem and counted reads derived from *gsdf* exons using SAMtools 1.9 (Li *et al.*, 2009) and ngsAssociation 0.2.4 (https://github.com/tplinderoth/ngsAssociation) summarize, which were also normalized to CPM.

697 Relationship between Y alleles and body size

698 Genetic PC1 was used as a proxy for the degree of admixture since this component clearly
699 separates fish based on their degree of benthic ancestry. Based on distinct clustering in the
700 genome-wide PCA plot, fish with PC1 > 0.04 were classified as genetically benthic and those

701 with PC1 < 0.04 as genetically littoral. We further classified fish with the lowest amounts of benthic ancestry as "low PC1" (PC1 < -0.02), those with more equal amounts of littoral and benthic ancestry as "middle PC1" (PC1 range -0.02 to 0.04), and the clear benthic cluster as 704 "high PC1" (PC1 > 0.04). The three Y alleles segregate in the littoral group only, which is composed of low and middle PC1 fish, yielding six possible Y and PC1 combinations when excluding the 0.7% of males that carry more than one type of Y. For all analyses related to fish size we considered only males that were heterozygous for their Y allele (except when we compared the length of *qsdf*-dup homozygotes to *qsdf*-dup heterozygotes). We tested the hypothesis that littoral Lake Masoko A. calliptera males with different ancestry backgrounds and 709 710 Y allele combinations differ in standard length using pairwise two-tailed t-tests in R. We investigated whether the size of littoral males is influenced by interactions between Y allele and ancestry regime by fitting linear models of standard length as a function of Y allele and PC1 class in R using glm(). We tested whether the interaction provides a significantly better fit with the anova() F-test by comparing the residual sums of squares between a model with only main effects to a model with main effects and an interaction between Y allele type and PC1 class. We also introduced a depth class variable into our models to investigate whether the depth at which fish were caught plays a role in explaining their length. Depths less than five metres were considered "shallow", depths ranging from 5-20 metres were "intermediate", and depths more than 20 metres were "deep". As before, we compared the fit of a saturated model including the three-way interaction between Y allele, PC1 class, and depth band to the same model but 721 without the three-way interaction using analysis of variance to determine if the joint interaction between all variables provides a significant amount of additional power for predicting fish length. Since the size of male fish is likely to influence fitness, we used log-linear models to look at whether the same factors affecting length could predict the frequency of males. Specifically, we fit models using glm() in R with family='poisson' for the frequency of males based on Y allele, 725 PC1 class, and depth band. We assessed whether the frequency of males belonging to categories based on these three variables are independent of one another, and if not, what interactions were involved by performing an analysis of variance on nested pairs of models. We tested whether the differences in the residual deviance between the models being compared 729 were significant using χ^2 tests. This enabled us to find the simplest model that predicts male 730 frequencies statistically as well as the saturated model that includes all main effects and their possible interactions. The significance of terms within the context of a particular model for which

- 733 they were fit was determined using a Wald test of the null hypothesis that a term's effect is equal
- 734 to zero.
- 735 Assessment of linkage disequilibrium around sex loci
- 736 We calculated LD in terms of r² between each of the most highly sex-associated GWAS SNPs
- 737 and their surrounding SNPs using PLINK 1.9 (Purcell, 2014; Purcell et al., 2007). We observed
- 738 high LD, $r^2 > 0.5$, between the strongest GWAS SNPs tagging chr19-ins and chr7-ins and
- 739 far-ranging surrounding SNPs, which we visualized using plot_zoom
- 740 (https://github.com/hmunby/plot zoom). In order to determine how unusual these long stretches
- of high LD were, we compared the variance in the pairwise physical distance between the top
- GWAS SNPs and all SNPs within one megabase and $r^2 > 0.5$ to an expected distribution. The
- 743 background distributions were generated by randomly sampling 5,000 focal SNPs from across
- 744 the genome having the same alternate allele frequencies as each of the top GWAS SNPs. For
- 745 each sampled SNP, we calculated the variance among pairwise distances with other SNPs in
- 746 the same way as we had done for the GWAS SNPs.

747 Acknowledgments

- 748 We are grateful to African collaborators who assisted in sample collection, particularly the staff
- 749 of the Tanzanian Fisheries Research Institute, as well as Alan Hudson. We thank the
- 750 sequencing core staff at the Wellcome Sanger Institute. This work was supported by the
- 751 Wellcome Trust (WT207492 and WT206194). Additional support was to MJG & GFT
- 752 Leverhulme Trust Royal Society Africa Awards (AA100023 and AA130107); to MJG
- 753 Leverhulme Trust award (RF-2014-686); to GFT Leverhulme Trust award (RPG-2014-214); to
- 754 EAM Wellcome Trust Senior Investigator award (104640/Z/14/Z and 219475/Z/19/Z) and CRUK
- 755 award (C13474/A27826). GV thanks Wolfson College, University of Cambridge and the
- 756 Genetics Society, London for financial support.

757 Competing interests

758 The authors declare that they have no competing interests.

759 References

- 760 Albertson, R. (2002). Genetic basis of adaptive radiation in East African cichlids [Doctoral
- 761 Thesis, University of New Hampshire]. https://scholars.unh.edu/dissertation/98
- 762 Alexander, D. H., Novembre, J., & Lange, K. (2009). Fast model-based estimation of ancestry in
- unrelated individuals. *Genome Research*, 19(9), 1655–1664.
- 764 https://doi.org/10.1101/gr.094052.109
- Altschul, S. F., Gish, W., Miller, W., Myers, E. W., & Lipman, D. J. (1990). Basic local alignment
- search tool. *Journal of Molecular Biology*, *215*(3), 403–410.
- 767 https://doi.org/10.1016/S0022-2836(05)80360-2
- 768 Bachtrog, D., Mank, J. E., Peichel, C. L., Kirkpatrick, M., Otto, S. P., Ashman, T.-L., Hahn, M.
- 769 W., Kitano, J., Mayrose, I., Ming, R., Perrin, N., Ross, L., Valenzuela, N., Vamosi, J. C.,
- Tree of Sex Consortium. (2014). Sex determination: Why so many ways of doing it?
- 771 *PLoS Biology*, *12*(7), e1001899. https://doi.org/10.1371/journal.pbio.1001899
- 772 Baroiller, J. F., Chourrout, D., Fostier, A., & Jalabert, B. (1995). Temperature and sex
- chromosomes govern sex ratios of the mouthbrooding Cichlid fish *Oreochromis niloticus*.
- Journal of Experimental Zoology, 273(3), 216–223.
- 775 https://doi.org/10.1002/jez.1402730306
- 776 Bezault, E., Clota, F., Derivaz, M., Chevassus, B., & Baroiller, J.-F. (2007). Sex determination
- and temperature-induced sex differentiation in three natural populations of Nile tilapia
- (Oreochromis niloticus) adapted to extreme temperature conditions. Aquaculture, 272,
- 779 S3–S16. https://doi.org/10.1016/j.aquaculture.2007.07.227
- 780 Blaser, O., Grossen, C., Neuenschwander, S., & Perrin, N. (2013). Sex-chromosome turnovers
- induced by deleterious mutation load. *Evolution*, 67(3), 635–645.
- 782 https://doi.org/10.1111/j.1558-5646.2012.01810.x
- 783 Brawand, D., Wagner, C. E., Li, Y. I., Malinsky, M., Keller, I., Fan, S., Simakov, O., Ng, A. Y.,
- Lim, Z. W., Bezault, E., Turner-Maier, J., Johnson, J., Alcazar, R., Noh, H. J., Russell, P.,
- 785 Aken, B., Alföldi, J., Amemiya, C., Azzouzi, N., ... Di Palma, F. (2014). The genomic
- substrate for adaptive radiation in African cichlid fish. *Nature*, *513*(7518), 375–381.
- 787 https://doi.org/10.1038/nature13726
- 788 Camacho, C., Coulouris, G., Avagyan, V., Ma, N., Papadopoulos, J., Bealer, K., & Madden, T. L.
- 789 (2009). BLAST+: Architecture and applications. *BMC Bioinformatics*, 10(1), 421.
- 790 https://doi.org/10.1186/1471-2105-10-421
- 791 Carleton, K. L., Conte, M. A., Malinsky, M., Nandamuri, S. P., Sandkam, B. A., Meier, J. I.,

792 Mwaiko, S., Seehausen, O., & Kocher, T. D. (2020). Movement of transposable elements 793 contributes to cichlid diversity. *Molecular Ecology*, 29(24), 4956–4969. 794 https://doi.org/10.1111/mec.15685 Clark, F. E., Conte, M. A., Ferreira-Bravo, I. A., Poletto, A. B., Martins, C., & Kocher, T. D. 795 796 (2017). Dynamic Sequence Evolution of a Sex-Associated B Chromosome in Lake 797 Malawi Cichlid Fish. Journal of Heredity, 108(1), 53-62. 798 https://doi.org/10.1093/jhered/esw059 799 Clark, F. E., Conte, M. A., & Kocher, T. D. (2018). Genomic Characterization of a B 800 Chromosome in Lake Malawi Cichlid Fishes. Genes, 9(12). 801 https://doi.org/10.3390/genes9120610 802 Clark, F. E., & Kocher, T. D. (2019). Changing sex for selfish gain: B chromosomes of Lake 803 Malawi cichlid fish. Scientific Reports, 9(1), 20213. 804 https://doi.org/10.1038/s41598-019-55774-8 Conte, M. A., & Kocher, T. D. (2015). An improved genome reference for the African cichlid. 805 806 Metriaclima zebra. BMC Genomics, 16(1), 724. 807 https://doi.org/10.1186/s12864-015-1930-5 Dechaud, C., Volff, J.-N., Schartl, M., & Naville, M. (2019). Sex and the TEs: Transposable 808 809 elements in sexual development and function in animals. Mobile DNA, 10(1), 42. 810 https://doi.org/10.1186/s13100-019-0185-0 811 Dechaume-Moncharmont, F.-X., Cornuau, J. H., Keddar, I., Ihle, M., Motreuil, S., & Cézilly, F. 812 (2011). Rapid assessment of female preference for male size predicts subsequent 813 choice of spawning partner in a socially monogamous cichlid fish. *Comptes Rendus* 814 Biologies, 334(12), 906-910. https://doi.org/10.1016/j.crvi.2011.08.004 Dobin, A., & Gingeras, T. R. (2015). Mapping RNA-seg Reads with STAR. Current Protocols in 815 816 Bioinformatics, 51, 11.14.1-11.14.19. https://doi.org/10.1002/0471250953.bi11114s51 Einfeldt, A. L., Kess, T., Messmer, A., Duffy, S., Wringe, B. F., Fisher, J., den Heyer, C., 817 818 Bradbury, I. R., Ruzzante, D. E., & Bentzen, P. (2021). Chromosome level reference of 819 Atlantic halibut *Hippoglossus hippoglossus* provides insight into the evolution of sexual 820 determination systems. Molecular Ecology Resources, 1755-0998.13369. 821 https://doi.org/10.1111/1755-0998.13369 822 Eshel, I. (1975). Selection of sex-ratio and the evolution of sex-determination. *Heredity*, 34(3), 823 351-361. https://doi.org/10.1038/hdv.1975.44 El Taher, A. E., Ronco, F., Matschiner, M., Salzburger, W., & Böhne, A. (2020). Dynamics of sex 824

chromosome evolution in a rapid radiation of cichlid fishes [Preprint]. bioRxiv.

825

826 https://doi.org/10.1101/2020.10.23.335596 827 Flynn, J. M., Hubley, R., Goubert, C., Rosen, J., Clark, A. G., Feschotte, C., & Smit, A. F. 828 (2020). RepeatModeler2 for automated genomic discovery of transposable element 829 families. Proceedings of the National Academy of Sciences of the United States of 830 America, 117(17), 9451-9457. https://doi.org/10.1073/pnas.1921046117 831 Fumagalli, M., Vieira, F. G., Linderoth, T., & Nielsen, R. (2014). ngsTools: Methods for population genetics analyses from next-generation sequencing data. *Bioinformatics*, 832 833 30(10), 1486–1487. https://doi.org/10.1093/bioinformatics/btu041 834 Furman, B. L. S., Metzger, D. C. H., Darolti, I., Wright, A. E., Sandkam, B. A., Almeida, P., Shu, 835 J. J., & Mank, J. E. (2020). Sex Chromosome Evolution: So Many Exceptions to the Rules. Genome Biology and Evolution, 12(6), 750–763. 836 837 https://doi.org/10.1093/gbe/evaa081 Hermann, C. M., Brudermann, V., Zimmermann, H., Vollmann, J., & Sefc, K. M. (2015). Female 838 preferences for male traits and territory characteristics in the cichlid fish *Tropheus moorii*. 839 840 Hydrobiologia, 748(1), 61–74. https://doi.org/10.1007/s10750-014-1892-7 Herpin, A., Braasch, I., Kraeussling, M., Schmidt, C., Thoma, E. C., Nakamura, S., Tanaka, M., 841 & Schartl, M. (2010). Transcriptional rewiring of the sex determining dmrt1 gene 842 843 duplicate by transposable elements. *PLoS Genetics*, 6(2), e1000844. 844 https://doi.org/10.1371/journal.pgen.1000844 845 Holzberg, S. (1978). A field and laboratory study of the behaviour and ecology of 846 Pseudotropheus zebra (Boulenger), an endemic cichlid of Lake Malawi (Pisces; 847 Cichlidae). Journal of Zoological Systematics and Evolutionary Research, 16(3), 171–187. https://doi.org/10.1111/j.1439-0469.1978.tb00929.x 848 849 Jiang, D. N., Yang, H. H., Li, M. H., Shi, H. J., Zhang, X. B., & Wang, D. S. (2016). *qsdf* is a 850 downstream gene of dmrt1 that functions in the male sex determination pathway of the 851 Nile tilapia. Molecular Reproduction and Development, 83(6), 497–508. 852 https://doi.org/10.1002/mrd.22642 Kaneko, H., Ijiri, S., Kobayashi, T., Izumi, H., Kuramochi, Y., Wang, D.-S., Mizuno, S., & 853 854 Nagahama, Y. (2015). Gonadal soma-derived factor (gsdf), a TGF-beta superfamily gene, induces testis differentiation in the teleost fish Oreochromis niloticus. Molecular 855 856 and Cellular Endocrinology, 415, 87–99. https://doi.org/10.1016/j.mce.2015.08.008 Kocher, T. D. (2004). Adaptive evolution and explosive speciation: The cichlid fish model. *Nature* 857 858 Reviews. Genetics, 5(4), 288–298. https://doi.org/10.1038/nrg1316 859 Korneliussen, T. S., Albrechtsen, A., & Nielsen, R. (2014). ANGSD: Analysis of Next Generation

860 Sequencing Data. BMC Bioinformatics, 15, 356. 861 https://doi.org/10.1186/s12859-014-0356-4 Kozlov, A. M., Darriba, D., Flouri, T., Morel, B., & Stamatakis, A. (2019). RAXML-NG: A fast, scalable and user-friendly tool for maximum likelihood phylogenetic inference. 863 864 Bioinformatics, 35(21), 4453–4455. https://doi.org/10.1093/bioinformatics/btz305 865 Lande, R., Seehausen, O., & Alphen, J. J. M. van. (2001). Mechanisms of rapid sympatric speciation by sex reversal and sexual selection in cichlid fish. Genetica, 112/113, 866 867 435–443. https://doi.org/10.1023/A:1013379521338 868 Lefort, V., Desper, R., & Gascuel, O. (2015). FastME 2.0: A Comprehensive, Accurate, and Fast 869 Distance-Based Phylogeny Inference Program. Molecular Biology and Evolution, 32(10), 870 2798–2800. https://doi.org/10.1093/molbev/msv150 871 Li, D., Luo, R., Liu, C.-M., Leung, C.-M., Ting, H.-F., Sadakane, K., Yamashita, H., & Lam, T.-W. (2016). MEGAHIT v1.0: A fast and scalable metagenome assembler driven by advanced 872 873 methodologies and community practices. *Methods*, 102, 3–11. 874 https://doi.org/10.1016/j.ymeth.2016.02.020 875 Li, H., Handsaker, B., Wysoker, A., Fennell, T., Ruan, J., Homer, N., Marth, G., Abecasis, G., Durbin, R., & 1000 Genome Project Data Processing Subgroup. (2009). The Sequence 876 877 Alignment/Map format and SAMtools. *Bioinformatics*, 25(16), 2078–2079. 878 https://doi.org/10.1093/bioinformatics/btp352 879 Li, Heng. (2013). Aligning sequence reads, clone sequences and assembly contigs with 880 BWA-MEM. ArXiv:1303.3997 [g-Bio]. http://arxiv.org/abs/1303.3997 881 Liao, Y., Smyth, G. K., & Shi, W. (2014). featureCounts: An efficient general purpose program 882 for assigning sequence reads to genomic features. Bioinformatics, 30(7), 923–930. 883 https://doi.org/10.1093/bioinformatics/btt656 Malinsky, M., Challis, R. J., Tyers, A. M., Schiffels, S., Terai, Y., Ngatunga, B. P., Miska, E. A., 884 885 Durbin, R., Genner, M. J., & Turner, G. F. (2015). Genomic islands of speciation separate 886 cichlid ecomorphs in an East African crater lake. Science, 350(6267), 1493–1498. 887 https://doi.org/10.1126/science.aac9927 Malinsky, Milan, Svardal, H., Tyers, A. M., Miska, E. A., Genner, M. J., Turner, G. F., & Durbin, 888 889 R. (2018). Whole-genome sequences of Malawi cichlids reveal multiple radiations 890 interconnected by gene flow. Nature Ecology & Evolution, 2(12), 1940–1955. 891 https://doi.org/10.1038/s41559-018-0717-x

Markert, J. A., & Arnegard, M. E. (2007). Size-dependent use of territorial space by a

rock-dwelling cichlid fish. *Oecologia*, 154(3), 611–621.

892 893 894 https://doi.org/10.1007/s00442-007-0853-5 895 Matsuda, M., & Sakaizumi, M. (2016). Evolution of the sex-determining gene in the teleostean 896 genus Oryzias. General and Comparative Endocrinology, 239, 80–88. 897 https://doi.org/10.1016/j.ygcen.2015.10.004 898 McKenna, A., Hanna, M., Banks, E., Sivachenko, A., Cibulskis, K., Kernytsky, A., Garimella, K., 899 Altshuler, D., Gabriel, S., Daly, M., & DePristo, M. A. (2010). The Genome Analysis 900 Toolkit: A MapReduce framework for analyzing next-generation DNA sequencing data. 901 Genome Research, 20(9), 1297–1303. https://doi.org/10.1101/gr.107524.110 902 Myosho, T., Otake, H., Masuyama, H., Matsuda, M., Kuroki, Y., Fujiyama, A., Naruse, K., 903 Hamaguchi, S., & Sakaizumi, M. (2012). Tracing the Emergence of a Novel 904 Sex-Determining Gene in Medaka, Oryzias luzonensis. Genetics, 191(1), 163–170. 905 https://doi.org/10.1534/genetics.111.137497 Nelson, C.M. (1995). Male size, spawning pit size and female mate choice in a lekking cichlid 906 907 fish. Animal Behaviour, 50(6), 1587–1599. 908 https://doi.org/10.1016/0003-3472(95)80013-1 909 Parnell, N. F., & Streelman, J. T. (2013). Genetic interactions controlling sex and color establish 910 the potential for sexual conflict in Lake Malawi cichlid fishes. Heredity, 110(3), 239-246. https://doi.org/10.1038/hdy.2012.73 911 912 Pennell, M. W., Mank, J. E., & Peichel, C. L. (2018). Transitions in sex determination and sex 913 chromosomes across vertebrate species. *Molecular Ecology*, 27(19), 3950–3963. 914 https://doi.org/10.1111/mec.14540 915 Peterson, E. N., Cline, M. E., Moore, E. C., Roberts, N. B., & Roberts, R. B. (2017). Genetic sex 916 determination in Astatotilapia calliptera, a prototype species for the Lake Malawi cichlid 917 radiation. Die Naturwissenschaften, 104(5–6), 41. 918 https://doi.org/10.1007/s00114-017-1462-8 Poplin, R., Ruano-Rubio, V., DePristo, M. A., Fennell, T. J., Carneiro, M. O., Van der Auwera, G. 919 920 A., Kling, D. E., Gauthier, L. D., Levy-Moonshine, A., Roazen, D., Shakir, K., Thibault, J., 921 Chandran, S., Whelan, C., Lek, M., Gabriel, S., Daly, M. J., Neale, B., MacArthur, D. G., 922 & Banks, E. (2017). Scaling accurate genetic variant discovery to tens of thousands of 923 samples [Preprint], bioRxiv. https://doi.org/10.1101/201178 Purcell, S. (2014). PLINK 1.9. http://pnqu.mgh.harvard.edu/purcell/plink/ 925 Purcell, S., Neale, B., Todd-Brown, K., Thomas, L., Ferreira, M. A. R., Bender, D., Maller, J., 926 Sklar, P., de Bakker, P. I. W., Daly, M. J., & Sham, P. C. (2007). PLINK: A tool set for 927 whole-genome association and population-based linkage analyses. American Journal of 928 Human Genetics, 81(3), 559–575. https://doi.org/10.1086/519795 929 R Core Team. (2020). R: A Language and Environment for Statistical Computing. R Foundation 930 for Statistical Computing. https://www.R-project.org/ 931 Rhie, A., McCarthy, S. A., Fedrigo, O., Damas, J., Formenti, G., Koren, S., Uliano-Silva, M., 932 Chow, W., Fungtammasan, A., Kim, J., Lee, C., Ko, B. J., Chaisson, M., Gedman, G. L., 933 Cantin, L. J., Thibaud-Nissen, F., Haggerty, L., Bista, I., Smith, M., ... Jarvis, E. D. 934 (2021). Towards complete and error-free genome assemblies of all vertebrate species. 935 Nature, 592(7856), 737–746. https://doi.org/10.1038/s41586-021-03451-0 936 Roberts, R. B., Ser, J. R., & Kocher, T. D. (2009). Sexual Conflict Resolved by Invasion of a 937 Novel Sex Determiner in Lake Malawi Cichlid Fishes. Science, 326(5955), 998-1001. https://doi.org/10.1126/science.1174705 938 939 Robinson, J. T., Thorvaldsdóttir, H., Winckler, W., Guttman, M., Lander, E. S., Getz, G., & 940 Mesirov, J. P. (2011). Integrative genomics viewer. *Nature Biotechnology*, 29(1), 24–26. 941 https://doi.org/10.1038/nbt.1754 942 Robinson, M. D., McCarthy, D. J., & Smyth, G. K. (2010). edgeR: A Bioconductor package for 943 differential expression analysis of digital gene expression data. Bioinformatics, 26(1), 944 139–140. https://doi.org/10.1093/bioinformatics/btp616 945 Ronco, F., Büscher, H. H., Indermaur, A., & Salzburger, W. (2020). The taxonomic diversity of 946 the cichlid fish fauna of ancient Lake Tanganyika. East Africa, Journal of Great Lakes 947 Research, 46(5), 1067–1078. https://doi.org/10.1016/j.jglr.2019.05.009 948 Saunders, P. A., Neuenschwander, S., & Perrin, N. (2018). Sex chromosome turnovers and 949 genetic drift: A simulation study. *Journal of Evolutionary Biology*, 31(9), 1413–1419. 950 https://doi.org/10.1111/jeb.13336 951 Schartl, M., Schories, S., Wakamatsu, Y., Nagao, Y., Hashimoto, H., Bertin, C., Mourot, B., 952 Schmidt, C., Wilhelm, D., Centanin, L., Guiguen, Y., & Herpin, A. (2018). Sox5 is 953 involved in germ-cell regulation and sex determination in medaka following co-option of 954 nested transposable elements. BMC Biology, 16(1), 16. 955 https://doi.org/10.1186/s12915-018-0485-8 956 Sefc, K. M. (2011). Mating and Parental Care in Lake Tanganyika's Cichlids. International 957 Journal of Evolutionary Biology, 2011, 1–20. https://doi.org/10.4061/2011/470875 958 Ser, J. R., Roberts, R. B., & Kocher, T. D. (2010). Multiple interacting loci control sex 959 determination in Lake Malawi cichlid fish. *Evolution*, 64(2), 486–501. 960 https://doi.org/10.1111/j.1558-5646.2009.00871.x 961 Theis, A., Bosia, T., Roth, T., Salzburger, W., & Egger, B. (2015). Egg-spot pattern and body

962 size asymmetries influence male aggression in haplochromine cichlid fishes. Behavioral 963 Ecology, 26(6), 1512–1519. https://doi.org/10.1093/beheco/arv104 Turner, G. F., & Huntingford, F. A. (1986). A problem for game theory analysis: Assessment and 965 intention in male mouthbrooder contests. Animal Behaviour, 34(4), 961–970. 966 https://doi.org/10.1016/S0003-3472(86)80155-5 967 Turner, G., Ngatunga, B. P., & Genner, M. J. (2019). The Natural History of the Satellite Lakes of Lake Malawi [Preprint]. EcoEvoRxiv. https://doi.org/10.32942/osf.io/sehdq 968 969 Van der Auwera, G., & O'Connor, B. (2020). Genomics in the Cloud: Using Docker, GATK, and 970 WDL in Terra (1st ed.). O'Reilly Media. 971 van Doorn, G. S., & Kirkpatrick, M. (2007). Turnover of sex chromosomes induced by sexual 972 conflict. Nature, 449(7164), 909-912. https://doi.org/10.1038/nature06178 973 van Doorn, G. S., & Kirkpatrick, M. (2010). Transitions between male and female heterogamety caused by sex-antagonistic selection. *Genetics*, 186(2), 629–645. 974 975 https://doi.org/10.1534/genetics.110.118596 976 Vicoso, B. (2019). Molecular and evolutionary dynamics of animal sex-chromosome turnover. 977 Nature Ecology & Evolution, 3(12), 1632–1641. 978 https://doi.org/10.1038/s41559-019-1050-8 979 Vieira, F. G., Lassalle, F., Korneliussen, T. S., & Fumagalli, M. (2016). Improving the estimation 980 of genetic distances from Next-Generation Sequencing data: Genetic Distances from 981 NGS Data. Biological Journal of the Linnean Society, 117(1), 139–149. 982 https://doi.org/10.1111/bij.12511 983 Werren, J. H., & Beukeboom, L. W. (1998). Sex determination, sex ratios, and genetic conflict. 984 Annual Review of Ecology and Systematics, 29(1), 233–261. 985 https://doi.org/10.1146/annurev.ecolsys.29.1.233 Williamson, D., Jackson, M. J., Banerjee, S. K., Marvin, J., Merdaci, O., Thouveny, N., 986 987 Decobert, M., Gibert-Massault, E., Massault, M., Mazaudier, D., & Taieb, M. (1999). 988 Magnetic signatures of hydrological change in a tropical maar-lake (Lake Massoko, 989 Tanzania): Preliminary results. Physics and Chemistry of the Earth, Part A: Solid Earth 990 and Geodesy, 24(9), 799-803. https://doi.org/10.1016/S1464-1895(99)00117-9 991 Zhou, X., & Stephens, M. (2012), Genome-wide efficient mixed-model analysis for association 992 studies. Nature Genetics, 44(7), 821–824. https://doi.org/10.1038/ng.2310 993 Zhu, Y., Meng, L., Xu, W., Cui, Z., Zhang, N., Guo, H., Wang, N., Shao, C., & Chen, S. (2018). 994 The autosomal Gsdf gene plays a role in male gonad development in Chinese tongue 995 sole (Cynoglossus semilaevis). Scientific Reports, 8(1), 17716.

https://doi.org/10.1038/s41598-018-35553-7

997 Supplementary Figures & Tables

996

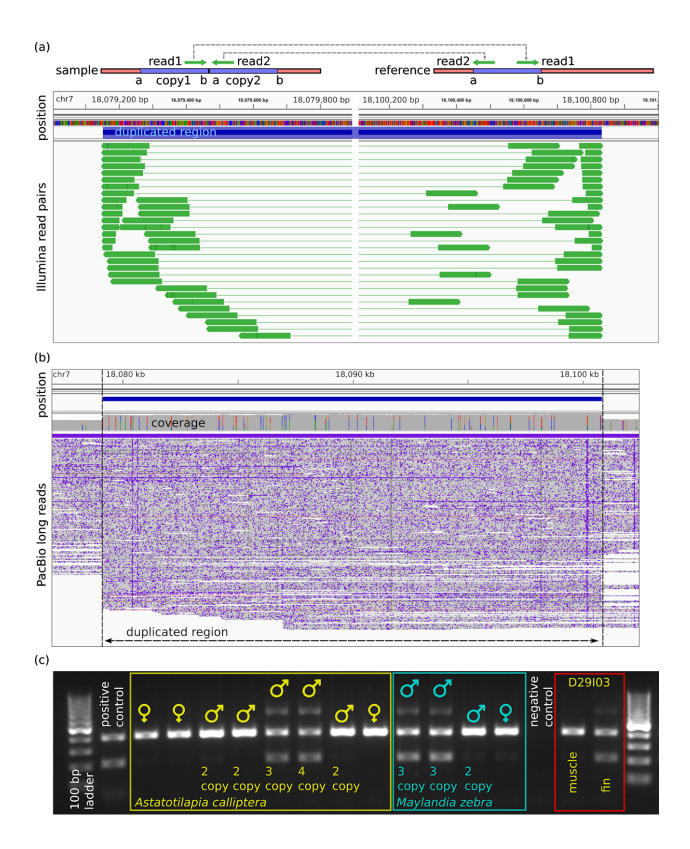
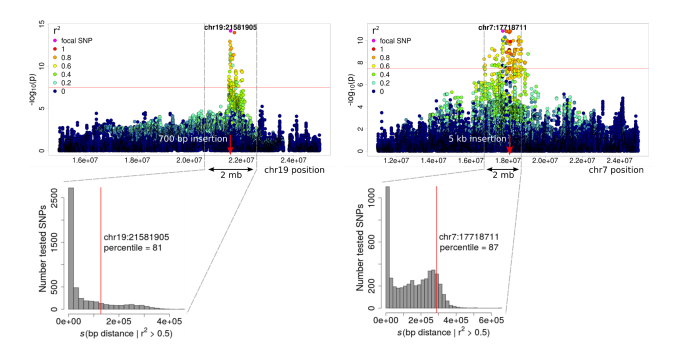


Figure S1: Characterization of the *gsdf* duplication. (a) Short Illumina reads from four
 Masoko male *A. calliptera* called homozygous for the *gsdf* duplication based on relative

sequencing depth that is approximately 2x higher than in ~38 kb of non-duplicated flanking
sequence. The mapping orientation of all read pairs to the fAstCal1.2 reference is consistent
with a tandem duplication as shown in the schematic at the top. (b) PacBio reads from a male
Tropheops 'mauve' mapped to the fAstCal1.2 reference. The sharp break in the alignment of
some of the reads at the edges of the gsdf duplication (blue horizontal bar) in conjunction with
elevated coverage signals that this individual is heterozygous for the same gsdf duplication
identified in Masoko A. calliptera. (c) Agarose gel image of PCR products from primers
designed to assay for the presence of the gsdf duplication. Based on this assay, individuals
positive for the gsdf duplication yield three distinct bands, whereas those negative for the
duplication produce a single band. The assay was used to confirm the presence of the
duplication in two male Maylandia zebra samples that were putative heterozygotes for gsdf-dup
based on sequencing depth. Two separate tissues for Masoko A. calliptera sample D29I03
produced different genotypes based on this PCR assay indicating a sampling error and resulted
in this individual being omitted from all analyses.



1014 **Figure S2: Elevated linkage disequilibrium around the chr19-ins and chr7-ins loci.** The top 1015 Manhattan plots are a regional view of the p-values for the likelihood ratio test from the GWAS 1016 for sex used to identify SNPs tagging chr19-ins (left) and chr7-ins (right). The positions of the 1017 insertions are denoted with red arrows. Elevated linkage disequilibrium (LD) between the SNP 1018 with the highest sex association in each GWAS and other surrounding SNPs extends far along

the respective chromosomes. This causes the variance in the pairwise physical distance among SNPs in high LD ($r^2 > 0.5$) with the top GWAS SNPs to be higher than typically expected throughout the genome, consistent with recent positive selection. The histograms show where this variance for the top GWAS SNPs fall along the expected distributions for Masoko *A.* calliptera, which were generated by randomly sampling 5,000 SNPs across the genome with the same alternate allele frequencies as the GWAS SNPs. The variance among the pairwise distances between each sampled SNP and their surrounding high-LD SNPs were calculated in the same manner as for the GWAS SNPs.

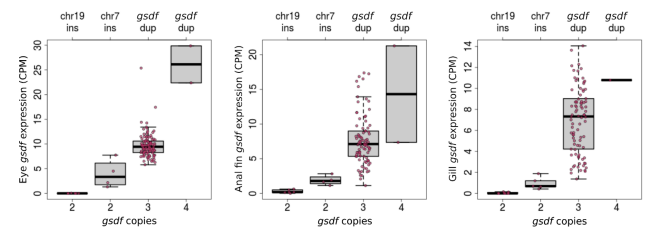


Figure S3: Expression of *gsdf* in somatic tissues for males with different Y alleles. The gsdf-dup and chr7-ins alleles are defined by a tandem duplication of the gsdf gene and an insertion directly upstream of gsdf, respectively. Levels of gsdf expression in eye, anal fin, and gill tissues from Masoko male *A. calliptera* demonstrate that males carrying putative Y alleles generated through mutations involving gsdf express this gene more than other males.

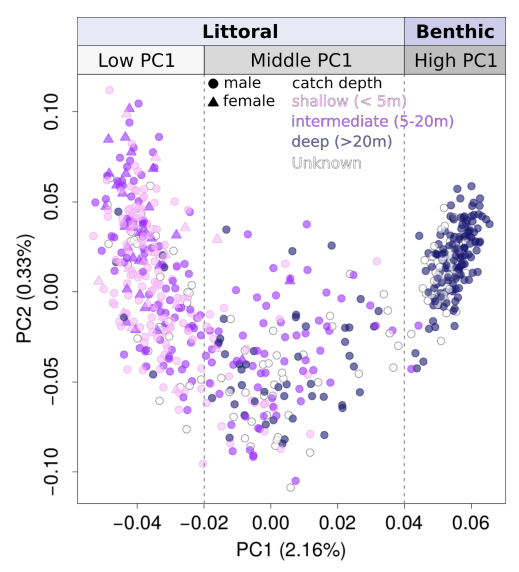
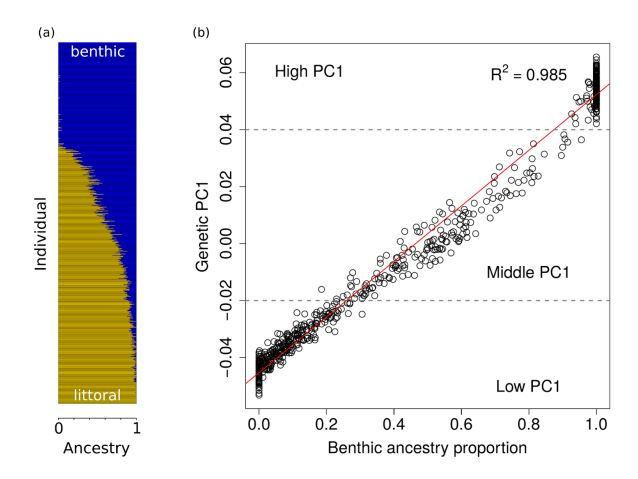


Figure S4: Relationship between genetic variation and catch depth. Lake Masoko *A*.

1033 *calliptera* distributed along the first two components of a principal component analysis of
1034 genome-wide variation reveals strong philopatry of high PC1 fish for deep depths. This
1035 coincides with nearly all high PC1 individuals conforming to the benthic ecomorph. In contrast,
1036 fish below PC1 values of 0.04 are almost all of the littoral ecomorph and exhibit far less
1037 constrained habitat preference. Among littoral fish (PC1 < 0.04), the most admixed individuals in
1038 the middle of PC1 (-0.02 to 0.04) regularly occupy all depth bands, while low PC1 littorals (PC1
1039 < -0.02) remain mostly at depths above 20 metres, though occasionally they are found deep.



1040 Figure S5: Ancestry characterization of Masoko *A. calliptera*. (a) Genome-wide ancestry 1041 proportions for individuals inferred using the program ADMIXTURE and ordered by their genetic 1042 PC1 rank shows the genetic distinctiveness of the benthic (high PC1) subgroup, a subset of 1043 littorals having low amounts of benthic ancestry (low PC1), and a highly admixed group (middle 1044 PC1). (b) The genetic PC1 scores of Lake Masoko individuals regressed against their 1045 proportion of benthic ancestry shows that PC1 almost perfectly describes the genetic structure 1046 of the Lake Masoko population in terms of the continuum between genetically benthic and 1047 littoral ancestries. The fitted linear regression line is shown in red and the low, middle, and high 1048 PC1 classification cutoffs are depicted with dashed grey lines.

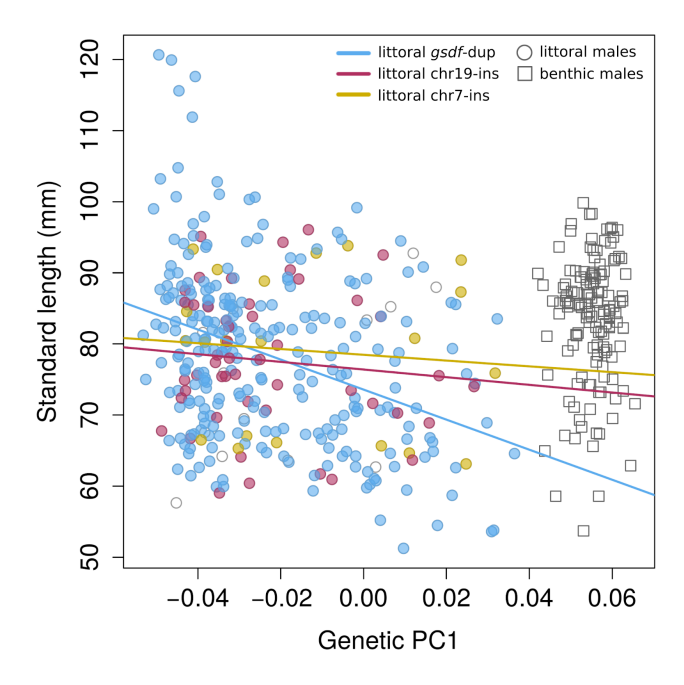
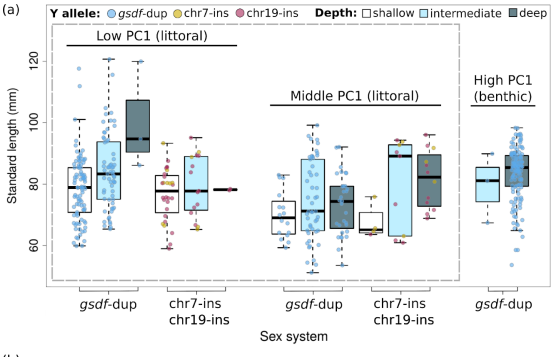


Figure S6: Interaction between genetic background and Y allele in predicting male size.

The standard lengths of male *A. calliptera* from Lake Masoko plotted against their position along PC1 of the principal component analysis of genome-wide variation shows a negative trend in the length among genetically littoral (PC1 < 0.04) males (circles) with increasing PC1 value.

Linear regression models of length predicted by PC1 were fitted separately for littoral males heterozygous for either *gsdf*-dup, chr19-ins, or chr7-ins corresponding to the colours blue, red, and yellow, respectively. Littoral males carrying more than one Y allele, homozygous for Y alleles, or which did not have an identified Y, are represented by uncoloured circles and were

1057 excluded from the regressions. Genetically benthic males, defined as fish with PC1 > 0.04, are 1058 plotted for comparative purposes as squares without any indication of their Y genotype. The 1059 distinctly more negative slope of the regression line fit to gsdf-dup males compared to chr19-ins 1060 and chr7-ins males shows that length is predicted to decrease much more drastically with more 1061 benthic admixture among gsdf-dup males. This difference is so great that males using gsdf-dup 1062 are predicted to switch from being longer than males using other Y alleles to actually being 1063 shorter above PC1 values of -0.02.



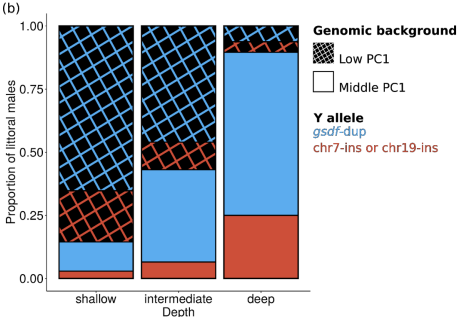


Figure S7: Male sizes and frequencies according to Y allele, genetic PC1, and catch depth. (a) Standard length comparisons across different PC1 genetic backgrounds and catch depths of Lake Masoko *A. calliptera* males heterozygous for only one of the Y alleles shows an interaction between Y allele, catch depth, and PC1 background in predicting size. Among the genetically littoral males (within the dashed grey box) those carrying *gsdf*-dup are smaller on middle PC1 versus low PC1 backgrounds regardless of what depth they are found at. In

1070 contrast, among males using the other Y alleles only middle PC1 males found in shallow waters
1071 are smaller than the low PC1 males, while at deeper depths their size remains constant across
1072 genetic backgrounds and may even show a subtle tendency to be larger with middle PC1
1073 benthic ancestry. **(b)** A comparison of the proportion of littoral males characterized by different
1074 genetic PC1 backgrounds and Y alleles at different catch depths shows that the proportion of
1075 males with middle PC1 ancestry increases with depth. However, within PC1 backgrounds, the
1076 fraction of males using the different Y alleles remains relatively stable across depths. Overall,
1077 *gsdf*-dup males dominate at all depths.

1078 **Tables S1 to S7** can be found in the attached Excel file:

supplementary_tables_differential_use_of_multiple_genetic_sex_determination_systems_in_div ergent_ecomorphs_of_an_African_crater_lake_cichlid.xls. For convenience the table legends are given below, and we also copy below the contents of tables S3 and S7, which are short.

1082 **Table S1: Lake Masoko** *Astatotilapia calliptera* **samples** Genetic, phenotypic, and collection 1083 information for all Lake Masoko *A. calliptera* samples.

Table S2: GWAS multilocus sex determination genotype frequencies Counts of Masoko *A.* 1085 *calliptera* individuals, stratified by sex and PC1 genetic background, for all observed combinations of *gsdf* copy number and genotypes at the most strongly associated SNPs in the 1087 serial GWAS for sex. 0 = reference allele, 1 = insertion allele, ./. = missing genotype.

1088 **Table S3: Average sizes of Masoko males** The mean standard length of Masoko *A. calliptera* 1089 males heterozygous for one type of Y allele stratified by PC1 genetic background and catch 1090 depth.

Lake-wide mean length (mm)					
Y allele	Low PC1	Middle PC1			
<i>gsdf</i> -dup	81.34	73.55			
chr7-ins or chr19-ins	77.68	78.73			

Shallow (< 5 m) mean length (mm)					
Y allele	Low PC1	Middle PC1			
gsdf-dup	78.55	69.91			
chr7-ins or chr19-ins	76.67	67.46			
,					
Intermediate (5-20 m) mean length (mm)					
Y allele	Low PC1	Middle PC1			
gsdf-dup	84.41	74.87			
chr7-ins or chr19-ins	79.50	79.96			
Deep (> 20 m) mean length (mm)					
Y allele	Low PC1	Middle PC1			
gsdf-dup	100.26	73.33			
chr7-ins or chr19-ins	78.22	81.56			

- 1091 **Table S4: Littoral male frequencies according to genetic type and catch depth** Counts of 1092 Lake Masoko *A. calliptera* littoral males heterozygous for one type of Y allele stratified by
- 1093 genetic PC1 background and depth at which they were caught.
- 1094 Table S5: Sex loci genotype calls for Lake Malawi cichlid radiation species The number of
- 1095 gsdf copies and genotype (GT) calls for chr19-ins and chr7-ins (0 = reference allele, 1 =
- 1096 insertion allele, ./. = missing genotype) for individuals of different species belonging to the Lake
- 1097 Malawi haplochromine cichlid radiation. The AC values indicate the number of "<reference
- 1098 allele>,<insertion allele>" sequencing reads observed for an individual.

Table S6: Frequency of chr7-ins in non-*calliptera* species from the Lake Malawi
haplochromine radiation Counts of individuals from all species apart from *Astatotilapia*calliptera in which chr7-ins was found, stratified by *gsdf* copy number and chr7-ins genotype.
Multilocus genotype calls are defined as <number of *gsdf* copies>/<number of chr7-ins alleles>:
for example, "3/1" denotes an individual possessing three *gsdf* copies and who is heterozygous
for the insertion allele at the chr7-ins locus. Genotype class cells with non-zero counts are
highlighted for readability.

1106 **Table S7: PCR primers for the detection of** *gsdf***-dup** All samples should undergo 1107 amplification for the 402 bp control fragment, whereas only samples positive for the gsdf 1108 duplication should show equally strong amplification for the 207 bp fragment (and an additional 1109 614 bp fragment which is not present when each primer pair is run in individual reactions).

primer	sequence	Tm (°C)	%GC	primer partner	amplicon size (bp)
dup_fwd	TGTCGCGTCATAACGAGGAG	59.9	55	dup_rev	207
dup_rev	AGCTGATCTGGTCCCTCACT	60.0	55	dup_fwd	
control_fwd	GCTGCCCACCTCGTAGTAAT	59.5	55	control_rev	402
control_rev	GCACGAGTGGGAACCAGTAA	60.0	55	control_fwd	
dup_fwd				control_rev	614