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Population genomics of introduced Nile tilapia Oreochromis niloticus (Linnaeus, 1758) in the Democratic Republic of the Congo: Repeated introductions since colonial times with multiple sources

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- 1 Population genomics of introduced Nile tilapia (Oreochromis niloticus
- 2 (Linnaeus, 1758)) in the Democratic Republic of the Congo: repeated
- 3 introductions since colonial times with multiple sources

5 Introduced Nile tilapia in the Congo Basin (running title)

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Abstract

During colonial times, Nile tilapia *Oreochromis niloticus* (Linnaeus, 1758) was introduced into non-native parts of the Congo Basin (Democratic Republic of the Congo, DRC) for the first time. Currently, it is the most farmed cichlid in the DRC, and is present throughout the Congo Basin. Although Nile tilapia has been reported as an invasive species, documentation of historical introductions into this basin and its consequences are scant. Here, we study the genetic consequences of these introductions by genotyping 213 Nile tilapia from native and introduced regions, focussing on the Congo Basin. Additionally, 48 specimens from 16 other tilapia species were included to test for hybridisation. Using RAD sequencing (27,611 SNPs), we discovered genetic admixture with other tilapia species in several morphologically identified Nile tilapia from the Congo Basin, reflects their ability to interbreed and the potential threat they pose to the genetic integrity of native tilapias. Nile tilapia populations from the Upper Congo and those from the Middle-Lower Congo are strongly differentiated. The former show genetic similarity with Nile tilapia from the White Nile, while specimens from the Benue Basin and Lake Kariba are similar to Nile tilapia from the Middle-Lower Congo, suggesting independent introductions using different sources. We conclude that the presence of Nile tilapia in the Congo Basin results from independent introductions, reflecting the dynamic aquaculture history, and that their introduction probably leads to genetic interactions with native tilapias, which could lower their fitness. We therefore urge avoiding further introductions of Nile tilapia in non-native regions and to use native tilapias in future aquaculture efforts.

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Keywords

- 72 Invasive species, cichlid, RAD sequencing, genetic integrity, genetic structure, independent
- 73 introductions

Introduction

76 Aquaculture production is one of the fastest-growing food-producing sectors in the world 77 (FAO, 2020). Together with fisheries, it plays a significant role in reducing hunger, promoting 78 health, and reducing poverty by providing jobs and livelihood to millions of people (Dugan et 79 al., 2010; FAO, 2020). Many people in Africa, especially those living near major rivers (Congo, 80 Nile, and Niger rivers) and the Great Lakes (lakes Tanganyika, Victoria and Malawi), depend primarily on fish as a source of animal protein (Brummett et al., 2008; FAO, 2016; Satia, 81 82 2017). 83 Tilapias are, after carps, the world's most important group of freshwater species used in 84 aquaculture (Eknath & Hulata, 2009), and they also have been introduced for capture fisheries 85 and sportfishing (Trewavas, 1983; Welcomme, 1988). In this study, we use 'tilapia' to refer to 86 a paraphyletic species assemblage, composed of several tribes (Dunz & Schliewen, 2013), 87 belonging to the so-called haplotilapiine lineage within the cichlids (Teleostei: Cichliformes: 88 Cichlidae). The most commonly farmed tilapia species is Nile tilapia, Oreochromis niloticus 89 (Linnaeus, 1758), which belongs to the tribe Oreochromini (Dunz & Schliewen, 2013; FAO, 90 2016; Lind et al., 2012) and comprises eight subspecies that are recognised based on 91 morphological characteristics, biogeography, behaviour, development, feeding, and analysis of 92 partial mitochondrial DNA sequences (Seyoum & Kornfield, 1992; Trewavas, 1983) (Table 1; 93 Figure 1). Nile tilapia is native to 22 countries, and its natural distribution roughly comprises 94 the Nile Basin, several river basins in West Africa (Senegal, Gambia, Volta, Niger, Benue and 95 Chad), various waterbodies of the East African Rift Valley (lakes Albert, Edward, Kivu, 96 Baringo, Turkana, some shallow parts of Lake Tanganyika, and the Omo and Suguta Basins), 97 Lake Tana in Ethiopia, and the Yarkon Basin in Israel (Trewavas, 1983) (Table 1; Figure 1). 98 However, because of worldwide introductions, both deliberate through stocking and

unintentional through aquaculture escapees (Welcomme, 1988), its presence is now reported in 105 countries (Froese & Pauly, 2021). The popularity of Nile tilapia in aquaculture stems from its fast growth and reproductive rate, and its ability to feed at a range of trophic levels and being tolerant to a range of environmental conditions (Canonico et al., 2005; Philippart & Ruwet, 1982; Zengeya et al., 2012). However, these same characteristics predispose it to be a successful invasive species (Canonico et al., 2005; Trewavas, 1983; Welcomme, 1988; Zengeya et al., 2012). Farmed fish can escape from aquaculture systems, establish themselves in local waterbodies and form feral populations (Lind et al., 2012). Here, they can predate on eggs and small fish, compete with native fishes for food and habitat resources, and introduce aquatic pathogens and parasites (Canonico et al., 2005; Deines et al., 2016; Jorissen et al., 2020; Lind et al., 2012; Naylor et al., 2001; Welcomme, 1988). These processes can cause a decline in the population size of native fish species (including native tilapias), which indirectly results in the loss of genetic diversity. The introduction of Nile tilapia can also have a direct genetic impact on native tilapia populations through hybridisation, a process that is often exploited for aquaculture purposes (Bezault et al., 2012; Brummett et al., 2004; Brummett & Ponzoni, 2009; Wohlfarth & Hulata, 1981). Unintentional hybridisation between escaped Nile tilapia and native tilapia species is a major concern for the genetic integrity of the latter and can cause a reduction of their overall degree of adaptation or fitness (Brummett & Ponzoni, 2009; Lind et al., 2012; Shechonge et al., 2018). Several cases of hybridisation have been recorded in the wild. The introduction of *O. niloticus* has been linked to the decline of native tilapias through hybridisation in Lake Victoria (Balirwa, 1992; Goudswaard et al., 2002), the Limpopo River system (D'Amato et al., 2007; Firmat et al., 2013; Moralee et al., 2000), and the Kafue River (Deines et al., 2014). Aquaculture in sub-Saharan Africa is a relatively new activity and is characterised by fluctuations caused by political instabilities and civil wars (Brummett et al., 2008; Toguyeni,

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2004). Tilapia aquaculture probably originated during the Second World War in the region of Lubumbashi, in the province Haut-Katanga in the then Belgian Congo (now the Democratic Republic of the Congo (DRC)) (Charpy, 1954; Micha, 2013; Robert, 1976; Toguyeni, 2004), producing mainly the native species O. macrochir and Coptodon rendalli (Boulenger, 1897) (Huet, 1957, 1959; Micha, 2013; Thys van den Audenaerde, 1964; Toguyeni, 2004). These species were also imported from Haut-Katanga into the Republic of the Congo and into the Central African Republic (then part of French Equatorial Africa), where they were used in aquaculture in the Middle Congo (Charpy, 1954; Lemasson, 1958). After WWII, aquaculture production in the Upper Congo increased by the creation of several fry production centres. At this point, the main cultured species were native O. macrochir and C. rendalli, and introduced O. niloticus of unknown origin, with the latter outperforming the former two by the end of the 1950s (Micha, 2013; Thys van den Audenaerde, 1988). However, after the independence of the country in 1960, aquaculture activities encountered numerous negative impacts due to the hasty departure of the Belgian supervisory staff, lack of trained personnel, and remaining political unrest (Brummett et al., 2008; Toguyeni, 2004). However, since 1996, fish production restarted and has been growing since (Toguyeni, 2004). Currently, tilapia production in the DRC is the highest among the Central African countries (Satia, 2017). Its annual tilapia production increased from 2000 tons/year to 3000 tons/year between 2000 and 2010, with Nile tilapia being the most farmed, followed by O. macrochir and C. rendalli (El-Sayed, 2013). The Congo River Basin covers almost the entire area of the DRC and parts of its neighbouring countries (Runge, 2007; Snoeks et al., 2011), and is divided into three main sections: the Upper Congo running from its source until the Boyoma Falls, upstream of Kisangani; the Middle Congo running from these falls until Pool Malebo near Kinshasa; and the Lower Congo running from the outlet of Pool Malebo until its estuary in the Atlantic Ocean (Brummett et al., 2011; Roberts & Stewart, 1976; Runge, 2007) (Figure 2). Nile tilapia is naturally present only in a

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small part of the Congo Basin (in some shallow parts of Lake Tanganyika and Lake Kivu) (Thys van den Audenaerde, 1964) (Figure 1). However, due to its extensive (historical) introduction for aquaculture purposes, and the possible secondary unintentional dispersal of aquaculture escapees, it has established itself throughout the entire basin (Decru et al., 2017a; Kisekelwa et al., 2020; Lunkayilakio et al., 2010). In view of the well-documented negative effects that the introduction of Nile tilapia can have upon native species, it is paramount to identify and trace introductions. Moreover, regarding the current efforts being made to boost Nile tilapia aquaculture (Micha, 2013), it is important to understand the distribution of genetic diversity and structure of introduced Nile tilapia in the Congo Basin as genetic diversity is a critical indicator for the evolutionary potential of populations, an attribute that could be of great value for the management of aquaculture stocks (Lind et al., 2012). We aim to gain insight into the historical introduction of Nile tilapia in the Congo Basin and to assess possible genetic consequences on native tilapias and on introduced Nile tilapia itself. We use a RAD sequencing approach to study the genetic structure of Nile tilapia populations from the Upper, Middle, and Lower Congo Basin, including farmed as well as feral populations. We hypothesise that: (i) a certain degree of genetic admixture exists in introduced Nile tilapia due to their ability to interbreed with other tilapia species, (ii) feral populations have higher genetic variation than farmed populations as a result of mixing of escapees from different sources of farmed populations in combination with inbreeding and artificial selection under farmed conditions, (iii) several (independent) introductions took place using populations with different genetic backgrounds due to a turbulent aquaculture history characterised by political instabilities and civil wars, and (iv) Nile tilapia populations from the Upper, Middle, and Lower Congo Basin are genetically similar, i.e. that there is no genetic differentiation between populations from the different sections, since aquaculture was first developed in the Upper Congo (Lubumbashi), following transfer of specimens from the Upper

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to the Middle Congo, as already reported for *O. macrochir* and *C. rendalli* (Charpy, 1954;
Lemasson, 1958).

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Materials and Methods

Sample areas

In the present study, we focus on the part of the Congo Basin in the DRC, excluding Lakes Tanganyika and Kivu (Figure 2), and will refer to this area as the 'Congo River Basin' (CRB). When referring to the 'Upper' Congo, we intend the sections of the Congo Basin that fall within the CRB. A total of 272 samples, consisting of fins, (dorsal) muscles, spleen, and gills stored in 99% ethanol (v/v), were selected. Of these, 96 museum specimens were morphologically identified as O. niloticus, and originated from different locations in the CRB: 33 from the Upper, 29 from the Middle, and 34 specimens from the Lower Congo. These include specimens from fish farms and feral specimens from rivers and lakes (Figure 2). Additionally, 74 specimens of O. niloticus from its native range were included (Nile River, Senegal River, the Albertine Rift Valley (lakes Albert, Edward, George, Tanganyika, and Kivu, and the Ruzizi River), Lake Tana, Lake Hashenge, and the Benue Basin), and 43 specimens from regions where it has been introduced (China, Jordan, Madagascar, Uganda (Lake Victoria), Benin, Togo, and Zimbabwe) to infer the origins of introduced specimens (Table S2: Table S3)... Further, 48 specimens of other tilapia species, present in the collection of the Royal Museum of Central Africa (RMCA), were included to study possible hybridisation between introduced Nile tilapia and other (native) tilapia species: Oreochromis aureus (Steindachner, 1864), O. macrochir, O. andersonii, O. upembae (Thys van den Audenaerde 1964), O. leucostictus (Trewavas, 1933), O. salinicola (Poll, 1948), O. schewebischi (Sauvage, 1884), Coptodon zillii (Gervais, 1848), C. rendalli, C. congicus (Poll & Thys van den Audenaerde, 1960), Congolapia bilineata (Pellegrin, 1900), Tilapia sparrmanii Smith, 1840, T. ruweti (Poll & Thys van den Audenaerde, 1965), Sarotherodon melanotheron Rüppell, 1852, S. galilaeus (Linnaeus, 1758), and Pelmatochromis ocellifer Boulenger, 1899. Additionally, five morphologically identified hybrid specimens obtained through crossing of O. aureus and O. niloticus were included, caught in a natural ecosystem in Israel (escapees, or their descendants, from aquaculture facilities), and six morphologically identified hybrid specimens from the Upper Congo River (five between O. niloticus and O. macrochir, and one between O. niloticus, O. macrochir and/or C. rendalli) (Table 1; Table S4).

DNA extraction

Total genomic DNA was extracted from the samples using the DNeasy® Blood & Tissue Kit (Qiagen) following the manufacturer's instructions. The concentration of DNA extracted from each individual was quantified with a Qubit® 2.0 Fluometer (Life Technologies, Paisley (UK)).

RAD library preparation

Seventeen RAD libraries, each including 16 individuals, were prepared according to the protocol described in Baird et al. (2008) (Baird et al., 2008) and Etter et al. (2011) (Etter et al., 2011). First, the DNA of each individual was enzymatically digested with *SbfI-HF*® (NEB, cut site 5'-CCTGCA^GG-3'). A first adaptor, containing forward amplification and Illumina sequencing primer sites, was ligated to each digested DNA fragment. The uniquely barcoded samples were then pooled into multiplexed libraries, followed by random mechanical shearing with the Covaris® S220 Focused-ultrasonicator. Subsequently, fragments between 250 and 700 bp were selected using a BluePippinTM device (Sage Science, Beverly, MA, USA). Next, the DNA was ligated to a second adaptor with a unique barcode ensuring PCR amplification

and the identification of different libraries. RAD libraries were 101 bp paired-end sequenced on an Illumina platform with a HiSeq 4000 system at Macrogen Korea (Seoul, South Korea).

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SNP discovery and genotyping

The overall quality of the reads in each library was checked with the software FastQC version 0.11.7 (Andrews et al., 2011). Raw reads were processed using Stacks v2.3b (Catchen et al., 2011; Catchen et al., 2013). The intactness of the RAD cut site was checked and reads were demultiplexed using the *process radtags* module. Reads with a dubious RAD cut site or a low quality score were discarded with the filtering options '-r', '-c', and '-q'. Next, PCR clones were identified and discarded with the *clone filter* module. Using the *kmer filter* module, reads were filtered according to the number of abundant k-mers they contained with the filtering option '--abundant'. Reads were then mapped against the reference genome of O. niloticus (O niloticus UMD NMBU, accession number MKQE02000000 (Conte et al., 2017)) using the BWA-MEM algorithm of the software BWA version 0.7.17 (Li & Durbin, 2009). Next, a sequence dictionary was made with the same command line tools, and the reference sequence and BAM files were indexed with SAMtools version 1.7. For the actual SNP discovery and genotyping, the software GATK version 4.0.0.0 (McKenna et al., 2010) was used. With this software, local realignment was performed with the 'RealignmentTargetCreator' and 'IndelRealigner' option so that the number of mismatching bases was minimised across all the reads. Finally, SNPs were called with the 'UnifiedGenotyper' option using a Bayesian genotype likelihood model. The resulting VCF file was filtered using VCFtools version 0.1.13 (Danecek et al., 2011) to include only high-quality SNPs with the following parameters: only bi-allelic SNPs with a quality score above 30, and only SNPs that were successfully genotyped in 80% of the individuals. Only one SNP per RAD tag was kept to minimise linkage disequilibrium. To remove possible paralogues, sites characterised by heterozygosity excess (*q*-value <0.05) were discarded. The final dataset included 27,611 SNPs.

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Genetic structure

To investigate genetic population structure, the software STRUCTURE version 2.3.4 (Pritchard et al., 2000) was used. For each value of K (number of clusters) ranging from one to ten, ten iterations were run using the admixture model (generations = 20 000; burn-in = 10 000). The optimal number of clusters K was inferred in Structure Harvester version 0.6.94 (Earl & vonHoldt, 2012) by the LnP(K) and the derived delta K calculated by the method of Evanno et al. (2005) (Evanno et al., 2005). Because independent iterations resulted in different outcomes, the optimal alignment of the three iterations with the highest estimated log probability was determined using CLUMPP version 1.1.2 (Jakobsson & Rosenberg, 2007). For each cluster, the individual's membership coefficient values (Q-values) were estimated. Plots were visualised in DISTRUCT version 1.1 (Rosenberg, 2004). In case of a bimodal support for different K values, plots with both K values were visualized. Genetic structure was further assessed by performing a non-scaled, non-centred Principal Coordinate Analysis (PCoA) using the R package 'adegenet' version 2.1.3 (Jombart, 2008) in R version 4.1.0 (R Core Team, 2021). For these analyses, the Albertine Rift Valley was split into the northern Nilotic (lakes Albert, George, and Edward) and southern Congolese part (lakes Tanganyika and Kivu, and Ruzizi River). Populations from Benin and Togo were considered together as both were sampled in the Mono Basin. Populations from the Nile River (Egypt and Sudan) were considered separately because of the large geographical distance between them.

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Discovery of admixed Nile tilapias

Species identification based on morphology alone can be challenging (Blackwell et al., 2020; Bradbeer et al., 2019; Ciezarek et al., 2021; Rhymer & Simberloff, 1996). In order to detect the possible source of Nile tilapia populations from the Congo Basin, and to not overestimate the genetic diversity and structure in these populations, we performed some additional analyses to exclude possible hybrids or misidentified specimens from further analyses. We use the term 'admixture' to refer to genetic introgression resulting from interspecific crossings. First, an exploratory STRUCTURE analysis was performed, revealing some aberrant individuals with a high membership coefficient value to clusters Q3 and Q4 (when K = 4) (see Results). Based on these results, individuals were selected as 'potential hybrids or misidentified specimens' when their membership coefficient values (Q-values) to the minor clusters (Q3 and Q4) were above 5%. Next, SNPs with large allele frequency differences between the population of purebred O. niloticus (specimens of O. niloticus excluding the ones selected in the previous step) and each of the other tilapia species (Table 1) were identified using PLINK version 1.9 (Purcell et al., 2007). Subsequently, all specimens were assigned to hybrid classes based on the selected SNPs using the R package 'Hybrid index estimation' or 'HIest' version 2.0 (Fitzpatrick, 2012). Given the large number of markers included in this study (27,611 SNPs), the assignment of an individual to a certain hybrid class was considered reliable when the loglikelihood of the best-fit class was over two units greater than the log-likelihood of the second best-fit class and within two units of the maximum log-likelihood (Fitzpatrick, 2012). Class '1' is purebred Nile tilapia, class '2' is the purebred other tilapia species, class '3' is a F1 hybrid, class '4' is a F2 hybrid, class '5' is a backcross to Nile tilapia, and class '6' is a backcross to the other tilapia species. In total, 17 HIest analyses were performed, each time with O. niloticus and another tilapia species from Table 1 as parental species. Individuals that were significantly assigned to one of the hybrid classes (class 3, 4, 5, or 6) or purebred other tilapia species (class 2) were considered to be admixed. The same applies to those that were not assigned to the class

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for purebred Nile tilapia (class 1) in any of the tests. We checked the performance of the HIest analysis on six specimens from the Upper Congo that were morphologically identified as hybrids.

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Genetic diversity and differentiation

Pairwise F_{st} values were calculated between specimens from each of the geographical regions included in this study (Table S2) with Arlequin version 3.5 (Excoffier & Lischer, 2010), using 1,000 permutations. To account for multiple testing, FDR adjusted p-values were calculated using the Benjamini and Hochberg procedure (Benjamini & Hochberg, 1995) with the R package 'BiocManager' version 1.30.16 (Morgan, 2021) and a significance level of 0.05. To explore whether the genetic differentiation between specimens within the Congo Basin increases with geographical distance between them ('isolation by distance'), a Mantel test, implemented in GenAlEx version 6.5 (Peakall & Smouse, 2006, 2012) with 999 permutations was performed between the matrices of Euclidean genetic distances and hydrological distances. The shortest hydrological distance was measured between each locality with OGIS version 3.18.1 by mapping the locations on a river network, splitting the network into segments, and measuring the length of the segments between each pair of specimens. For the Mantel test, a subset of 8,098 SNPs was randomly sampled with the '--thin 80 000' option in VCFtools, and farmed specimens were excluded as they are not free to move. Additionally, basic population genetic parameters (mean number of individuals typed per locus per population (N), mean observed heterozygosity per locus (H_0), and mean expected heterozygosity per locus (H_e)) were calculated using the R package 'diveRsity' version 1.9.90 (Keenan et al., 2013) for all locations, and for farmed and feral specimens of the different sections of the Congo Basin. Mean allelic richness per locus (A) and mean private allelic

richness per locus (A_{pr}) were estimated using the rarefaction algorithm implemented in HP-Rare version 1.1 (Kalinowski, 2005).

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Results

Detection and exclusion of admixed specimens of Nile tilapia

The exploratory STRUCTURE analysis, including only specimens morphologically identified as Nile tilapia (n = 213), showed a bimodal K value: K = 2 (highest delta K value), and K = 4(highest mean LnP(K)) (Table S5) (Figure 3). The following results are based on the optimal number of four clusters (K = 4). Most individuals had high membership coefficients to clusters Q1 and Q2. Overall, the membership coefficient to cluster Q1 was higher for individuals from the Middle and Lower Congo Basin than for individuals from the Upper Congo Basin, which had a high membership coefficient to cluster Q2. A total of 86 individuals were identified as 'potentially admixed' based on the estimated membership coefficient to Q3 and Q4 (Table S6). The HIest test resulted in the selection of 39 specimens of admixed Nile tilapia, each of which had also been identified as admixed in the STRUCTURE analysis: six from the Upper Congo (all farmed specimens), seven from the Middle Congo (all feral specimens), five from the Lower Congo (two farmed and three feral specimens), four from the northern Albertine Rift Valley, eight from the southern Albertine Rift Valley, two from Madagascar, two from the Senegal Basin (Senegal), one from the Nile River (Sudan), one from Lake Victoria (Uganda), and three from Lake Kariba (Zimbabwe) (Table S7). In the HIest test, five specimens were significantly classified as a purebred 'other' tilapia species (class 2): one from the Middle Congo (MC MS 36 1) was classified as O. macrochir and another (MC ULI 39 1) as C. bilineata; one from the northern Albertine Rift Valley (UG GRG 32 5) as O. upembae; one from the southern Albertine Rift Valley (DRC NYA 35 6) as S. melanotheron; and one from Lake Kariba (ZIM KAR 25 2) as C. rendalli. One specimen from the Lower (LC INK 51 1) and one from the Middle Congo (MC_BOO_43_1) were classified as an F2 hybrid (class 4) between *O. niloticus* and *O. upembae*. Some specimens (e.g. UC_KAT_DEP_1_1) were significantly assigned to class 1 in different HIest analyses, using a different tilapia species as second parental species. This outcome means that these specimens were significantly assigned to the class of purebred Nile tilapia in these analyses. The HIest test with morphologically identified hybrids (Table S8) assigned one of the morphologically identified hybrids to the class of purebred Nile tilapia (class 1). Also, specimens of other tilapia species were not always assigned to the class of purebred other tilapia species (class 2) (Table S8).

Genetic structure of purebred Nile tilapia

PCoA and STRUCTURE analyses were performed without the admixed specimens of Nile tilapia identified with the HIest analysis. The STRUCTURE analysis including only purebred specimens from the CRB had a bimodal optimal number of clusters: K = 2 (highest delta K), and K = 6 (highest mean LnP(K)) (Table S5). In the plot (Figure 4a), individuals from the Upper Congo had an overall high membership fraction to cluster Q2 (when K = 2) and Q3 (when K = 6). Individuals from the Middle and Lower Congo Basin had a high membership fraction to Q1 (when K = 2) and Q6 (when K = 6). In the PCoA results, a clear geographical genetic clustering was visible (Figure 4b; Figure S1). The first three principal components explained respectively 13.9%, 6.6%, and 3.6% of the variation. Individuals from the Upper Congo Basin were separated from individuals from the Middle and Lower Congo Basin by the first principal component PCo1. Individuals from the Middle and Lower Congo Basin were separated mainly by the third principal component PCo3. Individuals on the positive side of PCo1 space (Figure 4b) had a higher membership fraction to Q2 than to Q1 in the STRUCTURE analysis (when K = 2) (Figure 4a), while individuals on the negative side of PCo1 space had a higher membership fraction to Q1 than to Q2 (when K = 2) (Figure 4a).

The STRUCTURE analysis including purebred Nile tilapia from all geographical regions also had a bimodal optimal number of clusters: K = 3 (highest delta K), and K = 5 (highest mean LnP(K)) (Table S5). In the PCoA including purebred Nile tilapia from all geographical regions, the first three principal components explained respectively 11.4%, 6.0%, and 4.7% of the variation. In both STRUCTURE (Figure 5a), as well as the PCoA (Figure 5b; Figure S2) analyses, most individuals from the Upper Congo Basin clustered together with most native Nile tilapia from the Nile River (Sudan) and introduced Nile tilapia from feral populations from Madagascar and China. Most individuals from the Middle and Lower Congo Basin clustered with native Nile tilapia from the Nile River (Egypt), the Benue Basin (Cameroon), and the southern Albertine Rift Valley (Ruzizi River: populations 37, 39, and 41 in Figure 5a), and with introduced Nile tilapia from feral populations in the Mono Basin (Benin), and Lake Kariba (Zimbabwe). Individuals from Lake Tana (Ethiopia) formed a small cluster separate from all other locations (Figure 5b) with a high membership fraction to cluster Q5 (when K = 5) in the STRUCTURE analysis (Figure 5a). The population from Lake Tana was genetically distinct from all other Nile tilapia populations based on the PCoA (Figure 5b) and the STRUCTURE analysis with K = 5 (Figure 5a).

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Genetic differentiation

Pairwise F_{st} analysis of purebred Nile tilapia from the CRB indicated that individuals from the respective sections of the Congo Basin were significantly differentiated from each other. Individuals from the Middle Congo were genetically most similar to those from the Lower Congo ($F_{st} = 0.049$), and individuals from the Upper Congo were most differentiated from individuals from the Lower Congo ($F_{st} = 0.161$) (Table 2 and S9). When considering other introduced and native Nile tilapias (Table S9), pairwise F_{st} values indicated that individuals from the Upper Congo Basin were not significantly differentiated

from individuals from Sudan ($F_{st} = 0.010$), and that the genetic differentiation with individuals from Lake Hashenge ($F_{st} = 0.044$), the Benue Basin ($F_{st} = 0.040$), Lake Victoria ($F_{st} = 0.045$), and China ($F_{st} = 0.056$) was significant, but relatively low (Table 2). Individuals from the Middle Congo and Lower Congo Basin were not significantly differentiated from individuals from the Benue Basin ($F_{st} = -0.009$ and $F_{st} = 0.0100$, respectively), and individuals from the Lower Congo were not significantly differentiated from individuals from Lake Kariba (F_{st} = -0.010) (Table 2). The genetic differentiation between individuals from the Middle Congo and the Lower Congo ($F_{st} = 0.04923$), Nile Basin in Egypt ($F_{st} = 0.017$), Mono Basin ($F_{st} = 0.016$), and Lake Kariba ($F_{st} = 0.041$) was significant although relatively low (Table 2). Finally, the genetic differentiation between individuals from the Lower Congo and the Nile Basin in Egypt $(F_{st} = 0.081)$ and Mono Basin $(F_{st} = 0.048)$ was significant, though, relatively low (Table 2). The Mantel test including purebred feral Nile tilapia from the CRB demonstrated no significant correlation between the Euclidean genetic distances and the hydrological distances (R = 0.157, p-value = 0.090) (Figure S3). When considering fish from each section of the basin separately, again no significant correlation was found in the Middle Congo (R = 0.247, p-value = 0.230) and Lower Congo (R = 0.075, p-value = 0.400) (Figure S3). A Mantel test for feral individuals from the Upper Congo was not performed, as there was a negligible hydrological distance between the individuals.

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Genetic diversity

When considering only purebred Nile tilapia from the CRB, no significant differences were found between the respective sections of the basin in terms of genetic diversity (Table 3). Also, within each section, no significant difference was found in the genetic diversity between farmed and feral populations. (Table 3; Figure S4). When considering all other introduced and

native Nile tilapia, no statistically significant differences were found between geographic regions (Table 4; Figure S5).

Discussion

Traditional morphometric and molecular markers used in previous studies (Agnèse et al., 1997; Bezault et al., 2011; Seyoum & Kornfield, 1992; Trewavas, 1983; Vreven et al., 1998) have low resolving power to unveil genetic differentiation within and between populations. Because of the high number of SNPs that can be identified, the rise of NGS techniques provides an efficient approach to increase resolution in population genomic studies and has already proven its value in the assessment of population structure and diversity in cultured and feral populations of Nile tilapia in Tanzania (Kajungiro et al., 2019). In the present study, 27,611 SNPs were derived from RAD-seq data to investigate: (i) whether introduced Nile tilapia suffered from genetic contamination from other tilapia species due to their ability to interbreed, (ii) whether feral populations have higher genetic variation than farmed populations as a result of mixing of escapees from different sources of farmed populations in combination with inbreeding and artificial selection in farmed conditions, (iii) whether one or rather several (independent) introductions took place in the Congo Basin using different genetic backgrounds, and (iv) the possible source(s) of historical introductions in this river basin.

Genetic contamination of introduced Nile tilapia

The classification of Nile tilapia into eight subspecies was contradicted by studies using morphometric, allozyme, restriction fragment length polymorphism data, and microsatellite data (Agnèse et al., 1997; Bezault et al., 2011; Rognon & Guyomard, 2003; Tibihika et al., 2020; Vreven et al., 1998). Still, these studies based on traditional markers gave some

inconsistent results, which suggests that these genetic markers have insufficient resolving power to characterise variation and/or mixing between phyletic lineages. In addition to the debated taxonomy of Nile tilapia subspecies, morphological identification of tilapias is challenging because divergence of phenotypic traits can be influenced by environmental factors (Hornsby et al., 2013; Tibihika et al., 2018; Wohlfarth & Hulata, 1981). In addition, morphological divergence can be induced by anthropogenic activities, e.g. the introduction of populations with different genetic backgrounds, followed by intraspecific admixture (Tibihika et al., 2018). Consequently, misidentifications in the field are inevitable. Additionally, several taxonomic issues are known. *Coptodon zillii*, for example, is sometimes used when referring to C. rendalli and vice versa (Wohlfarth & Hulata, 1981). Also, O. aureus in Israel has been misidentified in the past as O. niloticus (Wohlfarth & Hulata, 1981). Interspecific and intergeneric hybridisation between different species of tilapia make identification based on morphology alone even more complicated, particularly after several generations of backcrossing (Bezault et al., 2012; Brummett et al., 2004; Brummett & Ponzoni, 2009; Rhymer & Simberloff, 1996; Wohlfarth & Hulata, 1981). Moreover, introgression resulting from hybridisation is not always reflected in morphology or in traits that can be easily measured (Blackwell et al., 2020; Bradbeer et al., 2019; Ciezarek et al., 2021; Rhymer & Simberloff, 1996; Shechonge et al., 2018). By performing a HIest test on our RAD-seq data, we classified about 20% of the morphologically identified Nile tilapia as admixed. For the CRB, this was the case for six individuals from the Upper (all farmed), seven from the Middle (all feral), and five from the Lower Congo (two farmed specimens and three feral) (Table S7). The results from the HIest analysis suggest that most of the admixed specimens probably were backcrosses, implying the viability of these hybrids and ongoing introgression. The presence of this hybrid swarm can

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potentially have a negative effect on the native species through genetic swamping (Facon et al., 2005; Gibson et al., 2019; Hohenlohe et al., 2013; Todesco et al., 2016). Our findings should, however, be interpreted with caution due to some methodological limitations that could have influenced our results. First, we selected a specimen as potentially admixed when the membership coefficient values (Q-values) to the minor clusters (Q3 and Q4) in the STRUCTURE analysis were above 5%. This selection could have influenced the selection of SNPs that are divergent between the two parental species, and, subsequently, the assignment of individuals to hybrid classes based on these SNPs. Secondly, results of the HIest test that included specimens that were phenotypically identified as hybrids, assigned one specimen to the class of purebred Nile tilapia (Table S8). This may reflect a high intraspecific divergence of phenotypic traits. It may also indicate introgression that has been masked by several generations of backcrossing, resulting in a low membership coefficient fraction for the introgressed specimen and, consequently, resulting in the wrong hybrid class assignment in the HIest analysis (Ciezarek et al., 2021). Additionally, classifying individuals in a limited set of hybrid classes is not suitable after many generations of hybridisation and backcrossing (Fitzpatrick, 2012). Also, the outcome of the HIest test is considered credible only if the loglikelihood of the best-fit class was over two units greater than the log-likelihood of the second best-fit class and within two units of the maximum log-likelihood (Fitzpatrick, 2012). This is an arbitrary cut-off, which might influence which specimens we then consider to be admixed. Unfortunately, we were not able to identify the exact hybrid status and respective parental species of the admixed specimens. Not all candidate parental species present in the basin (e.g. Oreochromis lepidurus (Boulenger 1899), O. mortimeri (Trewavas 1966), O. mweruensis Trewavas 1983, O. spilurus (Günther 1894) (Froese & Pauly, 2021)) and aquaculture strains (and their parental species) were included in our analyses. Therefore, the presence of admixed specimens might have been underestimated. Also, some specimens of the parental species

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might not themselves be pure species or even be misidentified, given that they were not all significantly assigned to 'class 2' in the HIest analysis (Table S8).

Another factor underestimating the presence of hybrids is the fact that we focused on

Another factor underestimating the presence of hybrids is the fact that we focused on morphologically identified Nile tilapia, ignoring those with a deviating morphology. In order to understand the real impact of Nile tilapia introduction on native tilapias in the Congo Basin, future research is required to more accurately identify the parental species of hybrids, and the direction and extent of introgression. To reach this goal, more specimens phenotypically resembling the respective native species and specimens with deviating morphology should be included, as well as more pure native species and commonly used aquaculture strains.

Genetic diversity of populations in the Congo Basin

We hypothesised that Nile tilapia from the Upper Congo were the donor population for aquaculture in the Middle and Lower Congo Basin, as aquaculture in the Congo Basin was first developed in the Upper Congo, and because transfer of other tilapia species from the Upper Congo to the rest of the Congo Basin has already been reported in the past (Charpy, 1954; Lemasson, 1958). If this was the case, we would expect a higher genetic diversity in the Upper Congo and a lower diversity in populations from the Middle-Lower Congo Basin due to founder effects and similar genotypes of Nile tilapia populations from the different sections of the basin.

Contrary to our expectations, Nile tilapia populations from the three sections of the CRB, as well as Nile tilapia from the other sampled regions, are not significantly different in terms of genetic diversity, suggesting multiple introductions into the Middle and Lower Congo (Table 3; Table 4; Figure S4: Figure S5). We also hypothesised that feral populations would have higher genetic variation than farmed populations because of interbreeding of escapees from different sources of farmed populations in combination with inbreeding and artificial selection

under farmed conditions. However, such an outcome was not apparent in our results (Table 3; Figure S4). Similar results were found, for example, for freshwater bream Abramis brama (Linnaeus 1758) (Hosseinnia et al., 2014) and Eurasian perch *Perca fluviatilis* Linnaeus 1758 (Khadher et al., 2016), where the relatively high genetic diversity in farmed populations was ascribed to the swapping of broodstock between different farms and the regular introduction of wild individuals (Khadher et al., 2016). In case of Nile tilapia, the relatively high diversity of farmed populations could have resulted from multiple introductions from different sources, as reported, for example, in Lake Victoria (Balirwa, 1992). Interestingly, the overall levels of observed and expected heterozygosities in our study are considerably lower than in prior genetic studies on Nile tilapia (Angienda et al., 2011; Dias et al., 2016; Hassanien & Gilbey, 2005; Kajungiro et al., 2019; Lind et al., 2019; Mireku et al., 2017; Moses et al., 2020; Romana-Eguia et al., 2005; Rutten et al., 2004; Sukmanomon et al., 2012; Tibihika et al., 2019) (Table S10). As most aforementioned studies used microsatellites as genetic markers, a direct comparison with these studies is not appropriate, as SNPs are biallelic. However, when comparing our results with previous studies using SNPs, heterozygosity values in the present study were still remarkably low (Kajungiro et al., 2019; Lind et al., 2019; Moses et al., 2020). This difference could possibly be caused by the exclusion of admixed individuals in the present study and by recent genetic bottlenecks of feral and

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The use of several source populations

inbreeding of cultured populations.

No significant correlation was found between genetic and hydrological distances within the CRB (Figure S3). Results from the PCoA (Figure 4b, 7, S1, and S2), STRUCTURE analyses (Figure 4a and 5a), and pairwise F_{st} comparisons (Table 2) suggested a clear genetic split between populations from the Upper and Middle-Lower Congo, and a high genetic similarity

between populations of the Middle and Lower Congo. The presence of waterfalls between the Middle and Upper Congo Basin (Runge, 2007) could preclude upstream migration of Nile tilapia. Also, the well-developed social behaviour (i.e. non-random mating) and substrate affinity (i.e., male territorial guarding and female parental care) makes Nile tilapia a rather sedentary species, influencing population differentiation on a small geographical and temporal scale (Bezault et al., 2011). This behaviour could cause a genetic divergence between populations living at large geographical distances from each other. However, given the relatively short history of modern aquaculture in the Congo Basin, it is implausible that this mechanism has caused the observed population differentiation in the Congo Basin. Combining the results from the PCoA, the STRUCTURE analyses, the pairwise F_{st} comparisons, and the fact that there is no significant difference between the genetic diversity of the different sections of the Congo Basin, we suggest that Nile tilapia from the Upper Congo was not the main donor for aquaculture in the Middle-Lower Congo. The current genetic structure of Nile tilapia in the CRB can be explained by human-mediated gene flow in the form of independent introductions, using different sources in the Upper and in the Middle-Lower Congo. The genetic differentiation of the population from Lake Tana from all other populations was also found by Tibihika et al. (2020) (Tibihika et al., 2020), and supports its status as a separate subspecies, as suggested by Seyoum and Kornfield (1992) (Seyoum & Kornfield, 1992).

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Possible source(s) of Nile tilapia populations in the Congo Basin

Documentation about historical Nile tilapia introductions in the Congo Basin is scant: only a few introductions have been reported from Sudan to Brazzaville (Lower Congo, Republic of the Congo) (Froese & Pauly, 2021) and from the Bouaké station (Ivory Coast) to Brazzaville (Lower Congo, Republic of the Congo) and to Bangui (Middle Congo, Central African Republic) (Thys van den Audenaerde, 1988). Introduction from the Lake Edward/George

system into the Middle Congo Basin was proposed by Decru et al. (2017a, 2017b) (Decru et al., 2017a; Decru et al., 2017b), though, this introduction was not formally registered. In recent years, aquaculture in the Upper Congo has been influenced by aquaculture activities in Southern Africa, introducing several aquaculture strains believed to include improved strains, such as the 'GIFT' (Genetically Improved Farmed Tilapia) and 'Chitralada' strain. These introductions are probably facilitated by the border position of the Lubumbashi area and the less-restrictive Congolese aquaculture policy (pers. obs., A. Chocha Manda, University of Lubumbashi). But, here too, the exact origin of Nile tilapia introductions is undocumented. The results from the PCoA (Figure 5b and S2), STRUCTURE analyses (Figure 5a), and pairwise F_{st} analysis (Table 2 and S9) restricted to purebred Nile tilapia from native and introduced populations suggest some possible source populations for aquaculture in the Congo Basin (Figures 5, and S2; Table 2 and S9). Native Nile tilapias from the Nile Basin in Sudan are genetically similar to introduced Nile tilapia from the Upper Congo. Native Nile tilapia from the Benue Basin (Cameroon) are genetically similar to introduced Nile tilapia from the Middle and Lower Congo Basin. In addition, introduced Nile tilapias from Lake Kariba (Zimbabwe) are genetically similar to introduced Nile tilapia from the Lower Congo. Possible introductions from Sudan or the Ivory Coast to the Lower Congo Basin, from the Ivory Coast to the Middle Congo Basin, from the Lake Edward/George system to the Middle Congo Basin, or from Southern Africa to the Upper Congo Basin, could not be validated in the present study because of our limited dataset and limited documentation of transfers of aquaculture stock. We also found relatively low (though significant) genetic differentiation between specimens from the Upper Congo and native specimens from Lake Hashenge (Ethiopia) and the Benue Basin (Cameroon), and between specimens from the Upper Congo and introduced specimens from Lake Victoria (Uganda) and the Songtao and Gaozhou Reservoirs (China) (Table 2). In Lake Victoria, Nile tilapia was introduced from Lake Edward, from fish ponds in Kajjansi

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592 (Uganda) and Lake Turkana (Balirwa, 1992; Fuerst et al., 2000; Pullin & Capili, 1988). In 593 China, only one introduction of Nile tilapia from Sudan has been documented (Pullin & Capili, 594 1988). 595 Nile tilapia from the Middle and Lower Congo are genetically relatively similar to each other 596 and to native individuals from the Nile (Egypt) and Benue basins (Cameroon), and to 597 introduced individuals from the Mono Basin (Benin and Togo) and Lake Kariba (Zimbabwe) 598 (Table 2). In the Mono Basin, introductions have been documented from stations in Ivory Coast 599 and Burkina Faso (Lazard, 1990; Lederoun et al., 2018; Montcho et al., 2015). In Lake Kariba 600 (Zimbabwe), one introduction was documented from Nakambala Estate Farm in Zambia 601 (Marshall, 1988) and recent research demonstrated the use of several strains of Nile tilapia used 602 in aquaculture in Lake Kariba (Makeche et al., 2020). 603 Clearly, Nile tilapia in most countries originate from multiple introductions using different 604 populations, of which the native source is often unknown. Even within a section of the Congo 605 Basin, e.g. the Upper Congo, multiple strains are currently used (e.g. 'Kipopo' and 'Israel' 606 strain at farm Kipopo). Furthermore, little introduction events are documented. Therefore, from 607 the data we have now, we cannot reach reliable conclusions upon the exact source of the strains 608 being cultured in the Congo Basin. 609 Besides the poorly documented introductions of Nile tilapia, our study was limited by the lack 610 of well-defined genetically improved aquaculture strains that are popular in Nile tilapia 611 aquaculture and the parental species used to produce them (such as 'GIFT', 'Chitralada', 612 'Ghana', etc.). The inclusion of these strains could help clarify our results. Especially the 613 inclusion of Nile tilapia strains from the Bouaké station (Ivory Coast) would be interesting, as 614 introductions from this station have been reported in the Republic of the Congo and the Central 615 African Republic (Thys van den Audenaerde, 1988). Also, the inclusion of specimens from 616 aquaculture facilities in southern Africa could highlight their current role in aquaculture in the area of Lubumbashi. Additionally, we have considered Nile tilapia populations coming from the native regions to be native. However, given the worldwide transportation of genetically improved strains, and the high chance of escapees through pond flooding or floating cage breakages (Lind et al., 2012), we cannot exclude the possibility that individuals from the native region are already products of admixture with other strains of Nile tilapia or other tilapia species. To avoid this problem, historical, pre-aquaculture samples should be included in the analyses.

To conclude, our genetic results reflect the complex history of frequent and rather careless introduction and translocation events of Nile tilapia throughout the Congo Basin, without considering the genetic consequences that now emerge. Whilst introduced Nile tilapia dominates tilapia culture in the DRC (Toguyeni, 2004), several native tilapia species, such as *O. macrochir*, *S. galilaeus*, and *C. rendalli*, have a proven aquaculture potential (Lind et al., 2012). To reconcile conservation with the growing demand for fish, future initiatives should promote the use of native tilapias that are most suitable to local conditions and use these species as a genetic resource for potential breeding programs (Lind et al., 2012).

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Data Accessibility and Benefit-Sharing

Genotype data are available on DataDryad (doi:10.5061/dryad.sxksn035k).

Author Contributions

T.H., M.P.M.V., and T.A. supervised the study. T.H., M.P.M.V., T.A., M.G., C.V., G.S., M.W.P.J., and K.S., helped in the setup of the sampling design. M.P.M.V., A.C.M., C.D.M., S.L.W., J.M.O.F., Y.T., M.V.S., E.V., and J.S. contributed to the collection and morphological identification of fish, and provided scientific background information. M.G. conducted the lab work and genetical analyses. C.V., G.S. and T.H. helped in the interpretation and discussion of the results. The first draft of the manuscript was written by M.G. All authors critically revised the draft and approved the final manuscript.

Tables and Figures

Table 1Species and number of tilapia specimens other than Nile tilapia and its hybrids included in this study with their introduction state (native or introduced) and sampling location

Tilapia species or hybrid	Number of individuals	Sampling location	Native/Feral
Congolapia bilineata (Pellegrin, 1900)	3	Lefini River (Middle Congo), DRC	Native
Coptodon congicus (Poll & Thys van den Audenaerde, 1960)	5	Lindi, Lefini and Sangha River (Middle Congo), and Inkisi River (Lower Congo), DRC	Native
Coptodon rendalli (Boulenger 1897)	4	Lindi and Kasaï River (Middle Congo), Lake Kipopo (Upper Congo), DRC	Native
Coptodon zillii (Gervais, 1848)	3	Lake Edward, Uganda and Epulu River (Middle Congo), DRC	Feral in Lake Edward, native in Epulu River

		Kabwe and	
Oreochromis andersonii (Castelnau 1861)	3	Kapabi Swamp (Upper Congo), Zambia	Feral
Oreochromis aureus (Steindachner, 1864)	3	Oued Draa River, Morocco	Native
Oreochromis leucostictus (Trewavas, 1933)	3	Ruzizi River and Lake George, Uganda Lufira, Kiswishi,	Native
Oreochromis macrochir (Boulenger 1912)	5	and Kimbeimbe River, and Lake Kipopo (Upper Congo), DRC	Native
Oreochromis salinicola (Poll, 1948)	3	Kalombe and Kabunda River (Upper Congo), DRC	Native
Oreochromis schwebischi (Sauvage, 1884)	1	Nyanga River, Republic of the Congo Lake Kabwe,	Native
Oreochromis upembae (Thys van den Audenaerde 1964)	3	Lake Kabele, and Fungwe River (Upper Congo), DRC	Native
Pelmatochromis ocellifer Boulenger, 1899	1	Congo River (Middle Congo), DRC	Native
Sarotherodon galilaeus (Linnaeus, 1758)	3	Inkisi Basin (Lower Congo), DRC	Native
Sarotherodon melanotheron Rüppell, 1852	3	Mono Basin, Benin	Native
Tilapia ruweti (Poll & Thys van den Audenaerde, 1965)	2	Kasaï River (Middle Congo), DRC	Native
Tilapia sparrmanii Smith, 1840	3	Lufira and Fungwe River, Mulenda Lake (Upper Congo), DRC	Native
O. niloticus x O. aureus	5	Coastal Levant, Israel	Feral
O. niloticus x O. macrochir	5	Lake Kipopo and Bumaki farm(Upper Congo), DRC	Feral
O. niloticus x O. macrochir x C. rendalli	1	Lake Kipopo (Upper Congo), DRC	Feral

 $\begin{tabular}{l} \textbf{Table 2}\\ \textbf{Matrix of population differentiation based on pairwise F_{st} estimators between purebred Nile tilapia populations of the different sections of the CRB and purebred Nile tilapia from the other geographical regions. The number of individuals per regions is given between parentheses. \\ \end{tabular}$

	Upper Congo, CRB (DRC)	Middle Congo, CRB (DRC)	CRB (DRC)
Upper Congo, CRB (DRC) [†] (n = 27)	/	0.12523***	0.16182***
Middle Congo, CRB $(DRC)^{\dagger}$ $(n = 22)$	0.12523***	/	0.04923***
Lower Congo, CRB (DRC) † (n = 29)	0.16182***	0.04923***	/
Senegal Basin (Senegal) (n = 3)	0.11235***	0.10721**	0.20813**

Nile Basin (Egypt) (n = 6)	0.07574***	<u>0.01650*</u>	0.08123**
Nile Basin (Sudan) (n = 2)	<u>0.01017</u>	0.13771**	0.19954**
Northern Rift Valley (Uganda) (n = 11)	0.14693***	0.20364***	0.28431***
Southern Rift Valley (DRC, Burundi) (n = 20)	0.11821***	0.08655***	0.15415***
Lake Tana (Ethiopia) (n = 9)	0.34724***	0.42356***	0.45884***
Lake Hashenge (Ethiopia) (n = 4)	0.04440*	0.08569**	0.14422**
Jordan Basin (Jordan) † (n = 3)	0.07516***	0.13976**	0.20413**
Mono Basin (Benin, $Togo)^{\dagger}$ (n = 8)	0.09486***	0.01626*	0.04755*
Benue Basin (Cameroon) (n = 4)	0.03951**	<u>-0.00918</u>	0.00990
Betsiboka, Rianila, Sofia Basin (Madagascar)† (n = 10)	0.06324***	0.14395***	0.18445***
Lake Victoria (Uganda)† (n = 4)	0.04500**	0.07374**	0.12459**
Lake Kariba (Zimbabwe)† (n = 2)	0.17749**	0.04051*	<u>-0.00996</u>
Songtao and Gaozhou Reservoir (China) † (n = 10)	0.05552***	0.15872***	0.19895***

Geographical regions with a '†' indicate the regions where Nile tilapia has been introduced. The five lowest values are underlined for each section of the CRB. Significancy levels (FDR adjusted p-values<0.05) are indicated by an asterisk (*<0.05, **<0.01, and ***<0.001)

Table 3

Summary of genetic diversity among all purebred Nile tilapia in the Upper, Middle and Lower Congo (CRB) as well as among farmed and feral populations for these three sections of the CRB. Values are given as the mean per locus (and standard deviation) for the mean number of individuals typed per population (N), mean allelic richness (A), private allelic richness (A_{pr}), observed heterozygosity (H_0), and expected heterozygosity (H_0)

Geographical region	Farmed/Fer al	N	A	$\mathbf{A}_{\mathbf{pr}}$	H_o	H_{e}
Lower Congo	All	25,6981	1,0239	0,0042	0,0164	0,0235
(CRB)	All	(1,8238)	(0,0761)	(0,0165)	(0,0607)	(0,0746)
	Feral	6,9134	1,0337	0,0178	0,0165	0,0310
	relai	(1,5385)	(0,0999)	(0,0604)	(0,0655)	(0,0920)
	Farmed	18,7847	1,0198	0,0064	0,0165	0,0193
	ranned	(0,8098)	(0,0750)	(0,0266)	(0,0676)	(0,0731)
Middle Congo	All	19,8472	1,0269	0,0051	0,0223	0,0262
(CRB)	All	(1,6870)	(0.0808)	(0,0185)	(0,0704)	(0,0789)
	Feral	14,8812	1,0281	0,0116	0,0220	0,0271
	relai	(1,6474)	(0.0839)	(0,0335)	(0,0696)	(0.0809)
	Farmed	4,9660	1,0222	0,0082	0,0223	0,0199
	ranneu	(0,2229)	(0.0881)	(0.0411)	(0.0967)	(0,0792)
Upper Congo	All	26,7629	1,0259	0,0046	0,0243	0,0256
(CRB)	All	(0,7865)	(0.0803)	(0,0188)	(0,0790)	(0,0789)
	Feral	4,9549	1,0267	0,0116	0,0260	0,0240
	1.0141	(0,2385)	(0.0957)	(0,0492)	(0,1011)	(0,0861)
	Farmed	21,8080	1,0256	0,0107	0,0240	0,0250
	ranned	(0,7927)	(0,0805)	(0,0342)	(0,0787)	(0,0789)

Table 4

Summary of genetic diversity of all purebred Nile tilapia from all geographical regions. Values are given as the mean per locus (and standard deviation) for the mean number of individuals typed per population (N), mean allelic richness (A), private allelic richness (A_{pr}), observed heterozygosity (H_o), and expected heterozygosity (H_e)

Geographical region	Introduced/ Native	N	A	$\mathbf{A}_{\mathbf{pr}}$	$\mathbf{H}_{\mathbf{o}}$	H_e
Upper Congo, CRB (DRC)	Introduced	26.7629 (0.7865)	1.0259 (0.0803)	0.0046 (0.0188)	0.0243 (0.0790)	0.0256 (0.0789)
Middle Congo, CRB (DRC)	Introduced	19.8472 (1.6870)	1.0269 (0.0808)	0.0051 (0.0185)	0.0223 (0.0704)	0.0262 (0.0789)

Lower Congo, CRB (DRC)	Introduced	25.6981 (1.8238)	1.0239 (0.0761)	0.0042 (0.0165)	0.0164 (0.0607)	0.0235 (0.0746)
Senegal Basin (Senegal)	Native	2.8719 (0.3890)	1.0299 (0.1135)	0.0075 (0.0491)	0.0232 (0.1021)	0.0256 (0.0982)
Nile River (Egypt)	Native	5.2802 (0.8465)	1.0290 (0.0976)	0.0056 (0.0317)	0.0227 (0.0833)	0.0261 (0.0879)
Nile River (Sudan)	Native	1.9708 (0.1738)	1.0207 (0.1061)	0.0034 (0.0381)	0.0189 (0.1071)	0.0164 (0.0843)
Northern Rift Valley (Uganda)	Native	10.4329 (0.8615)	1.0264 (0.0815)	0.0064 (0.0260)	0.0232 (0.0773)	0.0254 (0.0779)
Southern Rift Valley (DRC, Burundi)	Native	18.3266 (1.5671)	1.0277 (0.0830)	0.0051 (0.0190)	0.0218 (0.0663)	0.0269 (0.0808)
Lake Tana (Ethiopia)	Native	8.5739 (1.0865)	1.0083 (0.0514)	0.0035 (0.0378)	0.0085 (0.0611)	0.0103 (0.0694)
Lake Hashenge (Ethiopia)	Native	1.6927 (0.8171)	1.0192 (0.1111)	0.0085 (0.0755)	0.0079 (0.0753)	0.0306 (0.1488)
Jordan Basin (Jordan)	Introduced	2.9201 (0.2976)	1.0280 (0.1096)	0.0066 (0.0461)	0.0261 (0.1130)	0.0234 (0.0917)
Mono Basin (Benin, Togo)	Introduced	6.7902 (1.1501)	1.0279 (0.0920)	0.0055 (0.0281)	0.0188 (0.0686)	0.0257 (0.0848)
Benue Basin (Cameroon)	Native	1.6270 (1.1250)	1.0165 (0.0961)	0.0060 (0.0559)	0.0099 (0.0747)	0.1963 (0.3897)
Betsiboka, Rianila, Sofia Basin (Madagascar)	Introduced	9.9021 (0.4381)	1.0252 (0.0854)	0.0048 (0.0262)	0.0227 (0.0834)	0.0242 (0.0814)
Lake Victoria (Uganda)	Introduced	3.9550 (0.2241)	1.0253 (0.0968)	0.0034 (0.0259)	0.0203 (0.0865)	0.0222 (0.0849)
Lake Kariba (Zimbabwe)	Introduced	1.9821 (0.1423)	1.0172 (0.0969)	0.0025 (0.0296)	0.0160 (0.1005)	0.0142 (0.0810)
Songtao and Gaozhou Reservoir (China)	Introduced	9.9351 (0.3563)	1.0234 (0.0815)	0.0043 (0.0232)	0.0229 (0.0846)	0.0225 (0.0777)

Figure 1

 Map of Africa with rivers and lakes in black, regions with natural occurence of Nile tilapia shaded in grey (based on Trewavas (1983) (Trewavas, 1983) and Bezault et al. (2011) (Bezault et al., 2011)), and sampling locations depicted as red dots (sampling locations in China not shown): **a.** Genetic clusters as identified by Bezault et al. (2011) (see Discussion section), framed region is expanded in **b.**, **b.** Geographical distribution of subspecies of *O. niloticus* following Trewavas (1983) and Seyoum and Kornfield (1992)

Figure 2

Map of Africa (top left) with the framed region expanded. Outline of the Congo Basin in orange and the part of the basin that we focus on in this study shaded in grey. Sampling locations of introduced Nile tilapia within this area are indicated as red dots. Boyoma Falls and Pool Malebo define the transition from the Upper Congo to the Middle Congo, and the Middle Congo to the Lower Congo, respectively. Numbers refer to the population identifiers in Table S2. Kinshasa, Kisangani and Lubumbashi indicated by green stars. Main rivers and lakes in black (shapefiles downloaded from Figure.landscapeportal.org, maps created using QGis 3.18.1 software)

Figure 3

Population structure plot resulting from individual-based clustering using STRUCTURE with the two optimal K values (K=2, K=4). All native and introduced specimens that were morphologically identified as Nile tilapia were considered in the analyses. Geographical regions are shown at the top. Each bar represents one individual, which is partitioned into as many as K coloured segments. The length of a coloured bar represents the estimated membership coefficient fraction (Q-values) in each of the K inferred clusters. Numbers at the bottom of the STRUCTURE plot represent the population identifiers as in Table S2

Figure 4

- 734 Visualisation of population structure including only purebred Nile tilapia from the CRB. a.
- 735 Individual-based clustering using STRUCTURE with the two optimal K values (K = 2, K =
- 6). Geographical regions are indicated at the top. Each bar represents one individual, and is 736
- 737 partitioned into as many as K coloured segments. The length of a coloured bar represents the
- 738 estimated membership coefficient fraction (Q-values) in each of the K inferred clusters.
- Numbers at the bottom of the STRUCTURE plot represent the population identifiers as in 739
- 740 Table S2. b. Genetic scatter plot of PCo1 versus PCo3 resulting from the PCoA. Each dot
- 741 represents one individual. Colours represent different geographical regions. Ellipses are
- 742 drawn at a confidence level of 0.95

745 Figure 5

- 746 Visualisation of population structure plot including purebred Nile tilapia from all native and 747 introduced populations. a. Individual-based clustering using STRUCTURE with the two
- 748 optimal K values (K = 3, K = 5). Geographical regions are indicated at the top. Each bar
- 749 represents one individual, which is partitioned into as many as K coloured segments. The length
- 750 of a coloured bar represents the estimated membership coefficient fraction (Q-values) in each
- 751 of the K inferred clusters. Numbers at the bottom of the STRUCTURE plot represent the
- population identifiers as in Table S2. **b.** Genetic scatter plot of PCo1 versus PCo3 of the PCoA. 752
- 753 Each dot represents one individual. Colours represent different geographical regions. Ellipses
- 754 are drawn at a confidence level of 0.95

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Additional files

- 757 Additional supporting information may be found online in the Supporting Information
- 758 section.

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