Sex differentiation and a chromosomal inversion lead to cryptic diversity in Lake Tanganyika sardines

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- 4 Julian Junker^{*1,2}, Jessica A. Rick^{*3}, Peter B. McIntyre⁴, Ismael Kimirei⁵, Emmanuel A. Sweke^{5,7}, Julieth
- 5 B. Mosille⁵, Bernhard Wehrli^{1,6}, Christian Dinkel¹, Salome Mwaiko^{1,2}, Ole Seehausen^{1,2}, Catherine E.
- 6 Wagner³
- 7 * denotes equal contribution
- 8 1 EAWAG Swiss Federal Institute of Aquatic Science and Technology, CH-6047 Kastanienbaum,
 9 Switzerland
- 2 Division of Aquatic Ecology, Institute of Ecology & Evolution, University of Bern, CH-3012 Bern,
 Switzerland
- 3 Department of Botany and Program in Ecology, University of Wyoming, Laramie, Wyoming 82072
 USA
- 14 4 Department of Natural Resources, Cornell University, Ithaca NY 14850 USA
- 15 5 Tanzania Fisheries Research Institute (TAFIRI), Dar es Salaam, Tanzania
- 16 6 Institute of Biogeochemistry and Pollutant Dynamics, ETH Zurich, CH-8092 Zürich, Switzerland
- 17 7 Deep Sea Fishing Authority (DSFA), Zanzibar, Tanzania
- 18 Corresponding authors: Julian Junker (Julian.junker@eawag.ch), Jessica Rick (jrick@uwyo.edu) and

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- 19 Catherine E. Wagner (<u>Catherine.Wagner@uwyo.edu</u>)
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- 36 Abstract

37 Two endemic sardines in Lake Tanganyika, Limnothrissa miodon and Stolothrissa tanganicae, are 38 important components of the lake's total annual fishery harvest. These two species along with four 39 endemic Lates species represent the dominant species in Lake Tanganyika's pelagic fish community, 40 in contrast to the complex pelagic communities in nearby Lake Malawi and Victoria. We use reduced 41 representation genomic sequencing methods to gain a better understanding of possible genetic 42 structure among and within populations of Lake Tanganyika's sardines. Samples were collected along 43 the Tanzanian, Congolese, and Zambian shores, as well as from nearby Lake Kivu, where Limnothrissa 44 was introduced in 1959. Our results reveal unexpected cryptic differentiation within both Stolothrissa 45 and Limnothrissa. We resolve this genetic structure to be due to the presence of large sex-specific 46 regions in the genomes of both species, but involving different polymorphic sites in each species. 47 Additionally, we find a large segregating inversion in *Limnothrissa*. We find all inversion karyotypes 48 throughout the lake, but the frequencies vary along a north-south gradient within Lake Tanganyika, 49 and differ substantially in the introduced Lake Kivu population. Little to no spatial genetic structure 50 exists outside the inversion, even over the hundreds of kilometres covered by our sampling. These 51 genetic analyses show that Lake Tanganyika's sardines have dynamically evolving genomes, and the 52 analyses here represent a key first step in understanding the genetic structure of the Lake 53 Tanganyika pelagic sardines. 54 55 Keywords: Stolothrissa tanganicae, Limnothrissa miodon, Lake Tanganyika, Inversion, sex-specific 56 region

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59 Introduction

60	Identifying the genetic basis of ecological adaptation is a high priority in evolutionary biology and has
61	important implications for population management. Recent research in this field focuses on genomic
62	regions with reduced recombination rates, such as chromosomal inversions (e.g. Berg et al. 2017;
63	Christmas et al. 2018; Kirubakaran et al. 2016; Lindtke et al. 2017), sex chromosome regions
64	(Presgraves 2008; Qvarnstrom & Bailey 2009) or both (Connallon et al. 2018; Hooper et al. 2019;
65	Natri et al. 2019). The reduced recombination rates in such chromosomal regions enable local
66	adaptation even when gene flow is high (Kirkpatrick & Barton 2006). Furthermore, it appears that
67	these mechanisms for restricted recombination are more prevalent in sympatric than in allopatric
68	species, and fixation of inversions is faster in lineages with high rates of dispersal and gene flow (Berg
69	et al. 2017). These patterns are consistent with theory in which the presence of gene flow favours
70	diversification of chromosomal rearrangements caused by locally adapted loci (Berg et al. 2017;
71	Kirkpatrick & Barton 2006).
72	Pelagic habitats represent uniform environments that allow for high dispersal rates due to the lack of
73	physical barriers. Well known examples of species from pelagic habitats that carry chromosomal
74	inversions or sex loci include Atlantic cod (Gadus moruha) (Berg et al. 2017; Kirubakaran et al. 2016),
75	Atlantic herring (Clupea harengus) (Lamichhaney et al. 2017; Martinez Barrio et al. 2016) and
76	stickleback (Gasterosteus aculeatus) (Jones et al. 2012). In Atlantic cod and herring populations, low
77	genome-wide divergence is interspersed with highly divergent inverted regions. These inversions in
78	cod distinguish between resident and migrating ecotypes (Berg et al. 2017; Kirubakaran et al. 2016),
79	and in herring they separate spring and fall spawners (Lamichhaney et al. 2017; Martinez Barrio et al.
80	2016). Additionally, inverted genomic regions in sticklebacks are involved in the divergence between
81	lake and stream ecotypes (Marques et al. 2016; Roesti et al. 2015).

82 From management perspectives, pelagic mixed stocks are notoriously difficult (Belgrano & Fowler 83 2011; Botsford et al. 1997) and part of this challenge lies in identifying Management Units (MUs) 84 which are demographically independent and genetically distinct populations. In a uniform habitat 85 without physical barriers, low genetic differentiation is typical, as there exist few environmental 86 restrictions to gene flow. However, there are increasingly cases where small genomic differences 87 lead to important variation in life history, influencing population resilience to fishing pressure (Berg 88 et al. 2017: Hutchinson 2008: Kirubakaran et al. 2016). The use of next generation sequencing 89 methods is therefore needed to shed light on population structure, particularly in species with low 90 genetic differentiation, to facilitate the detection of chromosomal variants which may be linked to 91 important ecological or local adaptation or selection (Lamichhaney et al. 2017). This is because 92 detailed information on the population structure, ecology and life history of harvested species is 93 crucial for effective fisheries management. 94 Lake Tanganyika is volumetrically the second largest lake in the world consisting of deep basins in the 95 north (~1200 m) and south (~1400 m), and a shallower basin (~800 m) in the middle region (Fig 1A) 96 (McGlue et al. 2007). At 9-12 million years in age (Cohen et al. 1993), it hosts a long history of 97 evolution, which has produced remarkable animal communities consisting largely of endemic species 98 (Coulter 1991). Among these endemics are six fish species which comprise the bulk of the lake's 99 pelagic fish community. These are two sardines, Stolothrissa tanganicae and Limnothrissa miodon, 100 and four endemic relatives of the Nile perch, Lates stappersii, Lates mariae, Lates angustifrons and 101 Lates microlepis. While little is known about the evolutionary history of the Lates species, Wilson et 102 al. (2008) showed evidence that the sardines of Lake Tanganyika descend from relatives in western 103 Africa and diverged from a common ancestor about 8 MYA. The harvest of Stolothrissa, Limnothrissa 104 and L. stappersii account for up to 95% of all catches within the lake (Coulter 1976, 1991; Mölsä et 105 al. 2002), making the second largest inland fishery on the continent of Africa (FAO 1995). The fishing 106 industry provides employment to an estimated 160'000 (Van der Knaap et al. 2014) to 1 million

107	people (Kimirei et al. 2008) and is an important source of protein to additional millions living on the
108	shores of Lake Tanganyika and further inland (Kimirei <i>et al.</i> 2008; Mölsä <i>et al.</i> 2002; Sarvala <i>et al.</i>
109	2002; Van der Knaap et al. 2014). Due to human population growth and an increased demand for
110	protein, fishing pressure has increased during the last decades, resulting in a decline of pelagic fish
111	stocks (Coulter 1991; van der Knaap 2013; Van der Knaap et al. 2014; van Zwieten et al. 2002). Also,
112	long-term decrease in fish abundance is likely linked to the observed warming of Lake Tanganyika
113	since the early 1900s, and further warming-induced decline in the lake's productivity is expected
114	during the 21 st century (Cohen <i>et al.</i> 2016; O'Reilly <i>et al.</i> 2003; Verburg & Hecky 2003; Verburg <i>et al.</i>
115	2003). Consequently, there is increasing recognition of the need to develop sustainable management
116	strategies for the lake's pelagic fish stocks (Kimirei <i>et al.</i> 2008; Mölsä <i>et al.</i> 1999; Mölsä <i>et al.</i> 2002;
117	van der Knaap 2013; Van der Knaap <i>et al.</i> 2014; van Zwieten <i>et al.</i> 2002).
118	Despite the economic importance of the pelagic fisheries in this lake, very little previous work has
119	investigated the genetic and phenotypic diversity and population structure of the key pelagic fish
120	species or their evolutionary origins (but see De Keyzer et al. 2019; Hauser et al. 1995, 1998; Wilson
121	et al. 2008). Lake Tanganyika's enormous size and spatial heterogeneity (e.g. Kurki et al. 1999;
122	Loiselle et al. 2014) harbours the potential for spatial segregation that may lead to temporal
123	differences in spawning and life history timing between distant sites. There are indeed indications of
124	genetically differentiated stocks of some of the pelagic fish of Lake Tanganyika known from basic
125	genetic work conducted two decades ago. For the sardines, these studies found no clear genetic
126	population structure at a large geographical scale (Hauser <i>et al.</i> 1998; Kuusipalo 1999), but some
127	small scale differences were found for Limnothrissa (Hauser et al. 1998). However, the genetic
128	methods used in these older studies (RAPDs and microsatellites) have limited power, are known to
129	suffer from error (RAPD, Williams et al. 1990), and in addition, have severe limitations in their
130	resolution. De Keyzer et al. (2019) used a modern RAD sequencing approach to examine Stolothrissa

131 and found small, if any, spatial structure in Stolothrissa sampled from the north, middle, and south of 132 Lake Tanganyika.

133	In this study, we focus on both sardine species, sampled from 13 sites spanning from the north to the
134	south of Lake Tanganyika (Fig 1). We also included Limnothrissa individuals from the introduced
135	population of this species present in Lake Kivu. Our null hypothesis was extremely simple: the surface
136	water of a large lake is horizontally well mixed and therefore provide a homogeneous habitat. Pelagic
137	fish can move freely and therefore due to the uniform environment, we should expect a lack of
138	genetic structure of their populations due to free interbreeding. Using reduced representation
139	genomic sequencing (RAD, Baird et al. 2008) we indeed do not find spatial genetic structure in either
140	species, supporting this null hypothesis. However, many loci deviating from Hardy-Weinberg
141	equilibrium differentiated the sexes in our samples, suggesting that these species have large sex-
142	determining regions. Furthermore, we find additional cryptic diversity in Limnothrissa due to genetic
143	patterns consistent with a chromosomal inversion. The generally low spatial genetic structure within
144	these species facilitated the detection of the differentiated loci, which may be related to sex-specific
145	or local adaptation.

146

147 **Material and Methods**

148 Study system and sampling

149 Our samples from Lake Tanganyika come from Tanzanian, Congolese and Zambian sites. Additionally 150 we added Rwandan Limnothrissa from Lake Kivu, where the species was introduced during the 1950s 151 (Collart 1960, 1989; Hauser et al. 1995) (Fig 1 and Table 1). Each fish was processed according to 152 standard protocols, during which we take a cuvette photograph of the live fish and subsequently 153 euthanize the fish with an overdose of MS222, and take fin clips and muscle tissue samples for 154 genetic analysis and stable isotope analysis, respectively. The specimens are preserved in

155	formaldehyde and archived in	the collections at EAV	WAG (2016, 2017,	2018 samples),	the University
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156 of Wyoming Museum of Vertebrates (2015 samples), and the University of Wisconsin- Madison

157 (2015 samples). Many fish for this project were obtained from fishermen and were already dead, and

158 in this case we completed this same protocol without euthanasia.

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160 Phenotypic sexing

161 Tanganyikan sardines caught by fishermen are frequently dried after being landed at the beach and

- although this does not inhibit the extraction of high-quality DNA, desiccated individuals cannot be
- accurately sexed. Therefore, we dissected 34 *Limnothrissa* and 15 *Stolothrissa* that were euthanized

and preserved in formalin just after being caught. These individuals were fully mature and in

165 excellent condition to accurately phenotypically sex them. We used these phenotypically sexed

166 individuals to determine whether inferred genetic groups correlated to sex in each species.

167

168 RAD sequencing

169 We extracted DNA from 486 individuals (181 Stolothrissa; 291 Limnothrissa) and obtained genomic 170 sequence data of these individuals using a reduced-representation genomic sequencing approach 171 (RADseq). Both species were pooled, divided into 10 RAD libraries, and sequenced. The DNA from all 172 individuals was extracted using Qiagen DNeasy Blood and Tissue kits (Qiagen, Switzerland). For 190 individuals collected in 2015, this DNA was then standardized to $20 ng/\mu L$ at the University of 173 174 Wyoming, and then prepared for RAD sequencing by Floragenex Inc. (Eugene, Oregon), and 175 sequenced at the University of Oregon on an Illumina HiSeg2000 (100bp SE). Individuals were 176 multiplexed in groups of 95 individuals using P1 adapters with custom 10 base pair barcodes, and 177 fragments between 200 and 400bp were selected for sequencing. In order to avoid library effects, 178 each individual was sequenced in two different libraries and the reads were combined after

179	sequencing. The other 296 individuals collected in 2016 and 2017 were prepared for sequencing
180	following the protocol by Baird et al. (2008) with slight modifications, including using between 400ng
181	and 1000ng genomic DNA per sample and digesting with Sbfl overnight. We multiplexed between 24
182	and 67 of these individuals per library and used P1 adapters (synthesized by Microsynth) with custom
183	six to eight base pair barcodes. These six libraries were sheared using an S220 series Adaptive
184	Focused Acoustic (AFA) ultra-sonicator (Covaris Inc. 2012) with the manufacturer's settings for a 500
185	bp mean fragment size. We selected fragments with a size between 300 and 700bp using a SageElf
186	(Sage Scientific Electrophoretic Lateral Fractionator; Sage Science, Beverly, MA). The enrichment step
187	was done in eight aliquots with a total volume of 200 μ l. Volumes were combined prior to a final size
188	selection step using the SageELF. Sequencing was done by the Lausanne Genomic Technologies
189	sequencing facilities of University of Lausanne, Switzerland. We sequenced each of six libraries on a
190	single lane using an Illumina HiSeq2000 (Illumina Inc. 2010) (100bp SE) together with 7–20%
191	bacteriophage PhiX genomic DNA.
192	

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193 Sequence data preparation

- 194 We filtered raw sequencing reads from each library by first removing PhiX reads using bowtie2
- 195 (Langmead & Salzberg 2012). Then we filtered reads for an intact Sbfl restriction site, de-multiplexed
- 196 the fastq file, and trimmed the reads down to 84 nucleotides using process_radtags from Stacks
- 197 v1.26 (Catchen *et al.* 2013) and a custom bash script. The FASTX toolkit v.0.0.13
- 198 (<u>http://hannonlab.cshl.edu/fastx_toolkit/</u>) and custom python scripts were used for quality filtering.
- 199 In a first step, we kept only reads with 100% of the bases with quality score of at least 10 and in a
- second step, we removed all reads with at least 5% of the bases with quality score below 30.

201

202 Assembly to reference genome

203	We generated a reference genome from a male Limnothrissa individual collected near Kigoma,
204	Tanzania, in 2018. High molecular weight DNA was extracted from fin tissue using the Qiagen HMW
205	gDNA MagAttract Kit, and then libraries were prepared using 10X Genomics Chromium library
206	preparation at the Hudson-Alpha Institute for Biotechnology Genomic Services Laboratory
207	(Huntsville, AL). The sequencing libraries were then sequenced on the Illumina HiSeq Xten platform
208	(150bp PE reads). Read quality was checked using FASTQC (Andrews 2010), and then reads were
209	assembled using 10X Genomics' Supernova assembly software, using a maximum of 500 million
210	reads. Assembly completeness was assessed using QUAST-LG (Mikheenko et al. 2018), which
211	computes both standard summary statistics and detects the presence of orthologous gene
212	sequences.
213	Reads for all Limnothrissa and Stolothrissa individuals were aligned to the reference genome using
214	BWA mem (Li & Durbin 2009), following the filtering steps discussed above. Alignments were then
215	processed using SAMtools v1.8 (Li et al. 2009b). We then identified variable sites in three different
216	groups using SAMtools mpileup and bcftools v1.8 (Li et al. 2009a): (1) all individuals; (2) only
217	Limnothrissa individuals; and (3) only Stolothrissa individuals. We obtained consistent results using
218	different combinations of more stringent and relaxed filtering steps. The results shown here are
219	based on a filtering as follows: within the two monospecific groups, we filtered SNPs using VCFTOOLS
220	(Danecek et al. 2011) to allow no more than 50% missing data per site, removed SNPs with a minor
221	allele frequency less than 0.01, included only high-quality variants (QUAL > 19), and retained only
222	biallelic SNPs. For the dataset including both species, we relaxed the missing data filter to allow sites
223	with up to 75% missing data.

224

225 Population genetics and outlier detection

226 After removing individuals with more than 25% missing data, we used the combined dataset to 227 conduct principal component analysis (PCA) using the R package SNPrelate (Zheng et al. 2012). To 228 delineate and visualize distinct groups, we performed K-means clustering (kmeans in R) on the first 229 five principal component axes. The value for K was chosen using the broken-stick method based on 230 the within-group sums of squares. We then used these groupings to assign individual fish to species 231 and clusters within species. We combined these clustering results with sexed phenotypes to confirm 232 the identity of each of these clusters. 233 After observing that the primary axis of differentiation in both Stolothrissa and Limnothrissa was 234 based on sex, we used the single-species SNP datasets and the R package adegenet (Jombart 2008) 235 to conduct discriminant analysis of principal components (DAPC, Jombart et al. 2010) on males 236 versus females of Stolothrissa and Limnothrissa to identify loci contributing to these sex differences. 237 We visually inspected the DAPC loading plots to determine an appropriate threshold for loading 238 significance and pulled out loci with loadings above these thresholds in each species. We then 239 calculated heterozygosity for these sex-associated loci using adegenet in R. 240 If Stolothrissa and Limnothrissa have a shared origin of these sex-linked loci, then we would expect 241 them to occur in similar locations in the genome; however, if the sets of significant SNPs are located 242 on different scaffolds for each of the two species, then expect these regions to more likely originate 243 from independent evolution. We therefore checked whether the same genomic regions explain 244 genetic differentiation between sexes in the two species. For this, we compared the location of SNPs 245 identified in each of the Stolothrissa and Limnothrissa DAPC analyses, both using the species-specific 246 and combined SNP data sets. We assessed the proportion of scaffolds shared among the two sets of 247 significant SNPs. As an additional comparison between the two species, we calculated the proportion 248 of Limnothrissa sex-linked SNPs that were polymorphic in Stolothrissa, and vice versa, as well as the 249 observed heterozygosity of *Limnothrissa* individuals at *Stolothrissa* sex-linked SNPs, and vice versa.

250	For each species, we also investigated population structure beyond sex differences to determine
251	whether there is any geographic signal of differentiation within each of the species. For this we
252	removed the sex specific SNPs in the species-specific datasets of both species. In Limnothrissa we
253	additionally removed the SNPs linked to the inverted region. We then calculated F_{ST} between all
254	sampling site pairs using VCFTOOLS (Danecek et al. 2011). In addition, we calculated pairwise genetic
255	distances between populations and used these in a Mantel test (using mantel.randtest() from
256	adegenet in R) for each species, which tests for an association between genetic distances and
257	Euclidean geographic distances between sites. For the Mantel tests, we used Edwards' Euclidean
258	genetic distance (calculated using dist.genpop() from adegenet in R) and omitted Lake Kivu, as well as
259	locations with fewer than 10 samples.
260	In Limnothrissa, the secondary axis of genetic differentiation clearly split the populations into three
261	genetic groups. To investigate the genetic basis of these groupings, we used DAPC to identify the loci
262	with high loadings on the differentiation between the two most extreme groups, using the dataset
263	where variants were called on <i>Limnothrissa</i> individually. In addition, we omitted Lake Kivu individuals
264	from this DAPC analysis. Once again, we visually inspected the loading plots to determine an
265	appropriate threshold for significance. We then calculated heterozygosity for these significant loci
266	using adegenet in R.
267	After assigning all individuals to one of the three distinct groups based on K-means clustering
268	(kmeans in R), we counted the frequencies of the three groups at each sampling site. To determine
269	whether the distribution of individuals among the clusters varied between regions in Lake
270	Tanganyika, we conducted a two-proportion z-test (prop.test() in R) between the three general
271	regions in Lake Tanganyika, as well as between each of these and Lake Kivu. Because patterns of
272	heterozygosity were consistent with these three groups being determined by a segregating
273	chromosomal inversion, we then tested whether the three genotypes are in Hardy-Weinberg

274 Equilibrium across all sampling sites, and within distinct geographic regions using the online tool

275 www.dr-petrek.eu/documents/HWE.xls

276 Genetic diversity within and among clusters

277 We performed population genetic analyses, including calculating genetic diversity within and

278 divergence between the different intraspecific groups, on the aligned BAM files using ANGSD

279 (Korneliussen et al. 2014), again using the Limnothrissa genome as a reference. Methods employed

- in ANGSD take genotype uncertainty into account instead of basing analyses on called genotypes,
- 281 which is especially useful for low- and medium-depth genomic data (Korneliussen et al. 2014), such
- as those obtained using RAD methods. From these alignment files, we first calculated the site allele
- frequency likelihoods based on individual genotype likelihoods (option -doSaf 1) using the samtools
- 284 model (option -GL 1), with major and minor alleles inferred from genotype likelihoods (option -
- 285 doMajorMinor 1) and allele frequencies estimated according to the major allele (option -doMaf 2).
- 286 We filtered sites for a minimum read depth of 1 and maximum depth of 100, minimum mapping
- 287 quality of 20, and minimum quality (q-score) of 20. From the site allele frequency spectrum, we then
- 288 calculated the maximum likelihood estimate of the folded site frequency spectrum (SFS) using the
- ANGSD realSFS program. The folded SFS was used to calculate per-site theta statistics and genome-
- 290 wide summary statistics, including genetic diversity, using the ANGSD thetaStat program

291 (Korneliussen et al. 2013). We performed each of these steps on all fish from each of Limnothrissa

- and Stolothrissa, and then individually for each sampling site, sex, and group (for Limnothrissa)
- 293 within each species.

294

295 Linkage disequilibrium among loci

296 To investigate the extent to which the loci identified by DAPC are linked to one another, we used

297 PLINK v1.9 (Purcell et al. 2007) to calculate pairwise linkage disequilibrium between all pairs of SNP

298	loci in our Limnothrissa and Stolothrissa data sets. Linkage disequilibrium was measured as the
299	squared allelic correlation (R^2, Pritchard & Przeworski 2001). We then subsetted each of these
300	comparisons to only the sex-linked loci identified using DAPC and compared the distribution of
301	linkage values among the sex-linked loci to those values between all SNPs in the dataset for each of
302	the two species. We then performed the same comparison for loci implicated in differences among
303	the three groups in Limnothrissa. To determine whether sex and grouping loci are more linked than
304	average across the genome, we performed a Mann-Whitney U test (wilcox.test() in R).
305	

306 Results

307 Genome assembly and variant calling

308 The final assembly of the 10X Genomics Chromium-generated reference genome for Limnothrissa

309 *miodon*, based on ~56x coverage, comprised 6730 scaffolds of length greater than 10Kb. The

assembly had a scaffold N50 of 456Kb and a total assembly size of 551.1Mb. The genome contained

311 83.5% complete single-copy BUSCO orthologs, as well as 4.62% fragmented and 11.82% missing

312 BUSCO genes. We retained only scaffolds > 10Kb in length for the reference genome used in

313 downstream alignment of the RAD reads.

314 The Floragenex libraries yielded between 306 and 328 million reads including 21–23% bacteriophage 315 PhiX genomic DNA, while the libraries sequenced at the Lausanne Genomic Technologies sequencing 316 facilities yielded between 167 and 248 million reads. On average, the mapping rate for Stolothrissa 317 individuals' RAD reads to the Limnothrissa reference genome was 80.2%, whereas it was 80.0% for 318 Limnothrissa individuals. We removed six Stolothrissa individuals and 10 Limnothrissa individuals due 319 to low quality reads, or too much missing data. After filtering, our species-specific RAD datasets 320 contained 8,323 SNPs from 175 Stolothrissa samples and 12,657 SNPs from 281 Limnothrissa 321 samples. The final dataset for the combined species approach contained 35,966 SNPs.

322

323 **Population structure**

324	Principal component analysis revealed two distinct genetic clusters in each species (Fig 2A). These
325	clusters correspond to sexes identified through sexing of individuals by dissection (Fig 2A and Table
326	S1). In a DAPC to identify the loci underlying the strong genetic differentiation of the sexes for
327	Stolothrissa, we visually selected a loadings cut off of 0.0009 on PC1 (Fig S1), which resulted in a total
328	of 369 (4.4%) significant SNPs distributed over 123 scaffolds with high loadings on sex difference. In
329	Limnothrissa, we selected a cut off of 0.0016 on PC1 based on the distribution of loadings (Fig S2).
330	This cut off resulted in 218 (1.7%) SNPs across 85 scaffolds with high loadings on sex differences. All
331	of these loci show an excess of homozygosity in females and an excess of heterozygosity in males (Fig
332	2C and 2E).
333	The sampling sites generally had similar levels of genetic diversity (Θ_{W}) for both species (Table 2,
334	Table 3). We found no evidence for significant spatial population structure or isolation by distance
335	within either Stolothrissa or Limnothrissa (Fig 3A). Within Stolothrissa, we found little evidence for
336	additional genetic structure beyond the genetic structure linked to sex (Fig 3B). In contrast, we find
337	very strong genetic structure within each sex in Limnothrissa (Fig 3C), suggesting the existence of
338	three distinct genetic groups of Limnothrissa in Lake Tanganyika. However, these three groups do not
339	correspond to geographic location where the fish were sampled.
340	

341 Evidence for a segregating inversion in Limnothrissa

342 Limnothrissa from Lake Kivu are divergent from individuals in Lake Tanganyika, but this

343 differentiation is weaker than that between the three groups observed within Lake Tanganyika (Fig

- 344 3C). The Limnothrissa individuals from Lake Kivu form additional clusters that are distinct from, but
- 345 parallel to, the Tanganyika clusters along the second and third PC axis (Fig 3C). Within Lake

346	Tanganyika, we found individuals of all three clusters at single sampling sites, and there is no clear
347	geographic signal to these groups (Fig 3C). DAPC analysis of the two most differentiated groups
348	within Lake Tanganyika identified 25 SNPs across 15 scaffolds with high loadings (> 0.006; Fig 4C and
349	S3). Among these SNPs with high loadings, we found that two clusters of <i>Limnothrissa</i> individuals
350	were predominantly homozygous for opposite alleles, while the third group consisted of
351	heterozygotes at these loci (Fig. 4D). This suggests that the three distinct genetic groups we observe
352	are due to a segregating inversion, with two of the groups representing homokaryotypes and the
353	third a heterokaryotype for these SNPs (Fig 4 and S3).
354	With this suggestion of a segregating inversion within Limnothrissa, we tested for Hardy-Weinberg
355	equilibrium among the three groups within the lake as a whole and at each sampling site individually
356	(Fig 5). In Lake Tanganyika, all sampling sites, regions, and the lake as a whole were in HWE (X^2 , p >
357	0.05), while Lake Kivu frequencies differed significantly from HWE (X^2 , p = 0.005) (Fig 5A and 5B). We
358	additionally found that the proportions of all three karyotype groups differed significantly between
359	Lake Kivu fish and the fish found in each of the north, middle (Mahale), and south basins in Lake
360	Tanganyika (p = 0.012, p = 0.0036, p << 0.001) (Fig 5B). This result seems to be driven by a much
361	higher frequency of genotype group 3 in Lake Kivu samples than was found in Lake Tanganyika (Fig
362	5B). The only difference between the three basins within Lake Tanganyika was that the northern
363	basin had a greater frequency of fish with genotype group 3 than either the Mahale or southern
364	basins (p = 0.025; all others p > 0.05) (Fig 5B).

365

366 Linkage disequilibrium among identified loci

367 The distribution of pairwise linkage disequilibrium values among loci in the species-specific and

368 species-combined datasets were highly right-skewed, with the majority of loci pairs having low to no

linkage (mean $R^2 = 0.009$) (Fig 6). In contrast, the subsets of loci identified as sex-linked in *Stolothrissa*

 that these sets of loci are more tightly linked than expected based on the distribution of LD values for all loci (Mann-Whitney test; <i>Stolothrissa</i> W = 1167300000, p << 0.001; <i>Limnothrissa</i> W = 202550000, p << 0.001). In <i>Limnothrissa</i>, the group-delineating loci had a mean pairwise LD of 0.734, suggesting that they are also more linked than expected for random loci (Mann-Whitney test, W = 23862000000, p << 0.001), but less tightly linked than the sex-linked loci (Mann-Whitney test, W = 1862600, p << 0.001). 	370	and <i>Limnothrissa</i> had mean pairwise LD values of 0.858 and 0.844, respectively (Fig 6), suggesting
 p << 0.001). In <i>Limnothrissa</i>, the group-delineating loci had a mean pairwise LD of 0.734, suggesting that they are also more linked than expected for random loci (Mann-Whitney test, W = 23862000000, p << 0.001), but less tightly linked than the sex-linked loci (Mann-Whitney test, W = 1862600, p << 0.001). 	371	that these sets of loci are more tightly linked than expected based on the distribution of LD values for
 that they are also more linked than expected for random loci (Mann-Whitney test, W = 23862000000, p << 0.001), but less tightly linked than the sex-linked loci (Mann-Whitney test, W = 1862600, p << 0.001). 	372	all loci (Mann-Whitney test; Stolothrissa W = 1167300000, p << 0.001; Limnothrissa W = 202550000,
 23862000000, p << 0.001), but less tightly linked than the sex-linked loci (Mann-Whitney test, W = 1862600, p << 0.001). 	373	p << 0.001). In <i>Limnothrissa</i> , the group-delineating loci had a mean pairwise LD of 0.734, suggesting
376 1862600, p << 0.001).	374	that they are also more linked than expected for random loci (Mann-Whitney test, W =
	375	23862000000, p << 0.001), but less tightly linked than the sex-linked loci (Mann-Whitney test, W =
377	376	1862600, p << 0.001).
	377	

378 No overlap of sex linked loci in the two sardine species

379	To test if the sex-linked loci overlap between the species, we used the species-combined dataset to
380	perform DAPC between sexes for each species individually and identified loci with high loadings.
381	Using this approach, we identified 570 SNPs across 133 scaffolds in <i>Stolothrissa</i> (loading > 0.0006; Fig
382	S4) and 334 SNPs across 91 scaffolds in <i>Limnothrissa</i> linked to sex (loading > 0.001; Fig S5). These two
383	sets of loci were completely non-overlapping, suggesting that the sex-linked loci are unique in each
384	species. In addition, the scaffolds on which these loci were located were non-overlapping between
385	the species, suggesting that the discrepancy between identified loci is not simply due to different
386	coverage of the reference genome between Stolothrissa and Limnothrissa data. When looking at
387	Stolothrissa sex-linked SNPs in Limnothrissa individuals, only 2.5% are polymorphic, and only 0.8% of
388	Limnothrissa sex-linked SNPs are polymorphic in Stolothrissa. In addition, the sex loci for each species
389	do not show the same patterns of heterozygosity in the opposite species (Fig S6).

390

391 Discussion

392 Little to no spatial genetic structuring is a relatively common observation in pelagic fish species with

393 continuous habitats (e.g. Canales-Aguirre *et al.* 2016; Hutchinson *et al.* 2001; Momigliano *et al.*

394	2017). However, many studies show that pelagic fish species harbour genetic structure that does not
395	correspond with geographic distance, but instead correlates with ecological adaptation (Berg et al.
396	2017; Kirubakaran et al. 2016; Roesti et al. 2015). We present here the largest genomic data sets
397	analysed for the two freshwater sardines of Lake Tanganyika to date. We did not find evidence for
398	spatial genetic structure in Stolothrissa or Limnothrissa of Lake Tanganyika (Fig 3), despite the
399	immense size of this lake and extensive geographic sampling of populations of both species. Instead,
400	we find evidence for the existence of many sex-linked loci in both Stolothrissa and Limnothrissa,
401	including strong deviations from expected heterozygosity at these loci, with males being the
402	heterogametic sex (Fig 2). In Limnothrissa, we additionally find three cryptic genetic groups, and
403	patterns in heterozygosity indicate the presence of a segregating chromosomal inversion underlying
404	this genetic structure (Fig 4). All three inversion genotypes (homokaryotypes and heterokaryotype)
405	appear in Limnothrissa from both Lake Tanganyika and Lake Kivu, but relative frequencies of the
406	karyotypes differ among these populations (Fig 5).
407	

408 *Genetic sex differentiation in both species*

409 According to the canonical model of sex chromosome evolution, development of sex chromosomes 410 initiates with the appearance of a sex-determining allele in the vicinity of loci only favourable for one 411 of the sexes. Mechanisms reducing recombination, such as inversions, support the spread of the sex-412 determining allele in combination with the sexually antagonistic region due to high physical linkage. 413 Eventually neighbouring regions also reduce recombination rate and further mutations accumulate, 414 leading to the formation of a new sex chromosome (Bachtrog 2013; Gammerdinger & Kocher 2018; 415 Wright et al. 2016). Examples range from ancient, highly heteromorphic sex chromosomes, to recent neo-sex chromosomes, which are found in mammals (Cortez et al. 2014), avian species (Graves 416 417 2014), and fishes (Feulner et al. 2018; Gammerdinger et al. 2018; Gammerdinger & Kocher 2018;

418 Kitano & Peichel 2012; Pennell et al. 2015; Roberts et al. 2009; Ross et al. 2009; Yoshida et al. 2014). 419 Our results suggest that sex-linked regions of the genome in both Stolothrissa and Limnothrissa are 420 large and highly differentiated between males and females (Fig. 2). The results from our analyses of 421 linkage disequilibrium suggest that these loci are more tightly linked in both Limnothrissa and 422 Stolothrissa than SNP loci are on average (Fig 6). The high number of loci implicated in these genetic 423 sex differences, and high linkage among those loci, in addition to clear patterns of excess 424 heterozygosity in males and homozygosity in females, give strong indication of the existence of large 425 sex-determining regions in these species, which may form distinct sex chromosomes. However, the 426 structural arrangement of these loci remains unclear with our current reference genome. It is worth 427 noting that the assembly of sex chromosomes remains challenging due to the haploid nature of sex 428 chromosomes and therefore reduced sequencing depth, and existence of ampliconic and repetitive 429 regions and a high amount of heterochromatin (Tomaszkiewicz et al. 2017). Such challenges with 430 assembling sex chromosomes may lead to many scaffolds being implicated in sex determination in 431 initial attempts at assembly, as we see in our analysis, even if these species actually have distinct sex 432 chromosomes. 433 We also show that Stolothrissa and Limnothrissa SNPs linked to sex are entirely distinct, representing 434 strong evidence for rapid evolution in these sex-linked regions (Fig S1 and S2). This means that if the 435 common ancestor of these species had a sex-determining region, the variants on this sex-linked 436 region have entirely turned over and become distinct in the two species, during the approximately 437 eight million years since these species diverged (Wilson et al. 2008). Rapid turnover of sex 438 chromosomes in closely related species are known from a diversity of taxa (e.g. (Jeffries *et al.* 2018; 439 Kitano & Peichel 2012; Ross et al. 2009; Tennessen et al. 2018). The proposed mechanisms leading to 440 such rapid turnover rates are chromosomal fusions of an autosome with an already existing sex 441 chromosome, forming a "neo sex chromosome" (Kitano & Peichel 2012; Ross et al. 2009) or the

translocation of sex loci from one chromosome to another (Tennessen et al. 2018). Understanding

442

443 the mechanisms responsible for the high turnover rate of the sex chromosomes in the Tanganyikan

444 freshwater sardines is a fascinating area for future research.

445	Furthermore, it will be important for future work to investigate if the strong differentiation between
446	the sexes might also be associated with adaptive differences between the sexes. Ecological
447	polymorphism among sexes is known in fishes (Culumber & Tobler 2017; Laporte et al. 2018; Parker
448	1992) and can be ecologically as important as differences between species (Start & De Lisle 2018).
449	It is worth noting that the strong sex-linked genetic differentiation in Limnothrissa and Stolothrissa
450	could have been mistaken for population structure had we filtered our data for excess heterozygosity
451	without first examining it, and had we not been able to carefully phenotypically sex well-preserved,
452	reproductively mature individuals of both species to confirm that the two groups in each species do
453	indeed correspond to sex (Table S1, Fig 2A). Because of the strong deviations from expected
454	heterozygosity at sex-linked loci, any filtering for heterozygosity would remove these loci from the
455	dataset, explaining why one previous study in <i>Stolothrissa</i> using RADseq data (De Keyzer et al. 2019)
456	did not clearly identify this pattern despite its prevalence in the genome. For organisms with
457	unknown sex determination systems, and for whom sex is not readily identifiable from phenotype,
458	there is danger in conflating biased sampling of the sexes in different populations with population
459	structure in genomic datasets (e.g. Benestan et al. 2017). This underscores the importance of sexing
460	sampled individuals whenever possible when sex determination systems are unknown, when
461	analyzing large genomic datasets. The phenotypic and genetic sex of the sardine samples were in
462	agreement in all individuals except one <i>Stolothrissa</i> sample (Table S1, sample 138863.IKO02). This
463	fish was phenotypically identified as a male but genetically clustered with female individuals. We
464	believe that this individual was not yet fully mature, and therefore was misidentified phenotypically.
465	

No spatial genetic structure in Limnothrissa but cryptic diversity in sympatric Limnothrissa: 466

467	Our results reveal the existence of three distinct genetic groups of Limnothrissa. Intriguingly, we find
468	all three of these groups together within the same sampling sites, and even within the same single
469	school of juvenile fishes (Fig S7). Given patterns of heterozygosity at loci that have high loadings for
470	distinguishing among the genetic clusters (Fig 4D) together with the strong linkage (Fig 4E), this
471	structure is consistent with a chromosomal inversion. Chromosomal inversions, first described by
472	Sturtevant (1921), reduce recombination in the inverted region because of the prevention of
473	crossover in heterogametic individuals (Cooper 1945; Kirkpatrick 2010; Wellenreuther & Bernatchez
474	2018). Mutations in these chromosomal regions can therefore accumulate independently between
475	the inverted and non-inverted haplotype. Although early work on chromosomal inversions in
476	Drosophila has a rich history in evolutionary biology (Kirkpatrick 2010), inverted regions have
477	recently been increasingly detected with the help of new genomic sequencing technologies in many
478	species (e.g. Berg et al. 2017; Christmas et al. 2018; Kirubakaran et al. 2016; Lindtke et al. 2017;
479	Zinzow-Kramer et al. 2015), with implications for the evolution of the populations with distinct
480	inversion haplotypes. In Limnothrissa, the strong genetic divergence between the two inversion
481	haplotypes (Fig 3C, 4A and 4B) is consistent with this pattern, and indeed the substantial
482	independent evolution of these haplotypes is how the inversion is readily apparent even in a RADseq
483	dataset. The divergence of the haplotypes, and the high frequency of both of these haplotypes,
484	indicates that this inversion likely did not appear recently, although its apparent absence in
485	Stolothrissa indicates it has arisen since the divergence of these sister taxa eight million years ago
486	(95% reliability interval: 2.1–15.9 MYA; (Wilson <i>et al</i> . 2008)).
487	Given that both inversion haplotypes appear in relatively high numbers, it seems unlikely that drift
488	alone could explain the rise of the inversion haplotype to its current frequencies in the Limnothrissa
489	population. We expect that Limnothrissa have sustained large effective population sizes through
490	much of their evolutionary history since their split with Stolothrissa, meaning that drift would have

491 been a continually weak force. Although selection against inversions might occur due to an

492 inversion's disruption of meiosis or gene expression due to the position of the breakpoints

493 (Kirkpatrick 2010), selection may also act on inversions when they carry alleles that themselves are

494 under selection.

495 Due to the reduced recombination rates in inversions, these regions of the genome provide 496 opportunities for local or ecological adaptation despite ongoing gene flow (Kirkpatrick & Barton 497 2006). It is unclear given current data whether the inversion that we describe here in *Limnothrissa* is 498 tied to differential ecological adaptation. When we examine frequencies of the inversion karyotypes 499 pooled across all sampled populations, the observed frequencies do not differ from Hardy-Weinberg 500 expectations (chi-square = 3.51; p-value = 0.06). However, the Lake Kivu population does show 501 deviation from HWE (chi-square =7.74; p-value = 0.005) when we examine sampled populations 502 individually. Furthermore, frequencies of the inversion karyotypes among sampled populations 503 differ: the proportions of all three karyotypes differ significantly between Lake Kivu and Lake 504 Tanganyika populations, and within Lake Tanganyika, one of the homokaryotypes (represented as 505 group 3 in Fig 5), has a higher frequency in the northern basin than in the middle or southern basins 506 (Fig 5). In Lake Tanganyika, the southern and northern basin differ substantially in nutrient 507 abundance and limnological dynamics, and the Mahale Mountain (middle) region represents the 508 geographical transition between the two basins (Bergamino et al. 2010; Kraemer et al. 2015; Plisnier 509 et al. 1999; Plisnier et al. 2009). Thus, it is plausible that differential ecological selection could be 510 driving differences in the frequencies on the inversion karyotypes spatially within the lake. Genetic 511 drift is another possibility to explain the spatial differences in frequencies, and although this is highly 512 plausible in explaining the frequency differences between Lake Tanganyika and Lake Kivu (see 513 below), given the lack of spatial genetic structure in Lake Tanganyika it seems a less likely explanation 514 within this lake. Greater understanding of the ecology of these fishes in the north and the south of 515 Lake Tanganyika, and assessment of the genes within the inverted region, is needed to clarify this 516 question.

517 Comparing Limnothrissa populations in Lake Tanganyika to that introduced to Lake Kivu

518	We found substantial divergence between Limnothrissa in their native Lake Tanganyika and the
519	introduced population in Lake Kivu. This could derive from founder effects, from drift within this
520	population since their introduction in the absence of gene flow with the Lake Tanganyika population,
521	or from adaptive evolution in the Lake Kivu population since their introduction to this substantially
522	different lake environment. Future studies should examine these possibilities with a larger sample of
523	individuals from Lake Kivu. We identified individuals in Lake Kivu with all three inversion genotypes
524	that were detected in Lake Tanganyikan fish, suggesting that the inversion is also segregating in Lake
525	Kivu, and that the founding individuals likely harboured this genetic variation. That said, the
526	frequency of the karyotypes in Lake Kivu strongly deviates from Hardy-Weinberg expectations (Chi-
527	square =7.74; p-value =0.005). This suggests that the inversion locus may be under directional
528	selection. This finding is in contrast with frequent expectations for the behaviour of inversions:
529	typically, one would predict some sort of balancing selection (e.g. negative frequency dependent
530	selection) to maintain inversion haplotype diversity (Wellenreuther & Bernatchez 2018). Another
531	possibility is that the inversion locus is linked to non-random mating in the Kivu population. In
532	addition, the strong difference in the frequencies of the inversion haplotypes compared to Lake
533	Tanganyika populations has also likely been influenced by founder effects. Limnothrissa were
534	introduced to Lake Kivu in the 1950s (Hauser et al. (1995), and all introduced fish were brought from
535	the northern part of Lake Tanganyika. The homokaryotype, represented as group 3, is the prevalent
536	karyotype in Lake Kivu, and this karyotype also appears in highest frequencies in our samples from
537	northern Lake Tanganyika sites (Fig 5). Thus, it is plausible that founder effects could have led to the
538	increased frequency of this karyotype within the Lake Kivu population. This origin, however, does not
539	explain current deviations from HWE given that the population was introduced decades ago.

540 Conclusions

541 Genomic data from <i>Stolothrissa</i> and <i>Limnothrissa</i> reveal an interesting arra
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- 542 in chromosomal evolution. Modern fisheries management seeks to define locally adapted,
- 543 demographically independent units. We do not find significant spatial genetic structure within these
- 544 two freshwater sardine species from Lake Tanganyika. The genetic structure we find is all in
- 545 sympatry, namely as strong genetic divergence between the sexes, and evidence of a segregating
- 546 inversion in *Limnothrissa*. Further research should focus on the potential for adaptive differences
- 547 between the sexes and between the inversion genotypes in *Limnothrissa*. Such work will contribute
- to better understanding the role that these key components of the pelagic community assume in the
- 549 ecosystem of this lake, which provides important resources to millions of people living at its shores.
- 550

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723 Roesti M, Kueng B, Moser D, Berner D (2015) The genomics of ecological vicariance in threespine 724 stickleback fish. Nat Commun 6, 8767. 725 Ross JA, Urton JR, Boland J, Shapiro MD, Peichel CL (2009) Turnover of Sex Chromosomes in the 726 Stickleback Fishes (Gasterosteidae). PLOS Genetics 5, e1000391. 727 Sarvala J, Tarvainen M, Salonen K, Mölsä H (2002) Pelagic food web as the basis of fisheries in Lake 728 Tanganyika: A bioenergetic modeling analysis. Aquatic Ecosystem Health and Management 5, 729 283-292. 730 Start D, De Lisle S (2018) Sexual dimorphism in a top predator (Notophthalmus viridescens) drives 731 aquatic prey community assembly. Proc Biol Sci 285. 732 Sturtevant AH (1921) A Case of Rearrangement of Genes in Drosophila. *Genetics* 7, 235-237. 733 Tennessen JA, Wei N, Straub SCK, et al. (2018) Repeated translocation of a gene cassette drives sex-734 chromosome turnover in strawberries. PLoS Biol 16, e2006062. 735 Tomaszkiewicz M, Medvedev P, Makova KD (2017) Y and W Chromosome Assemblies: Approaches 736 and Discoveries. Trends Genet 33, 266-282. 737 van der Knaap M (2013) Comparative analysis of fisheries restoration and public participation in Lake 738 Victoria and Lake Tanganyika. Aquatic Ecosystem Health and Management 16, 279-287. 739 Van der Knaap M, Katonda KI, De Graaf GJ (2014) Lake Tanganyika fisheries frame survey analysis: 740 Assessment of the options for management of the fisheries of Lake Tanganyika. Aquatic 741 Ecosystem Health and Management 17, 4-13. 742 van Zwieten PAM, Roest FC, Machiels MAM, Van Densen WLT (2002) Effects of inter-annual 743 variability, seasonality and persistence on the perception of long-term trends in catch rates 744 of the industrial pelagic purse-seine fishery of northern Lake Tanganyika (Burundi). Fisheries 745 Research 54, 329-348. 746 Verburg P, Hecky RE (2003) Wind patterns, evaporation, and related physical variables in Lake 747 Tanganyika, east Africa. Journal of Great Lakes Research 29, 48-61. 748 Verburg P, Hecky RE, Kling H (2003) Ecological consequences of a century of warming in Lake 749 Tanganyika. Science **301**, 505-507. 750 Wellenreuther M, Bernatchez L (2018) Eco-Evolutionary Genomics of Chromosomal Inversions. 751 Trends Ecol Evol 33, 427-440. 752 Williams JGK, Kubelik AR, Livak KJ, Rafalski JA, Tingey SV (1990) DNA polymorphisms amplified by 753 arbitrary primers are useful as genetic markers. Nucleic Acids Research 18, 6531-6535. 754 Wilson AB, Teugels GG, Meyer A (2008) Marine incursion: the freshwater herring of Lake Tanganyika 755 are the product of a marine invasion into west Africa. PLoS One 3, e1979. 756 Wright AE, Dean R, Zimmer F, Mank JE (2016) How to make a sex chromosome. Nature 757 Communications 7, 12087. 758 Yoshida K, Makino T, Yamaguchi K, et al. (2014) Sex Chromosome Turnover Contributes to Genomic 759 Divergence between Incipient Stickleback Species. PLOS Genetics 10, e1004223. 760 Zheng X, Levine D, Shen J, et al. (2012) A high-performance computing toolset for relatedness and 761 principal component analysis of SNP data. Bioinformatics 28, 3326-3328. 762 Zinzow-Kramer WM, Horton BM, McKee CD, et al. (2015) Genes located in a chromosomal inversion 763 are correlated with territorial song in white-throated sparrows. Genes Brain Behav 14, 641-764 654. 765 766 **Data Accessibility Statement**

767 We are happy, to make our genetic data, including our reference genome, publically available by

768 submitting it to the European Nucleotide Archive (ENA). We intend to submit as soon as possible but

769 by the latest after acceptance of the manuscript.

- 771 Data Accessibility
- RAD sequences: will be uploaded to ENA as soon as possible but by the latest after acceptance
- Final DNA sequence assembly will be uploaded to ENA as soon as possible but by the latest after
- 774 acceptance
- 775
- 776
- 777 Author contributions:
- JJ: developing and writing SNSF grant, sampling and processing fish, identifying phenotypic sex of
- fish, DNA extractions, preparing RAD libraries, data analysis, writing on the manuscript
- JR: sampling and processing fish, DNA extractions, whole genome assembly, data analysis, writing onthe manuscript
- PBM: developing grant for The Nature Conservancy, contributing samples, discussing results,reviewing manuscript
- 784 IK: developing and writing SNSF grant, facilitating permission processes, providing logistics for
 785 fieldwork, reviewing manuscript
- EAS: sampling and processing fish, facilitating permission processes, providing logistics for fieldwork,
 enable collaboration with Tanzanian fishermen, discussing manuscript, reviewing manuscript
- JBM: sampling and processing fish, facilitating permission processes, providing logistics for fieldwork,
 enable collaboration with Tanzanian fishermen, discussing and reviewing manuscript
- 790 BW: developing and writing SNSF grant, reviewing and discussing manuscript
- 791 CD: developing SNSF grant, sampling and processing fish, facilitating logistics during fieldwork,792 reviewing manuscript
- SM: developing SNSF grant, facilitating permission process, RAD library preparation for sequencing,reviewing manuscript
- OS: developing and writing SNSF grant, identifying phenotypic sex of fish, facilitating permission
 process, reviewing and discussing manuscript
- 797 CEW: developing and writing SNSF and TNC grants, sampling and processing fish, contributing
- samples, whole genome assembly, data analysis, discussing results, writing manuscript, revisemanuscript
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Table 1. Fish collected from Democratic Republic of Congo (DRC), Tanzania (TNZ), Zambia (ZM) and Rwanda (RW).

Number of sequenced individuals

Stolothrissa	Limnothrissa	Site	Country		
0	21	Lake Kivu	RW		
7	0	Kilomoni	DRC		
15	0	Lusenda	DRC		
5	2	Kabimba	DRC		
15	17	Kagunga	TNZ		
61	49	Kigoma	TNZ		
25	37	North Mahale	TNZ		
6	38	South Mahale	TNZ		
18	11	Ikola	TNZ		
12	61	Kipili	TNZ		
0	41	Kasanga	TNZ		
17	1	Mbete	ZM		
0	13	Crocodile Island	ZM		
181 Samples	291 Samples	13 Sites	4 Countries		

Table 2. Genetic diversity within (Watterson's theta, Θ_w , along diagonal) and differentiation between (weighted F_{ST}, above diagonal) sampling sites (unshaded) and basins (shaded) for *Limnothrissa* populations included in this study.

	Kivu	NORTH	Kagunga	Kigoma	Kabimba	MIDDLE	North Mahale	South Mahale	Ikola	SOUTH	Kipili	Kasanga	Mbete	Crocodile Island
Kivu	0.0010													
NORTH		0.0040				0.173				0.117				
Kagunga			0.0017	0.0032	0.00022		0.0042	0.0055	0.0010		0.0039	0.0007	0.0019	0.0024
Kigoma				0.0014	0		0.0021	0.0017	0.0026		0.0043	0.0023	0.0052	0.0056
Kabimba					0.0011		0	0.0034	0.0004		0.0077	0	0	0.0019
MIDDLE						0.0043				0.211				
North Mahale							0.0007	0.0005	0.0037		0.0063	0.0033	0.0038	0.0053
South Mahale								0.0008	0.0043		0.0049	0.0039	0.0069	0.0142
Ikola									0.0014		0.0028	0.0028	0.0047	0.0106
SOUTH										0.0023				
Kipili											0.0032	0.0013	0.0079	0.0199
Kasanga												0.0018	0.0042	0.0015
Mbete													0.0015	0
Crocodile Island														0.0013

Table 3. Genetic diversity within (Watterson's theta, Θ_W , along diagonal) and differentiation between (weighted F_{sT}, above diagonal) sampling sites (unshaded) and basins (shaded) for *Stolothrissa* populations included in this study.

	NORTH	Kilomoni	Lusenda	Kagunga	Kigoma	Kabimba	MIDDLE	North Mahale	South Mahale	Ikola	SOUTH	Kipili	Mbete
NORTH	0.0044						0.243				0.315		
Kilomoni		0.0012	0.0038	0.0033	0.00001	0.0098		0.0002	0	0.0017		0.0002	0.0011
Lusenda			0.0007	0.0086	0.0105	0.0268		0.0088	0.0144	0.0115		0.0080	0.0008
Kagunga				0.0007	0.0033	0.0154		0.0045	0.0031	0.0014		0.0000	0.0056
Kigoma					0.0007	0.0107		0.0007	0	0.0040		0.0051	0.0075
Kabimba						0.0008		0.0140	0.0013	0.0142		0.0158	0.0218
MIDDLE							0.0014				0.358		
North Mahale								0.0014	0	0.0055		0.0061	0.0061
South Mahale									0.0014	0.0045		0.0064	0.0116
Ikola										0.001		0.0010	0.0089
SOUTH											0.0007		
Kipili												0.0011	0.0048
Mbete													0.0011

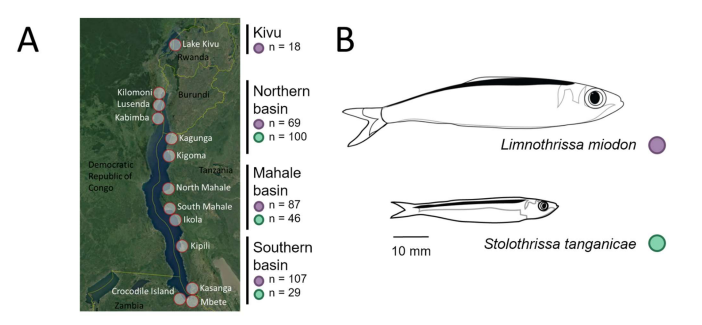


Figure 1. (A) Map of Lake Tanganyika, with sampling sites labeled and sample sizes from the three basins within Lake Tanganyika and Lake Kivu indicated for each species. (B) Drawings of *Limnothrissa miodon* and *Stolothrissa tanganicae*, with scale indicated for average mature sizes of the two species. Drawings courtesy of Jimena Golcher-Benavides.

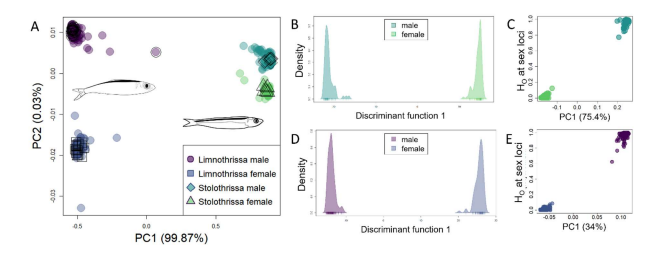


Figure 2. (A) Principal component analysis of all *Stolothrissa* and *Limnothrissa* individuals combined, colored by species identity and sex. Empty shapes denote individuals that were dissected and for whom sex was determined phenotypically. These dissection phenotypes group into genetic clusters, and therefore were used to identify the sex of each of the genetic clusters. In the combined PCA, the first axis generally corresponds to species, while the second axis corresponds to sex. Discriminant analysis of principal components (DAPC) results for (B) *Stolothrissa* and (D) *Limnothrissa* demonstrate distinct separation among males and females, with intraspecific differentiation (F_{ST}) between the two groups indicated. DAPC was used to identify loci associated with this differentiation (see Supplementary Figure S1 and S2); observed heterozygosity (H_{obs}) of each individual at those loci with high loadings is plotted against the first intraspecific PCA axis indicated in (C) for *Stolothrissa* and (E) for *Limnothrissa*, demonstrating both that sex dictates the first axis of differentiation in both species, and that males are the heterogametic sex at these loci in both species. There were 369 significant SNPs differentiating the sexes in *Stolothrissa* and 218 significant SNPs in *Limnothrissa*, with no overlap between the two species.

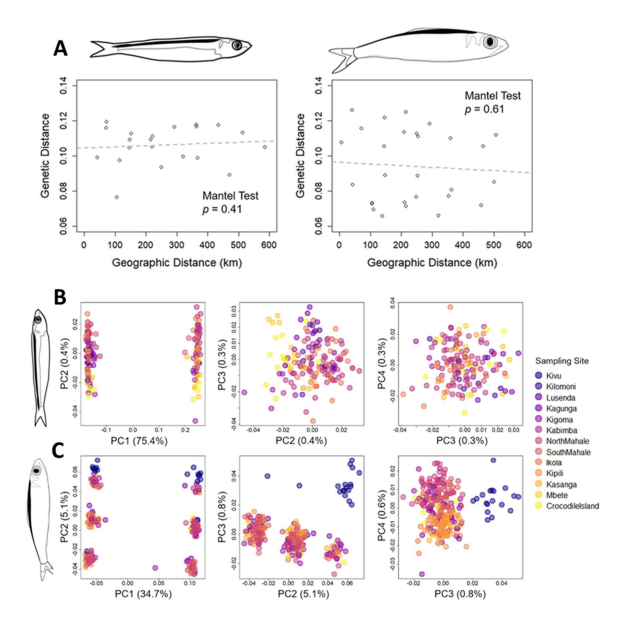


Figure 3. (A) shows the relationship between genetic and geographic distance. Neither species has evidence for statistically significant spatial population genetic structure using mantel test. (B) and (C) show Species-specific principal components analysis of *Stolothrissa* and *Limnothrissa* individuals, colored by sampling sites. In both species, PC1 differentiates the sexes; in *Limnothrissa*, PC2 separates each sex into three distinct groups, while PC3 separates individuals from Lake Kivu from those in Lake Tanganyika. Sampling sites are ordered from north (Kivu) to south (Crocodile Island).

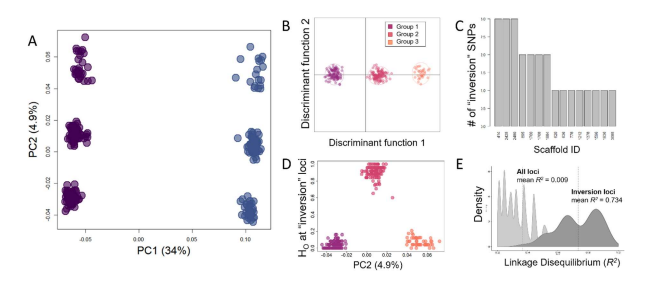


Figure 4. Evidence in *Limnothrissa* points to the existence of a segregating inversion. (A) Principal component analysis for all *Limnothrissa* individuals of Lake Tanganyika, demonstrating separation among sexes along the first principal component axis (PC1), and separation into three groups for both males (purple) and females (blue) on the second axis (PC2). For our discriminant analysis of principal components (B), we identified individuals according to these groups, and identified SNPs with high loadings along this axis. (C) Scaffold locations of SNPs with high loadings along this 'group' axis, showing that the 25 significant SNPs were found on 15 different scaffolds. At these significant loci, two groups were predominantly homozygous (D), while the third (intermediate) group was generally heterozygous. Despite the fact that these SNPs were spread out across several scaffolds, the distribution of pairwise linkage disequilibrium values (E) between just the inversion loci (dark) has a mean R² = 0.734, whereas the distribution for all loci in the data set for *Limnothrissa* (light) has a mean R² = 0.009.

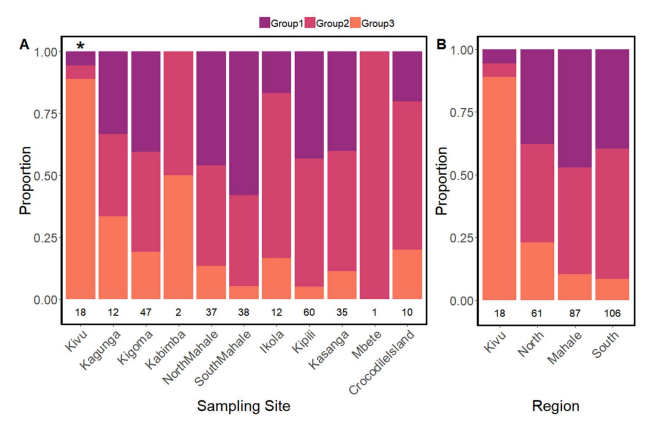


Figure 5. Proportion of individuals in each inversion karyotype group, by sampling site (A) and region (B). Sample sizes of individuals retained in analyses at each sampling site are indicated, as well as whether the distribution of the groups at each site differs from expectations under Hardy-Weinberg equilibrium (* indicates rejection of HWE at the site). Sites are ordered by geographic location, from north (Kivu, Kagunga) to south (Crocodile Island). The relative proportions of each haplotype observed differed significantly between Lake Kivu and all three regions in Lake Tanganyika, while only the relative proportion of Group 3 differed among the three regions within Lake Tanganyika.

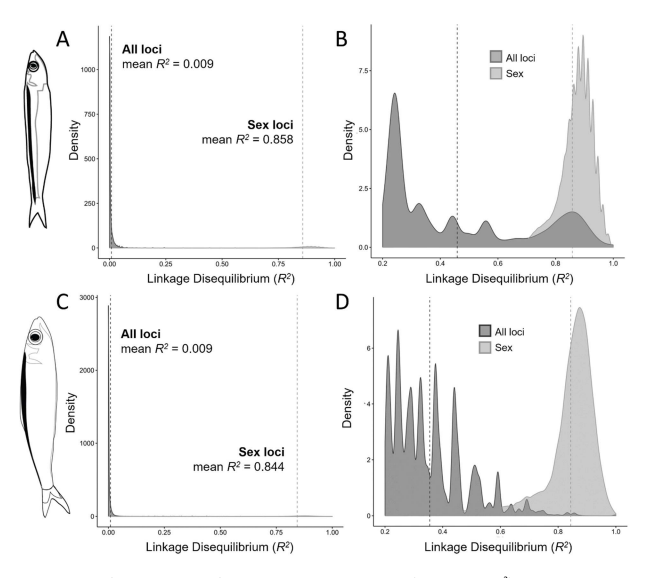


Figure 6. Results from an analysis of pairwise linkage disequilibrium (measured as R²) between SNPs in *Stolothrissa* (A-B) and *Limnothrissa* (C-D), demonstrating that loci associated with sex differences (light gray distributions) are more tightly linked than expected based on linkage values for all loci in the species-specific data sets (dark gray distributions). Panels (C) and (D) have been truncated at R2 = 0.2 to better visualize the distribution of LD values for sex-associated SNPs.

Supplemental Figures

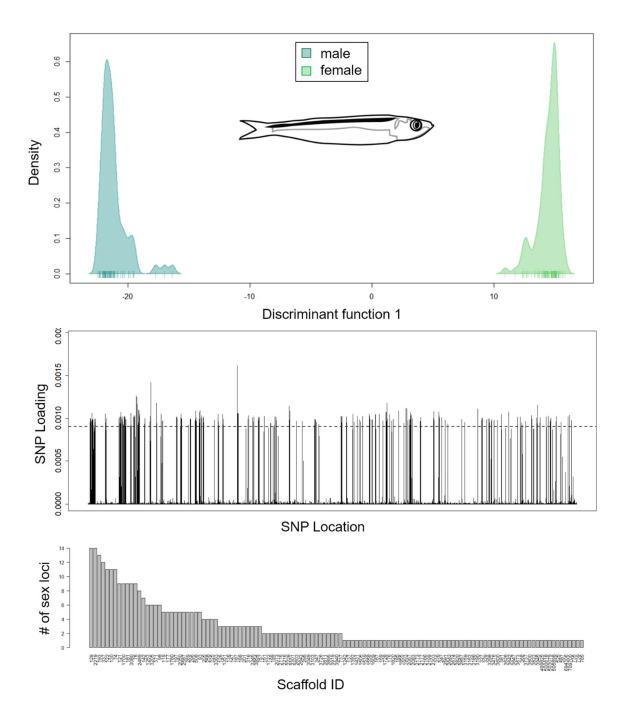


Figure S1. DAPC results for *Stolothrissa* differentiation between sexes, showing distribution of individuals on discriminant axis 1 (top), loadings for all SNPs (middle), and distribution of the 369 significant SNPs (loading > 0.0009) across scaffolds.

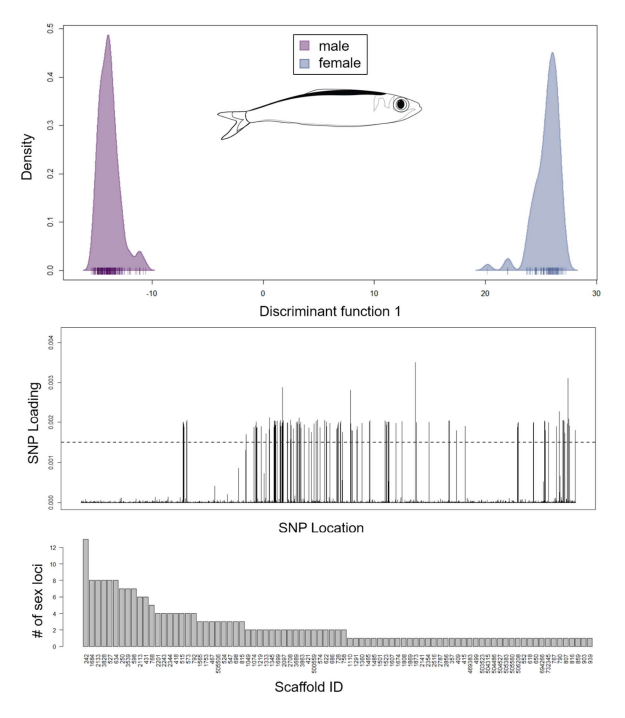


Figure S2. DAPC results for *Limnothrissa* differentiation between sexes, showing distribution of individuals on discriminant axis 1 (top), loadings for all SNPs (middle), and distribution of the 218 significant SNPs (loading > 0.0016) across scaffolds.

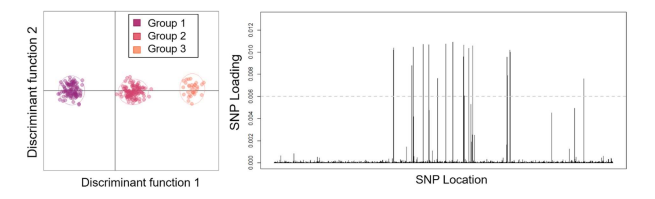


Figure S3. DAPC results for *Limnothrissa* differentiation between groups, showing distribution of locations on discriminant axis 1 and 2 (left), and loadings for all SNPs (right). We used a cutoff of 0.006 to identify SNPs with a significant loading on the differences between groups 1 and 3, resulting in a total of 25 significant SNPs.

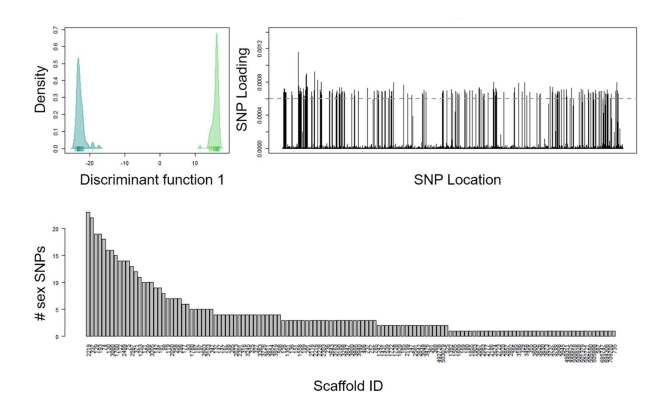


Figure S4. DAPC results for sex differentiation in *Stolothrissa* individuals using the combined species SNP dataset.

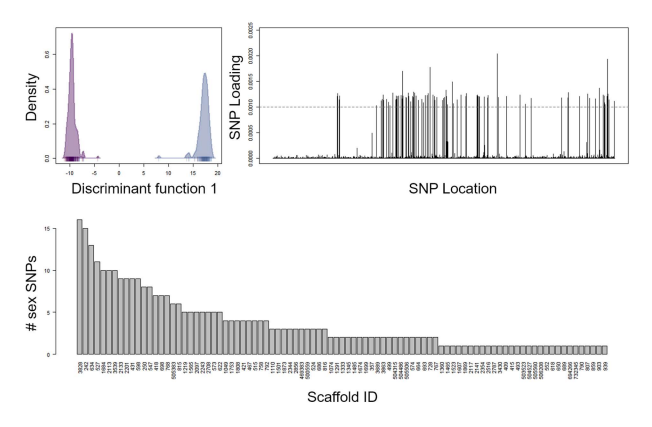


Figure S5. DAPC results for sex differentiation in *Limnothrissa* individuals using the combined species SNP dataset.

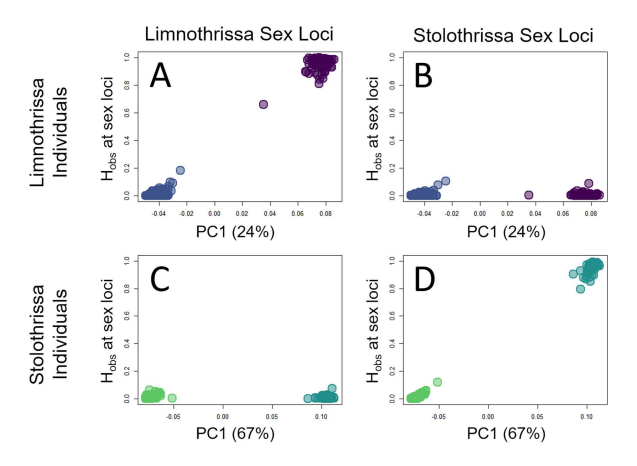


Figure S6. Observed heterozygosity of *Limnothrissa* individuals (A,B) and *Stolothrissa* individuals (C,D) at significant sex loci identified in *Limnothrissa* (A,C) and *Stolothrissa* (B,D), plotted against the first PC-axis for the species that the individuals belong to. Points are colored by genetically-identified sex, and PCAs were conducted on each species separately using the set of SNPs identified in the species-combined data set.

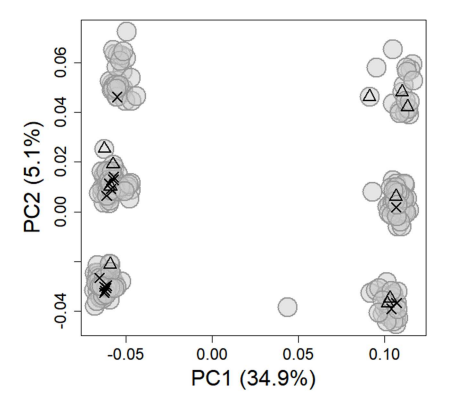


Figure S7. PCA of *Limnothrissa* individuals, highlighting a group of juvenile *Limnothrissa* (< 3cm) caught from the same school in Sibwesa (South Mahale) in 2015 (black X's), and nine Limnothrissa fry (< 2cm) caught in one scoop with a hand net in Kagunga in 2017 (black triangles). Both single-school samples included individuals from multiple different karyotypes.

Table S1 Sex determination in Stolothrissa and Limnothrissa

othrissa	
ple ID G	Phenotypic sex
863.IKO02 fe	male
864.IKO03 n	female
866.IKO05 fe	female
867.IKO06 fe	female
368.IKO07 n	female
869.IKO08 n	female
870.IKO09 fe	female
872.IKO11 n	female
873.IKO12 fe	female
874.IKO13 fe	male
883.IKO22 n	male
889.IKO28 n	female
217.KAG51 fe	female
219.KAG53 fe	female
	female
	female
	female
	male
	female
	female
	male
	male
	male
	female
	male
	male
	female
	female
	female
	male
	female
	male
	male
	female
	male
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