Limited hybridization between introduced and Critically Endangered indigenous tilapia fishes in northern Tanzania

Stephanie J. Bradbeer · Jack Harrington · Henry Watson · Abrahim Warraich · Asilatu Shechonge · Alan Smith · Rashid Tamatamah · Benjamin P. Ngatunga · George F. Turner · Martin J. Genner

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Abstract Hybridization between introduced and indigenous species can lead to loss of unique genetic resources and precipitate extinction. In Tanzania, the Nile tilapia (Oreochromis niloticus) and blue-spotted tilapia (Oreochromis leucostictus) have been widely introduced to non-native habitats for aquaculture and development of capture fisheries. Here, we aimed to quantify interspecific hybridization between these introduced species and the indigenous species Oreochromis esculentus, Oreochromis jipe and Oreochromis korogwe. In the Pangani basin, several hybrids were observed (O. niloticus × O. jipe, O. leucostictus × O. jipe, O. niloticus × O. korogwe), although hybrids were relatively uncommon within samples relative to purebreds. Hybrids between the native O. jipe × O. korogwe were also observed. In the Lake Victoria basin, no evidence of hybrids was found. Analysis of body shape using geometric morphometrics suggested that although purebreds could be discriminated from one another, hybrids could not be readily identified on body and head shape alone. These results provide the first evidence of hybridization between the introduced species and the Critically Endangered O. jipe in Tanzania. Given uncertainty regarding benefits of introduced species over large-bodied indigenous species in aquaculture and capture fisheries, we suggest that future

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introductions of hybridization-prone species should be carefully evaluated.

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**Introduction**

Introduced species are recognized as a major driver of biodiversity loss in aquatic environments (Clavero & García-Berthou, 2005), and many aquatic introduced species have been associated with substantial economic and ecological impacts (Lowe et al., 2000; Pimentel et al., 2005; Lowell et al., 2006). The spread of these species is predicted to continue as natural biogeographic barriers are overcome, either accidentally through unintended transport, or through deliberate introductions (Levine & D’Antonio, 2003; Taylor & Irwin 2004; Hulme et al., 2008; Westphal et al., 2008). Globally, there are few regions that have not been invaded by at least one introduced aquatic species (Molner et al., 2008; Leprieur et al., 2008), with global hotspots being highly correlated with human activity and trade (Drake & Lodge, 2003; Perrings et al., 2005; Meyerson & Mooney, 2007). Freshwater environments are considered especially vulnerable to invasion (Sala et al., 2000; Cox & Lima, 2006). A key concern is the inability to reverse invasions and the subsequent impacts, and only a few species have ever been successfully eradicated from an aquatic environment after establishment (Hill & Cichra, 2005; Williams & Grosholz, 2008; Leprieur et al., 2009; Hill & Sowards, 2015).

The range of ecological impacts associated with introduced species is extensive, including predation, competition and habitat alteration (Canonic et al., 2005). There is also growing concern surrounding the threat of the loss of indigenous unique genetic diversity through hybridization between native and introduced species, potentially even to the extent of species extinction (Levin et al., 1996; Rhymer & Simberloff, 1996). Hybridization is of particular concern in cases where one of the species are considered to be a threatened species. For example, in North America, vulnerable endemic Pecos pupfish *Cyprinodon pecosensis* (Echelle & Echelle 1978) have hybridized with the invasive sheepshead minnow *Cyprinodon variegatus* (Lacepède 1803) with potentially no pure populations remaining due to the apparent vigour of hybrid individuals (Rosenfield et al., 2004).

Hybridization has been invoked as a potential driver of biodiversity loss in the tilapiine cichlid fish of the genus *Oreochromis*. Several species have been introduced to non-native habitats in Africa with a view to developing and improving capture fisheries and aquaculture. Among the most widely distributed species is the Nile tilapia (*Oreochromis niloticus* L. 1758), native to the Nile basin and West Africa (Trewavas, 1983), but introduced into at least 15 African countries outside its native range (FAO, 2012). *Oreochromis* species are known for interspecific hybridization (Scribner et al., 2001), and hybrids are commonplace in aquaculture where they are selected for desirable characteristics such as salinity tolerance (Agresti et al., 2000; Kamal & Mair, 2005) and pigmentation (McAndrew et al., 1988; Romana-Eugia et al., 2004). In the natural environment, introduced *O. niloticus* has been documented as hybridizing with several species including *Oreochromis mossambicus* (Peters 1852) in South Africa (D’Amato et al., 2007), *Oreochromis aureus* (Steindachner 1864) in Egypt and West Africa (Rognon & Guymard, 2003; Bakhoum et al., 2009), *Oreochromis andersonii* (Castelnau 1861) and *Oreochromis macrochir* (Boulenger 1912) in Zambia (Deines et al., 2014), and *Oreochromis esculentus* (Graham 1928) and *Oreochromis leucostictus* (Trewavas 1933) in Kenya (Nyingi & Agne`se 2007; Angienda et al., 2011; Ndiwa et al., 2014). However, the underlying factors mediating levels of hybridization between *Oreochromis* are poorly understood.

In southern and eastern Africa, *O. niloticus* is now present in multiple major drainage systems where it was historically absent, including the Pangani, Rufiji, Ruvuma, Limpopo, Zambezi and Lake Victoria basins (Genner et al., 2013; Zengeya et al., unpublished data). This spread of *O. niloticus* has been accompanied by blue-spotted tilapia (*O. leucostictus*), native to Lakes Edward, George and Albert in the Nile system (Trewavas 1983; Shechonge et al. unpublished data). The consequences of introducing these species for native fauna in their new range are largely unknown (Deines et al., 2016). Concerns exist for both the conservation of native endangered populations (Gregg
et al., 1998; Moralee et al., 2000; Nyingi & Agnèse, 2007; Angienda et al., 2011) as well as the preservation of wild genetic resources for aquaculture purposes (Lind et al., 2012).

In this study, we focus on the extent of hybridization between native and introduced Oreochromis in the Lake Victoria and Pangani basins of northern Tanzania. Both basins are characterized by endemic species that are threatened by negative interactions, including hybridization with introduced O. niloticus and O. leucostictus. Specifically, in the Lake Victoria basin, the Critically Endangered endemic O. esculentus persists only in satellite water bodies of Lake Victoria, although it has been translocated to multiple locations in surrounding countries (Trewavas, 1983; Shechonge et al. unpublished data). In the Pangani system, the Critically Endangered endemic Oreochromis jipe (Lowe 1955) is distributed from Lake Jipe to the Pangani Falls dam where it co-occurs with the lowland native species Oreochromis korogwe (Lowe 1955) (Trewavas, 1983; Shechonge et al. unpublished data). Given an apparent threat to Critically Endangered species, here we assess the extent of hybridization with introduced Oreochromis species using genetic (microsatellite) markers. We also asked if genetically identified hybrids could be identified using geometric morphometric data that capture variation in external head and body shape, potentially informing future field survey work on hybrid abundance.

Methods

Sampling

We collected our focal Oreochromis samples from one site in the Lake Victoria catchment and four sites in the Pangani river drainage (Fig. 1; Table 1). Sampling took place in 2015 and 2016 using seine nets, gill nets and purchasing from artisanal fisherman. Samples of additional reference species were obtained from five other sites in Tanzania (Table 1), where the species were found alone, or co-existing with species without any morphological or genetic evidence of hybridization. Fish captured using netting were immediately euthanized on landing by an overdose of clove oil anaesthetic. Specimens were pinned out and imaged in the field prior to preservation in absolute ethanol. Whole fish were placed in 70% ethanol for long-term storage. Genetic samples (fin clips) were taken and preserved in absolute ethanol.

Microsatellite genotyping

DNA was extracted from fin tissues using the Promega Wizard DNA extraction kit. Samples were screened at 17 microsatellite loci (Supporting Information Table 1). PCR was performed in a volume of 10 μl PCR solution consisting of 1 μl DNA (~ 5 ng), 5 μl Mastermix and 4 μl primer mix (10 mM). PCR amplifications were conducted on BioRad MyCycler thermal cycler, with conditions consisting of one denaturation step of 15 min at 95°C, followed by 35 cycles of 30 s at 94°C, 90 s at 57°C and 1 min at 72°C, followed by a final extension step of 30 min at 60°C. PCR products were sized on an ABI 3500 automated sequencer against a LIZ 500 size standard using GeneMapper 4.1 (Applied Biosystems).

Microsatellite data analysis

All individuals were amplified at a minimum of 11 of the 17 microsatellite markers (Supporting Information Tables 2 & 3). Genetic diversity estimates, and tests of deviation from Hardy–Weinberg Equilibrium, were calculated in Arlequin 3.5 (Excoffier & Lischer, 2010). To estimate the genetic composition of individuals within sites, we used a two-step process. First we made an initial assignment of focal and reference individuals to species group using find clusters in the R package adegenet (Jombart & Ahmed, 2011), selecting the maximum number of Principal Components possible, and a K value reflecting the number of species present, based on their phenotypes. We then used this initial assignment to groups as prior (LOCPRIOR) in Structure 2.3.4 (Pritchard et al., 2000), selecting the admixture model, and 10 runs, each with a burn-in of 100,000 steps and 100,000 recorded iterations. Next, Clumpak (Kopelman et al., 2015) was used to summarize the Structure output. Individuals with an assignment probability lower than 0.9 to any one of the focal species were considered of hybrid origin.
Geometric morphometrics

The left-hand side of the specimens was photographed with a scale. Images were loaded into tpsDIG v.2.22 (Rohlf, 2005) and 21 landmarks were digitized (Fig. 2). Landmarks were chosen based on landmarks commonly used in morphometric studies (Genner et al., 2007). Shape was quantified using MorphoJ 1.06 (Klingenberg, 2011). In each analysis, a Procrustes superimposition was applied to landmarked data. To visualize shape differences among purebred individuals at the sites, we used a Canonical Variates Analysis in MorphoJ 1.06. To determine repeatability of landmarking, we landmarked a set of 24 randomly selected individuals a second time. Following Procrustes alignment in MorphoJ 1.06, coordinates were subjected to Procrustes Anova the R package Geo- morph (Adams et al., 2017) that revealed 54.85% of the total variance to be among individuals, 0.05% of the total variance between sets (original vs. repeat), and 45.10% representing residual variance.

To determine if hybrid individuals possessed intermediate morphology between parent species, we used a discriminant function analysis on procrustes scores in SPSS v.23.0 (IBM), using stepwise variable removal process that retained only the most informative variables. Genotypically “purebred” individuals were assigned to species groups, while putatively hybrid individuals were left unclassified. Individuals from Lake Kumba and Kerenge were pooled for morphological analyses due to their close geographic proximity and overlapping species composition.
Individuals from the other three sites were analysed separately.

Results

Microsatellite results

At each of the five sites, the complete samples showed significantly higher heterozygosity relative to expectations from Hardy–Weinberg equilibrium at most loci (Supporting Information Table 2). Analyses of these data supported the presence of *O. niloticus* at all five sites (Fig. 3), Lake Malimbe (*n* = 14), Kerenge (*n* = 30), Lake Kumba (*n* = 71), Nyumba-ya-Mungu (*n* = 14) and Pangani Falls Dam (*n* = 26). *O. leucostictus* was resolved as present at two sites, Lake Malimbe (*n* = 31) and Kerenge (*n* = 9). *O. esculentus* was present at two sites, Lake Malimbe (*n* = 31) and Nyumba-ya-Mungu (*n* = 3). *O. jipe* was present at three sites, Lake Kumba (*n* = 13), Nyumba-ya-Mungu (*n* = 18) and the Pangani Falls Dam (*n* = 9). *O. korogwe* was only found at the Pangani Falls Dam (*n* = 3). We found no evidence of hybrid individuals within Lake Malimbe or Lake Kumba. Evidence supporting the presence of hybrids was found at three

![Fig. 2](image)

Fig. 2 Landmarks used in geometric morphometric analyses

![Fig. 3](image)

Fig. 3 Posterior probabilities of assignment by structure to species groups of individuals collected at the five study sites, relative to the putatively purebred reference individuals. Each individual is represented by one vertical bar, with colours representing assignment probabilities to the species group. Reference individuals are underlined in black. Asterisk indicates putatively hybrid individuals
sites (Fig. 3). One hybrid individual was identified at Kerenge sample (O. leucostictus × O. jipe). Two O. niloticus × O. jipe hybrids were in the Nyumba-ya-Mungu sample, and eight hybrid individuals were found in the sample from Pangani Falls (one O. korogwe × O. niloticus, and seven O. korogwe × O. jipe).

Morphological characterization of purebred and hybrid individuals

Geometric morphometrics demonstrated significant differences in geometric morphometric space among purebred individuals (Table 2). Both CVA and discriminant analysis were consistent with sympatric populations being separable on shape variables (Figs. 4, 5), with evident differences primarily in body depth and eye size among species. In discriminant analyses, individuals identified as genetic hybrids were typically, but not exclusively, within the phenotypic space of parental purebred individuals (Fig. 5).

Discussion

We found no evidence of hybridization between O. esculentus, O. niloticus and O. leucostictus at any sites where pairs of these species co-occurred, including Lake Malimbe in the Lake Victoria catchment. Within the Pangani system, we found evidence for the presence of hybrid individuals of the Critically Endangered O. jipe with introduced species at two locations. At Kerenge one individual O. leucostictus × O. jipe was discovered, while at Nyumba-ya-Mungu two O. niloticus × O. jipe were found. Notably, at all these sites hybrid individuals between introduced and native species were uncommon relative to purebred individuals. The relatively low frequency of hybridization in some of our sites is consistent with evidence from other Oreochromis systems, such as introduced O. leucostictus and native O. niloticus in Kenya (Nyingi & Agnèse, 2007; Ndiwa et al., 2014). These patterns contrast with observations of extensive hybridization between introduced O. niloticus and native O. mossambicus in South Africa (D’Amato et al., 2007), and between introduced O. niloticus and native Oreochromis in Zambia (O. macrochir and O. andersonii; Deines et al., 2014).

The relative rarity of hybrids between introduced and indigenous species can perhaps be explained by strong prezygotic isolating mechanisms. Sexual selection acting on male traits such as breeding colour, courtship displays and spawning “bower” phenotypes have all been suggested to promote reproductive isolation in mouthbrooding cichlids (Seehausen et al., 1997, 2008). Additionally, there is a possibility that populations may mate assortatively due to separate breeding periods or different breeding habitat preference, but at present no information on the habitat choice of these species is available. Postzygotic mechanisms may also have contributed to an apparent absence of hybrid individuals in our samples, if hybrid individuals suffer from low viability or fertility relative to purebred individuals. The extent of viability and fertility among the Oreochromis in our study is uncertain, although we note that hybrids of several

### Table 2

| Site              | Function | Wilks’ λ | χ²     | df  | P       |
|-------------------|----------|----------|--------|-----|---------|
| Kerenge + Lake Kumba | 1 through 2 | 0.100    | 265.09 | 22  | < 0.001 |
|                   | 2        | 0.443    | 93.64  | 10  | < 0.001 |
| Nyumba-ya-Mungu   | 1 through 2 | 0.092    | 71.57  | 10  | < 0.001 |
|                   | 2        | 0.394    | 27.93  | 4   | < 0.001 |
| Pangani Falls dam | 1 through 2 | 0.099    | 69.38  | 10  | < 0.001 |
|                   | 2        | 0.717    | 9.97   | 4   | 0.041   |
| Lake Malimbe      | 1 through 2 | 0.014    | 212.88 | 16  | < 0.001 |
|                   | 2        | 0.201    | 79.39  | 7   | < 0.001 |
Oreochromis species have been generated within aquaculture producing viable and fertile offspring (Bartley et al., 2001). In principle, the extent of inviability will be dependent on the extent of genomic incompatibility linked to the timescale since divergence (Bolnick & Near, 2005; Stelkens et al., 2009). However, the fitness of hybrid fish will also be dependent on the specific phenotypes of hybrids relative to the parental lines within the local selective regime. For example, phenotypically intermediate hybrids of cyprinids (Nilsson et al., 2017) and cichlids (Maan et al., 2017) have been demonstrated to have reduced survival relative to parental forms, while hybrids of two centrarchid species have an inferior feeding performance relative to their parental species (McGee et al., 2015).

The evidence of hybridization between the sympatric native species *O. korogwe* and *O. jipe* was notable in the Pangani Falls Dam, constructed in 1994. Surveys have reported these species that are otherwise allopatric in their distributions, with *O. korogwe* being distributed in low altitude coastal stretches of the Pangani and Zigi rivers, while *O. jipe* is primarily a higher altitude inland species (Trewavas, 1983; Shechonge et al., unpublished data). Further work is needed to map the distributions of both species within the lower reaches of the Pangani river system. It is possible that there is a natural hybrid zone, but it is possible that hybridization has been promoted by the habitat modification either linked to the dam construction, or the presence of *O. niloticus*. Dam construction has resulted in hybridization in other freshwater fishes (Hasselman et al., 2014), while introduced species have been suggested to affect the natural reproductive behaviour of native fish species (Doupe et al., 2009), including driving the loss of unique genetic diversity though hybridization (Velema et al., 2012).

### Fig. 5

Discriminant function analysis of geometric morphometric shape variation among all individuals. Genetic purebreds were assigned to species, while hybrids were not preassigned. Individuals from Kerenge and Lake Kumba are grouped.

**Timescale of spread of non-native species**

An important factor determining the extent of hybridization and negative ecological effects on native fauna is the timescale of invasion. If hybridization is
dependent on the density of the invader, then evidence of hybridization may be absent or rare until the invader becomes established and passes a threshold density, resulting in a lag time between the appearance of the non-native species and production of hybrids (Crooks & Soule, 1999). *Oreochromis niloticus* and *O. leucostictus* are likely to have been first introduced into the Pangani system within the last 40 years. *Oreochromis niloticus* was notably absent from extensive surveys of Nyumba-ya-Mungu in 1974 (Bailey et al., 1978). At that time, *O. esculentus* was already established and abundant in the dam and forming an important part of the fishery. Given that *O. esculentus* had been reared in ponds in the lower Pangani near Korogwe as early as 1950 (Lowe-McConnell, 2006), it is possible that it arrived in the upper Pangani region prior to construction of the Nyumba-ya-Mungu dam in 1967–1969, and expanded in population size due to favourable conditions. Introductions of *O. leucostictus* and *O. niloticus* into Lake Victoria took place in 1953 (Pringle, 2005; Lowe-McConnell, 2006), but neither had been recorded in satellite Lake Malimbe as recently January 2003 (Kutanzi & Kishe, 2004). Given the relatively recent timeline of the arrival of introduced species, future monitoring of genetic structure in these habitats may provide evidence of how shifts in density and time affect the frequency of hybridization.

**Morphological evaluation of hybrids**

In natural systems, it can be possible to identify hybrid individuals on the basis of morphological characters. For example, F1 hybrids between the European cyprinids roach *Rutilus rutilus* (L. 1758) and bream *Abramis brama* (L. 1758) in Ireland can be identified using geometric morphometric analysis of body shape (Hayden et al., 2010), while hybrids of Atlantic salmon *Salmo salar* (L. 1758) and brown trout *Salmo trutta* (L. 1758) also typically exhibit intermediate morphology when measured using geometric morphometric approaches (Solem et al., 2014). We found that it was possible to separate purebred individuals of different species in sympatry using shape information, but hybrids overlapped in morphospace with parental individuals. Thus, we suggest that conclusive assignment of some *Oreochromis* hybrids may not be possible from geometric morphometric data of the gross body shape alone, although perhaps the method may be useful when used in combination with other phenotypic traits such as melanin patterning.

**Biodiversity and fisheries implications**

Hybridization can result in biodiversity loss through genetic swamping, where hybridization leads to the loss of unique genetic diversity, or demographic swamping, where the numeric increases in hybrids result in negative demographic consequences for the parental species (Todesco et al., 2016). The absence or low frequency of hybrids between introduced and indigenous species suggests neither of these scenarios are likely under current environmental regimes at our study sites. However, hybridization at low frequencies can lead to the introduction of novel alleles that either reduce fitness of native species (Muhlfield et al., 2009), or promote traits such as fast maturation and small body sizes that could compromise fish production. By contrast, hybridization can promote the sharing of beneficial alleles, reducing vulnerability to inbreeding and disease and facilitating increased niche width by both native and introduced populations (Hall, 2016; Pfennig et al., 2016). In principle, this can lead to net benefits to capture fisheries through improving survivorship. In *Oreochromis*, it is unclear if the presence of hybridization between native and non-native species has affected production in either capture fisheries or aquaculture, but these questions could be investigated through further work on the genomic composition of these *Oreochromis* communities, together with common-garden experiments investigating traits related to fish production.

**Concluding remarks**

Fisheries in East Africa are essential for local livelihoods and food security (Muir et al., 2005; Heck et al., 2007; Musaka & Musonda, 2013), and the introduced species *O. niloticus* and *O. leucostictus* are now important components of demersal fisheries in northern Tanzania (e.g. Kolding et al., 2014). However, any benefits from further spread of introduced species in East Africa must be weighed against potential risks to biodiversity, existing stocks, and future potential fisheries yields. We propose that fisheries managers adopt the precautionary principle, that suggests future aquaculture and capture fisheries development should be based primarily on indigenous large-bodied
species, unless there is compelling evidence that the economic and societal benefits will outweigh risks to biodiversity and existing artisanal fisheries. Further information on the likelihood of hybridization among species under different environmental conditions would help to guide policy and fisheries development in the region.

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