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# Phylogenomics reveals extensive introgression and a case of mito-nuclear discordance in the killifish genus *Kryptolebias*

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#### ABSTRACT

Introgression is a widespread evolutionary process leading to phylogenetic inconsistencies among distinct parts of the genomes, particularly between mitochondrial and nuclear-based phylogenetic reconstructions (e.g., mitonuclear discordances). Here, we used mtDNA and genome-wide nuclear sites to provide the first phylogenomicbased hypothesis on the evolutionary relationships within the killifish genus Kryptolebias. In addition, we tested for evidence of past introgression in the genus given the multiple reports of undergoing hybridization between its members. Our mtDNA phylogeny generally agreed with the relationships previously proposed for the genus. However, our reconstruction based on nuclear DNA revealed an unknown lineage - Kryptolebias sp. 'ESP' - as the sister group of the self-fertilizing mangrove killifishes, K. marmoratus and K. hermaphroditus. All individuals sequenced of Kryptolebias sp. (ESP) had the same mtDNA haplotype commonly observed in K. hermaphroditus, demonstrating a clear case of mito-nuclear discordance. Our analysis further confirmed extensive history of introgression between Kryptolebias sp. 'ESP' and K. hermaphroditus. Population genomics analyses indicate no current gene flow between the two lineages, despite their current sympatry and history of introgression. We also confirmed introgression between other species pairs in the genus that have been recently reported to form hybrid zones. Overall, our study provides a phylogenomic reconstruction covering most of the Kryptolebias species, reveals a new lineage hidden in a case of mito-nuclear discordance, and provides evidence of multiple events of ancestral introgression in the genus. These findings underscore the importance of investigating different genomic information in a phylogenetic framework, particularly in taxa where introgression is common as in the sexually diverse mangrove killifishes.

#### 1. Introduction

Estimating the evolutionary relationships among species is a crucial goal of evolutionary biology. With the unprecedented availability of large numbers of loci brought by the genomics era, it has become increasingly clear that organisms generally have a more complex evolutionary history than previously acknowledged, with biological processes such as recombination, incomplete lineage sorting, introgression, and genome rearrangements (Mallet et al., 2016; Nakhleh, 2013) contributing to different phylogenetic signals among topologies generated from different sets of data for the same group of organisms (Bravo et al., 2019).

Although phylogenetic incongruence may have appeared as a problem at first (Jeffroy et al., 2006; Maddison, 1997), evolutionary biologists now embrace heterogeneity of phylogenetic signals (Bravo et al., 2019; Hahn and Nakhleh, 2016), recognizing that phylogenetic incongruences offer a unique opportunity to investigate the biological phenomena underlying discordance. Among these, reticulate evolution through introgression is today commonly accepted as a widespread evolutionary process contributing to phylogenetic discordance (Bravo

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et al., 2019; Mallet, 2005; Nakhleh, 2013; Taylor and Larson, 2019). A striking example of how introgression can affect phylogenetic congruence is mito-nuclear discordance, which arises when phylogenetic reconstructions based on mitochondrial or nuclear loci for the same group of organisms substantially differ in their topologies (Bonnet et al., 2017). Although other biological factors (e.g., incomplete lineage sorting, selection on mtDNA, sex-biased dispersal) are also known to generate mito-nuclear discordances, introgression is commonly pointed out as a major source of mito-nuclear phylogenetic incongruences (Toews and Brelsford, 2012).

Differences in mating systems (i.e., defined as the proportion of selfing versus outcrossing in organisms with hermaphrodites (Barrett, 2014)) are expected to influence the extent and direction of hybridization (Pickup et al., 2019) and in the long-term the degree of introgression. For instance, prior selfing (i.e., eggs are self-fertilized before the window for outcrossing) is expected to provide a strong barrier for hybridization given the limited reproductive opportunity for outcrossing (Brys et al., 2016). The variety of mating systems (e.g., predominantly-selfing, mixed-mating, obligately outcrossing) found in the killifish genus *Kryptolebias* provides an ideal opportunity to investigate: i) how different mating systems may affect the extent of hybridization (Berbel-Filho et al., 2021); and ii) the role that introgression between lineages with different mating systems may have on phylogenetic congruence between mitochondrial and nuclear genomes.

*Kryptolebias* is a rivulid genus of killifishes (Order Cyprinodontiformes) (Costa, 2011b; Murphy et al., 1999; Thompson et al., 2021), currently composed of seven valid non-seasonal oviparous species (Costa, 2004; Costa, 2011a; Vermeulen and Hrbek, 2005). Previous phylogenetic analyses (based on mtDNA and/or few nuclear genes) proposed two distinct clades within *Kryptolebias*. The 'freshwater' clade

composed of narrowly distributed freshwater species living in shallow streams and pools in South America: *K. campelloi* (Costa 1990), *K. sepia* Vermeulen & Hrbek 2005, *K. gracilis* Costa 2007, *K. brasiliensis* (Valenciennes 1821). The second clade, known as the 'mangrove killifishes clade', is composed of three androdioecious species (i.e., populations consisting of males and hermaphrodites) living on mangrove forests along the tropical and subtropical western Atlantic basin: *K. marmoratus* (Poey 1880), *K. hermaphroditus* Costa 2011, and *K. ocellatus* (Hensel 1868)) (Berbel-Filho et al., 2020; Costa et al., 2010; Murphy et al., 1999; Tatarenkov et al., 2017; Tatarenkov et al., 2009; Vermeulen and Hrbek, 2005) (Fig. 1). *Kryptolebias marmoratus* and *K. hermaphroditus* are the only two known vertebrate species capable of self-fertilization (selfing) (Avise and Tatarenkov, 2015; Costa et al., 2010; Tatarenkov et al., 2009).

In the selfing mangrove killifish species (K. marmoratus and K. hermaphroditus), most of the eggs laid externally are already fertilized via selfing (Harrington, 1971; Lomax et al., 2017), leaving a limited window of opportunity for outcrossing either by intra or heterospecific males. Despite this expected limitation, recent studies identified cases of undergoing hybridization involving the selfing Kryptolebias species. Tatarenkov et al. (2018) (and later expanded by Tatarenkov et al. (2021)) reported hybridization between highly-selfing lineages of K. marmoratus and K. hermaphroditus 'Central clade' (a lineage closely related to K. hermaphroditus present in the southern portions of the Caribbean, Central America and northern South America, with taxonomic status still under debate (Lira et al., 2021; Tatarenkov et al., 2017)). In Southeast Brazil, two hybrid zones formed by interbreeding between K. hermaphroditus (predominantly-selfing, (Berbel-Filho et al., 2019)) and K. ocellatus (exclusively outcrossing, (Berbel-Filho et al., 2020)) represented the first known case of hybridization between



Fig. 1. Approximated geographic distribution of known *Kryptolebias* species and lineages. Geographic distributions for the species/lineages were on the literature (Berbel-Filho et al., 2020; Costa, 1990, 2004; Costa, 2007; Costa, 2006, 2016; Lira et al., 2021; Tatarenkov et al., 2017; Vermeulen and Hrbek, 2005) as well as online databases for sampling records (GBIF; https://www.gbif.org) and museum collections (CRIA - SpeciesLink; https://splink.cria.org.br/). Symbols next to species name represent species inhabiting mangrove (mangrove tree) or freshwater (blue spot) habitats. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

species with different mating systems in vertebrates (Berbel-Filho et al., 2021). These unlikely hybridization cases called for further research on the role of past introgression in the diversification of the genus *Kryptolebias*.

Although K. hermaphroditus populations are mostly composed of selfing hermaphrodites, outcrossing occasionally happens (Berbel-Filho et al., 2019), most likely involving rare males and hermaphrodites (Furness et al., 2015). Despite historical sampling, particularly in Southeast Brazil (Costa, 2011a), males of K. hermaphroditus were only reported recently (Berbel-Filho et al., 2016; Costa, 2016). Costa (2016) reported a relatively high frequency of K. hermaphroditus males (e.g., three out of 20 individuals) in a single population in the Brazilian state of Espírito Santo. In rivulids, the male color pattern is the most conspicuous character to diagnose species (Costa, 2009). This is particularly true for the mangrove killifish clade, in which hermaphrodites are remarkably similar morphologically (Costa, 2011a, 2016). Therefore, Costa (2016) argued that the color patterns observed (in two different 'morphs') in K. hermaphroditus males from this Espírito Santo and other locality in the Rio de Janeiro state represented an important diagnostic trait to the morphological identification of species in the group (Costa, 2009). However, the pattern of coloration of the *K*. hermaphroditus males reported by Costa (2016) differed substantially from the male color reported for K. hermaphroditus in Berbel-Filho et al. (2016) as well as other males reported for the species (Supplementary Figure S1; Amorim et al., 2022). Particularly the 'dark morph' (Costa, 2016), which exhibited a dark body flank with broad black margin along the whole caudal fin, while the 'light morph' and the other K. hermaphroditus males found in other populations exhibited an orange pattern of pigmentation along its body and often had faded black margins in the caudal fin (Supplementary Figure S1). The relatively high frequency of males, the presence of male two color morphs and the disparity in their coloration, together with the multiple evidence for hybridization in mangrove killifishes prompted further research on the identity and evolutionary history of the unusual Espírito Santo locality in Brazil.

The reports of undergoing hybridization (Berbel-Filho et al., 2021; Tatarenkov et al., 2018, 2021), as well as the recent advances in the knowledge of natural history and distribution of Kryptolebias species (Berbel-Filho et al., 2019; Berbel-Filho et al., 2020; Costa, 2016; Guimarães-Costa et al., 2017; Lira et al., 2021; Sarmento-Soares et al., 2014; Tatarenkov et al., 2017) highlight the need of an updated hypothesis regarding the evolutionary relationships within the genus *Kryptolebias.* Using a phylogenomic approach together with a higher number of loci and taxonomic sampling than previous phylogenetic reconstructions, our study aimed to provide the first phylogenomicbased hypothesis for the species relationships in Kryptolebias. In addition, we aimed to investigate the hypothesis that reticulation and past introgression events contributed to the diversification of Kryptolebias lineages. We reveal a previously unknown lineage/species with strong evidence of ancestral introgression hidden in a case of mito-nuclear discordance. Our findings highlight how the use of a phylogenomic approach can shed light on the phylogenetic history of groups with common history of interspecific hybridization and challenging taxonomy.

#### 2. Material and methods

#### 2.1. Mitochondrial DNA dataset

We generated a cytochrome oxidase 1 (*cox1*) dataset of 423 sequences from 50 sampling localities and five out of the seven species (with exception of *K. sepia* and *K. campelloi*) formally described as *Kryptolebias* species. Given the high nuclear divergence found in the Espírito Santo locality in the 'Southern clade' with nuclear data (see results below), we incorporated sequences for 18 individuals from this population generated here. Three additional *cox1* sequences for species in the 'freshwater' clade, namely *K. gracilis* and *K. brasiliensis* were also

generated here, while the remaining samples were extracted from previously published data. The samples processed for this study followed primers and PCR protocols described in Tatarenkov et al. (2017). Both forward and reverse DNA strands were sequenced and assembled using Geneious v. 9.1.8 (https://www.geneious.com). A detailed list of samples used in the mtDNA analyses is presented in Supplementary Table S1.

#### 2.2. Mitochondrial phylogeny and haplotype network

Our dataset containing 423 *Kryptolebias* individuals was reduced to unique 49 cox1 haplotypes of 591 bp. A cox1 sequence from *Atlantirivulus santensis* (Köhler, 1906) (GenBank accession number GU701924.1) was used as an outgroup for the phylogenetic reconstructions. We identified the best partition scheme and substitution models using ModelFinder in IQ-Tree2 v. 2.1.0 (Kalyaanamoorthy et al., 2017; Minh et al., 2020). We used the suggested partition scheme to infer a maximum likelihood reconstruction and inferred uncertainty with 1000 standard non-parametric bootstrap iterations. Given the evidence of mito-nuclear discordance within the selfing mangrove killifishes clade (see results below), we isolated the 27 haplotypes within this clade and used POPART v. 1.7 (https://popart.otago.ac.nz/) to generate a TCS haplotype network (Clement et al., 2002).

#### 2.3. Nuclear DNA dataset

We combined newly-generated and previously-published data to generate a nuclear DNA dataset across *Kryptolebias* species. First, we sampled populations of *K. ocellatus* (sensu Costa 2011), *K. hermaphroditus* (sensu Costa 2011), *K. brasiliensis* and *K. gracilis* during a field trip in Southeast Brazil between August and September 2017. We collected the fish using hand nets. *Kryptolebias ocellatus* and *K. hermaphroditus* are syntopic in their type-locality (GUA in Supplementary Fig. S3) (Berbel-Filho et al., 2020). Sampling was conducted under license ICMBio/SISBIO 57145-1/2017 and approved by Swansea University Ethics Committee reference SU-Ethics-Student-250717/245.

We build a genotype-by-sequencing library (GBS) for a total of 96 fin clips samples. DNA was extracted the using Qiagen DNeasy Blood and Tissue kit (Qiagen, Hilden, Germany) following the manufacturer's protocol. GBS libraries were prepared as described in Kitimu et al. (2015). In brief, extracted DNA was digested using the restriction enzymes EcoRI and HpaII and ligated to sequencing adapters. Those enzymes were selected based on successful sampling of many restriction sites in K. hermaphroditus populations (Berbel-Filho et al., 2019). An aliquot of 200 ng of genomic DNA were digested using a EcoRI (cutsite: GAATTC) and HpaII (cutsite: CCGG). Digested DNA was ligated to individually barcoded adapters with a HpaII cut site overhang and a common EcoRI Y adapter. Ligation products were individually cleaned to remove excess of adapters using Agencourt AMPure XP purification system (#A63880, Beckman Coulter, Brea, CA, USA) at a v/v ratio of 0.85 following the manufacturer's instructions. A single library was produced by pooling 20 ng of digested DNA from each restriction/ ligation product and amplified in eight separate PCR reactions which were pooled after amplification, size-selected (range 200-350 bp) and sequenced in a single lane of an Illumina NextSeq500 sequencer.

Out of the original 96 samples included in the library, 61 (36 *K. ocellatus* and 25 *K. hermaphroditus*) had been analysed in Berbel-Filho et al. (2021), while the remaining 35 (13 *K. gracilis*, 11 *K. brasiliensis* and 11 individuals from the Espírito Santo population) were generated specifically for the present study. We extracted 11 *K. hermaphroditus* and 9 *K. ocellatus* GBS samples from Guaratiba (type-locality for both species) previously published in Berbel-Filho et al. (2021). Furthermore, to expand our taxonomic sampling of *Kryptolebias* species, we incorporated raw whole-genome sequencing data for *K. marmoratus* individuals from Florida, Belize, Honduras, and San Salvador Island obtained by Lins et al. (2018). Additional raw whole-genome sequencing data for

localities from Guantanamo Bay in Cuba (from Lins et al. (2018)) and Panama (from Choi et al. (2020)), representing individuals from the 'Central clade' lineage were also included. These samples represent a lineage closely related to K. hermaphroditus present in the Greater Antilles, Lesser Antilles, southern Central America, and northern portions of South America (Lira et al., 2021; Tatarenkov et al., 2017). The formal taxonomic status of the 'Central clade' lineage (i.e., either as a distinct species or a differentiated lineage of K. hermaphroditus) is still under debate (Lira et al., 2021; Tatarenkov et al., 2021; Tatarenkov et al., 2017). To simultaneously highlight its proximity and divergence with K. hermaphroditus, we refer to mtDNA and SNPs data from individuals of the 'Central clade' as "K. hermaphroditus 'Central clade'" throughout the manuscript. Data from K. hermaphroditus individuals from Southeast Brazil is referred as "K. hermaphroditus 'Southern clade'" following the classifications in Lira et al. (2021) and Tatarenkov et al. (2017). Similarly to the mtDNA dataset, we were unable to get hold of samples from the freshwater species K. sepia and K. campelloi. Those species are only known from very limited geographical distributions in creeks in the Amazon Forest (Costa, 1990; Vermeulen and Hrbek, 2005) (Fig. 1). No further reports for neither of those species have been found since the sampling reported in the original description (2003 for K. sepia in Vermeulen et al., (2006); 1974 for K. campelloi in Costa (1990)). We also incorporated raw whole-genome sequencing data for Nematolebias whitei (Myers 1942) from Thompson et al., (2022), to be used as an outgroup in the phylogenetic reconstruction based on concatenated nuclear sites.

#### 2.3.1. Nuclear DNA data processing

We used GBSX v. 1.3 (Herten et al., 2015) to demultiplex the pairedend reads data from the GBS library allowing for one mismatch in the barcodes (-mb 1), no mismatch in the enzyme cut-site (-me 0) and ensuring that no common sequencing adapter was to be removed (-ca false). We then filtered (-qtrim r; -minlength 25) and merged the GBS reads by individuals using BBmap tools (Bushnell, 2014). All samples (both from GBS and whole-genome sequencing) were mapped to the assembled Kryptolebias hermaphroditus reference genome (Choi et al., 2020) using either BWA v. 0.7.17 (for the phylogenetic reconstruction – Dataset I) or Bowtie 2 v. 2.3.5 (for analyses within Kryptolebias - Datasets II to VIII) using default parameters (Langmead and Salzberg, 2012) and generated filtered and indexed individual BAM files using samtools v 1.10.0 (Li et al., 2009). The different aligners were used due to their different mapping algorithms. While Bowtie2 tends to have faster throughput, it does that at the expense of mapping a lower number of reads when compared to BWA (Hatem et al., 2013). This can dramatically decrease the number of sites shared across samples, especially when a distantly related sample is incorporated in the dataset (i.e., an outgroup). For this reason, we used BWA v. 07.17 (Li and Durbin, 2009) as the aligner in the dataset incorporating the Nematolebias whitei outgroup sample (Dataset I), while the remaining datasets including only Kryptolebias samples (Datasets II to VIII) had Bowtie2 v2.3.5 (Langmead and Salzberg, 2012) as an aligner. For the whole-genome samples extracted from the literature, we removed sequencing adapters using AdapterRemoval v. 2.2.2 (Schubert et al., 2016), mapped samples to K. hermaphroditus genome using either BWA v. 0.7.17 (for phylogenetic reconstruction - Dataset I) or Bowtie 2 v. 2.3.5 (for analysis within Kryptolebias - Datasets II to VIII) and filtered and indexed individual BAM files using samtools v. 1.10.0 within a pipeline in Paleomix v1.3.2 (Schubert et al., 2014). We limited our dataset to samples with  $\geq 500~k$ reads. These resulted in a dataset of 48 (out of 61) Kryptolebias samples. A detailed list of the samples and sampling localities is provided in Supplementary Table S2.

#### 2.3.2. Variant calling

For all datasets analysed (details provided in Table S4), we inferred genotypes using ANGSD v. 0.9.32 (Korneliussen et al., 2014). Due to the methylation sensitivity of HpaII, we constrained our variant calling to a

maximum of 5 % of missing data per loci across all individuals. We used ANGSD with the following parameters: minimum mapping quality (-minMapQ 30), minimum base quality (-minQ 20), missing data (-minInd 95 %), Global Depth (-setMaxDepth 600 \* number of individuals), minimum genotype posterior probability (-postCutoff 0.95), single and double-tons were accordingly removed based on minimum minor allele frequencies (-MinMaf), anomalous reads (-remove\_bads 1; SAM flag above 255), adjusted mapping quality for excessive mismatches (-C 50), performed BAQ computation (-baq 1), minimum coverage for genotype calling (-geno\_minDepth 3), use of SAMtools genotype likelihood model (-GL 1), and estimated posterior genotype probabilities assuming a uniform prior (-doPost 2). In addition, we used the ANGSD SNP calling method (-SNP\_pval 1e-6), where a Likelihood Ratio Test is used to compare between the null (maf = 0) and alternative (estimated maf) hypotheses by using a  $X^2$  distribution with one degree of freedom.

#### 2.4. Nuclear DNA phylogeny

Our first dataset (Dataset I) consisted of the full set of GBS sites (all sites recovered in our library passing the filtering scheme - both constant and variable) from 48 *Kryptolebias* individuals and *Nematolebias whitei* as an outgroup. This dataset consisted of 174,282 nuclear sites. We ran the ModelFinder algorithm (Kalyaanamoorthy et al., 2017) implemented in IQ-Tree2 v. 2.1.0 (Minh et al., 2020) to infer the best optimal substitution model for the concatenated dataset. Then, we ran IQ-Tree2 to infer the maximum likelihood (ML) tree for the concatenated alignment and to assess the support of internal branches using the Shimodaira-Hasegawa-like procedure support (SH-aLRT) (Guindon et al., 2010), the Bayesian-like transformation of SH-aLRT support (uFBoot) (Hoang et al., 2018) with 1,000 replicates.

To visualize the most common phylogenetic signal between *Kryptolebias* species while considering uncertainty that may derive from reticulation, we ran a NeighborNet analysis based on uncorrected p-distances among individuals from Dataset II (115,397 GBS sites across 48 *Kryptolebias* samples with coverage between 4.77X and 444.95X (mean 153.48X) and missing data per loci ranging from 0 % to 1.83 % (mean 0.25 %); Supplementary Table S3) was conducted in SplitsTree v. 4.18.2 (Huson and Bryant, 2006).

#### 2.5. Phylogenetic networks and ancestral introgression analysis

The bifurcating nature of phylogenetic trees may not accurately describe the phylogenetic history of a particular group, especially when introgression events are common (Olave and Meyer, 2020). Given the reduced representation nature of our GBS library and the high genetic divergence between the freshwater (*K. brasiliensis and K. gracilis*) and the remaining *Kryptolebias* species, we limited our introgression analysis to the species composing the 'mangrove killifishes clade,' namely *K. marmoratus, K. hermaphroditus* (Central and Southern clades), *Kryptolebias* sp. 'ESP' and *K. ocellatus*.

We used two approaches to assess reticulation and ancestral introgression events in *Kryptolebias*. First, to evaluate the incidence of reticulation events across the *Kryptolebias* phylogenetic tree, we used the *julia* package PhyloNetworks v. 0.14.2 (Solfs-Lemus et al., 2017). This package uses concordant factor tables to infer networks using pseudolikelihood under the multispecies network coalescent model. SNP-based concordant factors were inferred using the program SNPs2CFs v.1.4 (Olave and Meyer, 2020). SNPs2CFs requires phased and unliked SNPs data. We called SNPs for 33 individuals of the mangrove killifishes clade, following the parameters described in the variant calling section. This call resulted in a dataset containing a total of 9,532 SNPs (Dataset III). To phase the data, we limited our dataset to SNPs located only in the 24 chromosomes of the *K. hermaphroditus* reference genome, filtering out all SNPs located in unplaced scaffolds. To minimize linkage amongst SNPs, we further filtered out our dataset to SNPs separated by a minimum distance of five thousand base-pairs, resulting in a dataset containing 5,813 SNPs with an average distance of 110,809 base-pairs among SNPs (Dataset IV). We phased this dataset using Beagle v. 5.2 (Browning et al., 2021) and generated concordance factors using SNPs2CFs. The full number of quartets in our dataset is too large to be processed in PhyloNetworks (Solfs-Lemus et al., 2017). Therefore, we limited our sample to 1,000 alleles per species quartet (n.quartets = 1,000 on SNPs2CFs), resulting in a total 5,000 quartets. We estimated phylogenetic networks with a hmax (maximum number of hybridization



**Fig. 2.** Maximum-likelihood reconstruction for 49 unique cox1 haplotypes extracted from 423 *Kryptolebias* individuals. (a) Full tree containing the relationships among the 49 haplotypes with tip labels colored by species/lineages names. Tip labels show haplotypes and number of individuals sequenced (in parenthesis). (b) Haplotype network for the 27 cox1 haplotypes for individuals belonging to the selfing mangrove killifishes clade with their respective distribution. Details for all samples used for these analyses are provided in Table S1.

events) value ranging from zero to seven. Our starting network (hmax = 0) was represented by a concatenated ML tree ran on IQ-Tree2 using the full set of GBS loci (both constant and variable) for this dataset (Dataset V - 1,631,872 nuclear sites with no missing data: Supplementary Fig. S5). The resulting network for each hmax value was used for every subsequent run. We plotted the log-pseudolikelihood and selected the networks that resulted in substantial pseudolikelihood improvements.

To further evaluate the evidence of ancestral introgression in Kryptolebias, we used the software Dsuite v. 0.4 r28 (Malinsky et al., 2021) to calculate Patterson's D statistics (ABBA-BABA test) and f4-ratios (an estimate of admixture fraction). ABBA-BABA rely on comparisons between bi-allelic SNPs for four taxa (e.g., T1, T2, T3, O) which are related to each other by a rooted tree (e.g. (((T1, T2), T3), O)). 'A' and 'B' represents the ancestral and derived alleles, respectively. Under a no gene flow scenario, the patterns of ABBA (sharing of alleles between T2 and T3) and BABA (sharing of alleles between T1 and T3) are expected to occur with equal frequencies, while significant deviation from equal frequencies is consistent with introgression between T3 and either T2 (ABBA) or T1 (BABA). We formally tested for three possible past introgression events within the mangrove killifishes clades in Kryptolebias based on either previous or current evidence: i) between K. marmoratus and K. hermaphroditus 'Central clade' as suggested by Tatarenkov et al. (2018, 2021) with the following tree topology: (((Kher\_South, Kher\_-Central), Kmar), KspESP)); ii) between K. hermaphroditus 'Southern clade' and Kryptolebias sp. 'ESP' (see results) with the following tree topology (((Kmar, Kher\_South), KspESP), Koce)); and between (iii) K. hermaphroditus 'Southern clade' and K. ocellatus, given the ongoing hybridization found in Berbel-Filho et al. (2021) with the following tree topology: (((Kher\_South, KspESP), Koce), Kbra). 'Kmar', 'Kher\_Central', 'Kher\_South', 'KspESP', 'Koce', and 'Kbra' refer to K. marmoratus, K. hermaphroditus 'Central clade', K. hermaphroditus 'Southern clade', Kryptolebias sp. 'ESP', K. ocellatus, and K. brasiliensis, respectively. For the first two introgression tests (i and ii), we used Dataset III while for test iii we called SNPs for a dataset containing one representative per species (for maximize the number of sites given the inclusion of K. brasiliensis as an outgroup) of the mangrove killifish clade and an individual of K. brasiliensis as an outgroup (Dataset VI - 10,648 SNPs with no missing data).

#### 2.6. Genetic structure of the mangrove killifishes clade

Our results indicated (see below) the existence of a previously unknown lineage of Kryptolebias in a single coastal sampling site in of Espírito Santo State in Brazil (referred above as Kryptolebias sp. 'ESP') (Supplementary Fig. S4). We further explored the nuclear genomic structure Kryptolebias sp. 'ESP' in comparison to the other lineages in the mangrove killifishes clade (see Fig. 3) using Dataset III. To estimate individual ancestries, we used ngsAdmix v. 3.2 (Skotte et al., 2013) with K values ranging between 2 and 10 for 100 replicates using default parameters, except for tolerance for convergence (-tol  $1 \times 10^{-6}$ ), log likelihood difference in 50 iterations (-tolLike 50 1  $\times$  10<sup>-3</sup>), and a maximum number of EM iterations (-maxiter 10,000). We used StructureSelector (Li and Liu, 2018) to estimate the most likely number of genetic clusters. A pairwise genetic distance matrix between individuals matrix was computed directly from the genotype likelihoods using ngsDist v1.0.2 (Vieira et al., 2015) and was then used for Multidimensional Scaling (MDS) using the R function cmdscale. To calculate heterozygosity, we called the full set of GBS sites (both constant and variable) for a dataset containing all 33 individuals of the mangrove killifish clade (Dataset VII - 863,662 nuclear sites for 33 individuals with 5% of missing data). We used ANGSD to compute the unfolded global estimate of the Site Frequency Spectrum (SFS) using one individual of K. brasiliensis as the source of ancestral sequence. Heterozygosity was calculated as the proportion of heterozygous sites by the total number of sites per individual.

### 2.7. Introgression between Kryptolebias sp. 'ESP' and K. hermaphroditus 'Southern clade'

To gain further insights into the structure of the hybrid zone between *Kryptolebias* sp. 'ESP' lineage and *K. hermaphroditus* 'Southern clade' (as indicated in our results), we called SNPs for a dataset containing only *Kryptolebias* sp. 'ESP' and *K. hermaphroditus* 'Southern clade' individuals (Dataset VIII – 5,688 SNPs for 18 individuals with no missing data). With this dataset, we addressed the patterns of allele distribution (e.g., number of fixed and/or shared alleles) between the two lineages.

#### 3. Results

#### 3.1. Mitochondrial phylogeny

The phylogenetic reconstruction based on 49 unique *cox1* haplotypes extracted from 423 Kryptolebias individuals was largely consistent with previously suggested phylogenetic relationships in the genus (Berbel-Filho et al., 2020; Costa, 2004; Costa, 2007; Costa et al., 2010; Kanamori et al., 2016; Murphy et al., 1999; Tatarenkov et al., 2017; Tatarenkov et al., 2009; Vermeulen and Hrbek, 2005) (Fig. 2a). The freshwater species K. gracilis and K. brasiliensis formed a clade which is sister group of the mangrove killifishes clade, composed of K. ocellatus, K. hermaphroditus and K. marmoratus. The latter two formed a wellsupported clade within the mangrove killifishes clade (the selfing mangrove killifishes). As previously indicated, there were two K. hermaphroditus mtDNA clades, one comprising samples from San Salvador Island, the Caribbean and northern portions of South America (the 'Central clade' in Tatarenkov et al., 2017; Lira et al., 2021), and another composed of samples from Northeast and Southeast Brazil (the 'Southern clade' in Tatarenkov et al., 2017; Lira et al., 2021). All the 23 individuals from the Espírito Santo locality exhibited a single cox1 haplotype (Hap22, Fig. 2), which is widespread in many K. hermaphroditus populations along approximately 2,600 km of the Northeast and Southeast regions of the Brazilian coast (Lira et al., 2021) (Fig. 2b).

#### 3.2. Nuclear phylogeny

Our ML phylogenetic reconstruction based on 174,282 nuclear DNA sites (Dataset I) from 48 Kryptolebias individuals was generally concordant with the phylogenetic relationships proposed in the mtDNA tree, with two main exceptions (Fig. 3b). Although the freshwater species K. gracilis and K. brasiliensis grouped together, samples from the former formed a monophyletic group within a non-monophyletic composed of K. brasiliensis samples. The other exception consisted of a previously unknown and well-supported clade containing five individuals from the Espírito Santo locality. This clade (hereafter named as Kryptolebias sp. 'ESP') formed a sister clade to the selfing mangrove killifishes, consisting of K. marmoratus and K. hermaphroditus (both Central and Southern clades). Two additional individuals from the Espírito Santo locality clearly belonged to K. hermaphroditus 'Southern clade', suggesting this population consisted of two sympatric species. All 23 individuals sequenced for mtDNA from Espírito Santo locality (including the five Kryptolebias sp. 'ESP' individuals) had the same mtDNA haplotype typically observed in K. hermaphroditus populations in Northeast and Southeast Brazil (Hap22, Fig. 2), representing a clear case of mitonuclear discordance in Kryptolebias.

Our phylogenetic network reconstruction using SplitsTree largely agreed with the lineages found in our mtDNA and ML phylogenetic reconstruction (Figs. 2a and 3b). However, it also indicated the highest levels of site tree discordance in *Kryptolebias* are within the mangrove killifish clade (Fig. 3c). This finding suggests events of introgression may have been common during the evolutionary history of this clade.



**Fig. 3.** Phylogenetic reconstructions of the genus *Kryptolebias* using IQ-Tree 2 v. 2.0.1. (a) Schematic representation of the maximum-likelihood phylogenetic tree based on 49 unique mtDNA cox1 haplotypes extracted from 423 *Kryptolebias* individuals. Node circles represent nonparametric bootstrap values = 100. The full mtDNA phylogenetic reconstruction is provided on Fig. 2. (b) Maximum-likelihood phylogenetic tree based on 174,842 GBS nuclear sites (Dataset I). Node circles represent SH-aLRT (%), aBayes, and ultrafast bootstrap (%) support values, respectively. Only nodes with high support (SH-aLRT  $\geq$  90, aBayes = 1, and ultrafast bootstrap > 90) are shown. Branch lengths are shown in substitutions per site. Intraspecific clades were collapsed to facilitate visualization. The full nuclear phylogenetic reconstruction is provided on Supplementary Fig. S4. (c) 95 % confidence phylogenetic network (Neighbor-Net) constructed using SplitsTree based on all sites from Dataset II. All species, with exception of *K. marmoratus* and *K. hermaphroditus* (represented by hermaphrodites), are represented in the figure by male individuals.

#### 3.3. Phylogenetic networks and ancestral introgression

Our PhyloNetworks analysis indicated that the largest improvement in pseudolikelihood scores across the number of reticulation events evaluated (ranging from 1 to 7) occurred between zero (the original tree) and one reticulation event (Fig. 4a). The network generated with one reticulation indicated ancestral introgression from *K. hermaphroditus* 'Southern clade' and *Kryptolebias* sp. 'ESP', with the latter as hybrid lineage inheriting 40 % of its genomic content from the former, despite the fact these two lineages are relatively far from each other in the phylogenetic tree (Fig. 4b).

Introgression events between K. hermaphroditus 'Central clade' and K. marmoratus (D = 0.22; Z-score = 8.71; F<sub>4</sub> ratio = 0.22; p < 0.001)

(Fig. 4c), between *K. hermaphroditus* 'Southern clade' and *Kryptolebias* sp. 'ESP' (D = 0.62; Z-score = 50.15;  $F_4$  ratio = 1.07; p < 0.001) (Fig. 4d) and between *K. hermaphroditus* 'Southern clade' and *K. ocellatus* (D = -0.76; Z-score = 62.77;  $F_4$  ratio = -0.13; p < 0.001) (Fig. 4e) were all confirmed by our ABBA-BABA test, revealing extensive introgression events in several *Kryptolebias* lineages.

#### 3.4. Genetic structure of the mangrove killifishes clade

Admixture analysis indicated the presence of five genetic clusters (Fig. 4a), each representing the mangrove killifish clade lineages recovered by the phylogenetic reconstruction based on nuclear sites (Fig. 2b). All four metrics generated by StructureSelector further



**Fig. 4.** Phylogenetic networks and introgression analysis in the mangrove killifishes clade. Genetic structure analysis for mangrove killifish species based on 9,532 SNPs (Dataset III). (a) Log-pseudolikelihood scores per number of reticulation events tested using PhyloNetworks v. 0.14.2 (Solfs-Lemus et al., 2017). The red dot indicates the network chosen based on a large improvement of the pseudolikelihood score. (b) Phylogenetic network with one reticulation event. Numbers indicate inheritance proportions in the hybrid lineage. (c-e) ABBA-BABA results for introgression tests between: (c) K. hermaphroditus 'Central clade' ('Kher Central') and K. marmoratus ('Kmar'); (d) K. hermaphroditus 'Southern clade' ('Kher South') and Kryptolebias sp. 'ESP' ('Ksp. ESP'); (e) K. hermaphroditus 'Southern clade' ('Kher South') and K. ocellatus ('Koce'). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article.)

suggested five as the most likely number of genetic clusters (Supplementary Fig. S7; all clusters shown in Supplementary Fig. S8). As also indicated by our phylogenetic reconstruction, the Espírito Santo population is composed of two highly different lineages at the nuclear genome, with two of the individuals sequenced belonging to K. hermaphroditus 'Southern clade', and the remaining five belonging to the previously unknown lineage Kryptolebias sp. 'ESP', despite the fact that all those individuals (and other 18 individuals sequenced from the same population) have the same mtDNA haplotype (Hap22 in Fig. 2) commonly found in individuals from the K. hermaphroditus 'Southern clade' in Northeast and Southeast Brazil. Our admixture analysis further indicated that these two lineages are quite differentiated from each other, with no evidence of current admixture (or early hybrid generation individuals) between them (Fig. 5a). This result can also be observed in our MDS analysis (Fig. 5b), in which the clusters representing the species (with exception of K. hermaphroditus Central and Southern clades highlighting the proximity between these two lineages) occupied different portions of the eigenspace, with Kryptolebias sp. 'ESP' and K. hermaphroditus occupying opposite sides of the first dimension of genetic distance variation. In terms of genetic diversity, Kryptolebias sp. 'ESP' individuals had in average 4.25x higher proportion of heterozygous sites (average: 0.45) than the outcrossing species K. ocellatus (average: 0.10), and 11.7x higher than the selfing (and sympatric) K. hermaphroditus 'Southern clade' (average: 0.04) (Fig. 5c). Possibly due to long-term generation of selfing and/or low-coverage nature of sequencing, K. marmoratus and K. hermaphroditus 'Central clade' individuals had an extremely low proportion of heterozygous sites (<0.001) across the GBS sites sampled in Dataset VII.

## 3.5. Introgression between Kryptolebias sp. 'ESP' and K. hermaphroditus 'South clade'

Out of 5,688 SNPs in Dataset VIII, 4,976 (87.48 %) are fixed and homozygous across all 13 *K. hermaphroditus* 'Southern clade' individuals, reflecting the highly selfing nature of the species. Out of those,

4,186 (84.12 %) SNPs are present in a heterozygous state in *Kryptolebias* sp. 'ESP', another strong indication that the genome of *K. hermaphroditus* 'Southern clade' introgressed into the genome of a previously unknown *Kryptolebias* lineage, resulting in *Kryptolebias* sp. 'ESP'. This latter lineage is highly heterozygous (4,714 out of 5,688 (82.87 %) SNPs are heterozygote). Further indication that an unknown and highly differentiated species was originally involved in the introgression with individuals of the *K. hermaphroditus* 'Southern clade' is the fact that 74.68 % (4,248 out of 5,688) of the SNPs contained alleles exclusive to individuals of *Kryptolebias* sp. 'ESP'.

#### 4. Discussion

Our study provided the first phylogenomic-based hypothesis for the phylogenetic relationships within killifish genus *Kryptolebias*, involving five out of the seven currently valid species. Our results (based on both mtDNA and nuclear markers) largely agreed with previously proposed phylogenetic relationships within the genus, comprising two major monophyletic groups: one grouping the freshwater fishes (*K. brasiliensis and K. gracilis*), while the other grouping the 'mangrove killifish clade', comprising *K. ocellatus*, *K. hermaphroditus* (Central and Southern clades) and *K. marmoratus* (Berbel-Filho et al., 2020; Costa et al., 2010; Murphy et al., 1999; Tatarenkov et al., 2017; Tatarenkov et al., 2009; Vermeulen and Hrbek, 2005). In addition, our results revealed an extensive history of introgression in *Kryptolebias*. Our findings revealed yet a highly differentiated (and previously unknown) lineage with elevated levels of heterozygosity and a history of admixture with the predominantly-selfing and sympatric *K. hermaphroditus*.

#### 4.1. Kryptolebias phylogenetic relationships and introgression

Previous attempts to reconstruct the phylogenetic relationships within *Kryptolebias* were based either exclusively on mtDNA (Vermeulen and Hrbek, 2005), and/or were exclusively focused on the 'mangrove killifish clade' (Berbel-Filho et al., 2020; Kanamori et al., 2016; Murphy



**Fig. 5.** Genetic structure plots for the lineages in the mangrove killifish clade. (a) Admixture plot for K = 5, indicated by StructureSelector (Li and Liu, 2018) as the most likely number of genetic clusters based on Dataset III (9,532 SNPs). (b) Multidimensional scaling plot based on the pairwise genetic distances between individuals extracted from Dataset III. (c) Proportion of heterozygous sites per individual based on the site frequency spectrum for Dataset VII (863,662 nuclear sites). For ease of visualization, species based on data extracted from whole-genome sequences (*K. marmoratus* and *K. hermaphroditus* 'Central clade') were omitted from the plot given a very low number of heterozygous sites (see Results). All plots follow the color scheme described in (b). Across all plots, individuals marked with asterisks represent K. hermaphroditus 'Southern clade' sympatric to Kryptolebias sp. 'ESP'.

et al., 1999; Tatarenkov et al., 2017; Tatarenkov et al., 2009; Weibel et al., 1999). Our phylogenetic reconstruction not only expanded the number of genomic loci used, but also widened the taxonomic sampling of Kryptolebias, particularly by including the freshwater species: K. gracilis and K. brasiliensis; representatives of recently uncovered lineages (K. hermaphroditus Central and Southern clades); and a lineage revealed by the present study (Kryptolebias sp. 'ESP'). Overall, our phylogenetic reconstruction generally agrees with the topologies previously proposed by Vermeulen and Hrbek (2005) based on mtDNA loci, where two monophyletic Kryptolebias clades were represented by species living in freshwater streams or brackish environments close to mangrove forests (the freshwater clade and mangrove killifishes clade, respectively). Although we have not sampled the Amazonian freshwater species K. sepia and K. campelloi, those species are thought to be closely related to K. brasiliensis (Costa, 1990; Vermeulen and Hrbek, 2005), which in our phylogeny grouped together with K. gracilis in the 'freshwater clade'. It is important to highlight that our K. brasiliensis samples have not formed a monophyletic clade within the freshwater clade of our nuclear phylogeny, with K. gracilis being nested within K. brasiliensis (Fig. 3b) Those two species are morphologically very similar, with K. brasiliensis being distributed in broader area along lowland streams and creeks in the state of Rio de Janeiro (Costa, 2007). Our study thus calls for further research on the taxonomic status of K. brasiliensis and K. gracilis, with the possibility of the former representing a species complex.

Our nuclear phylogeny revealed some differences from previous reconstructions within the mangrove killifishes clade. So far, all studies that tried to reconstruct the phylogenetic relationships within this clade, regardless of whether it was based on mtDNA (Berbel-Filho et al., 2020; Murphy et al., 1999; Tatarenkov et al., 2017; Tatarenkov et al., 2009; Vermeulen and Hrbek, 2005; Weibel et al., 1999) or nuclear markers (Kanamori et al., 2016), have supported the obligated outcrossing species K. ocellatus (Berbel-Filho et al., 2020) as the sister-species of the clade containing the selfing species (K. marmoratus and K. hermaphroditus). Our phylogenetic reconstruction revealed a clear case of mitonuclear discordance (see discussion below) including a previously unknown lineage more closely related to the selfing mangrove killifishes than to K. ocellatus. This change in topology may have implications for understanding the evolution of mating systems within the genus. As all the other known Kryptolebias species are dioecious and inhabit freshwater habitats, the classical view based on the phylogenetic mapping of reproductive traits in Kryptolebias suggested that synchronous hermaphroditism has emerged in the common ancestor of all mangrove killifish species (K. ocellatus, K. hermaphroditus and K. marmoratus), with the self-fertilization evolving later in the common ancestor of the sisterspecies K. hermaphroditus and K. marmoratus (Avise and Tatarenkov, 2015; Costa et al., 2010). However, the phylogenetic positioning of Kryptolebias sp. 'ESP' as the sister-group of the selfing species raises the discussion of whether self-fertilization may have evolved earlier in the genus. Notwithstanding, it is important to note that self-fertilization tends to reduce heterozygosity levels in half every generation (Avise, 2008). The fact that the individuals from Kryptolebias sp. 'ESP' examined here had over 11x higher level of heterozygosity when compared to K. hermaphroditus, together with the evidence of ancestral introgression from K. hermaphroditus 'Southern clade' into Kryptolebias sp. 'ESP', suggests that Kryptolebias sp. 'ESP' may not undergo self-fertilization. All non-male individuals captured in the sampling locality of Kryptolebias sp. 'ESP' (and previously by Costa, 2016) had a typical external appearance of hermaphrodites of the selfing mangrove killifishes, suggesting this may be another androdioecious, but not self-fertilizing, species in the genus, similarly to K. ocellatus (Berbel-Filho et al., 2020). Nonetheless, our limited sample size makes imperative the need for further life-history and behavioural evaluation of the mating system of Kryptolebias sp. 'ESP' individuals. Until then, the possibility that selffertilization may have evolved earlier (in the common ancestor between Kryptolebias sp. 'ESP', K. hermaphroditus, and K. marmoratus) in

Kryptolebias must be considered.

Another major goal of our study was to evaluate the possibility that reticulate evolution may have played a role in the diversification of the genus Kryptolebias. Our phylogenetic networks analyses showed complex history of reticulate evolution in the mangrove killifish clade (Fig. 3c), and revealed an ancient introgression event from K. hermaphroditus 'Southern clade' into Kryptolebias sp. 'ESP'. This reticulation event may explain the fact that all individuals of Kryptolebias sp. 'ESP' had high heterozygosity levels and the same mtDNA haplotype commonly found in K. hermaphroditus 'Southern clade' in Northeast Brazil. Contrary to the prediction that highly-selfing taxa (such as K. hermaphroditus) provide a low opportunity for hybridization and introgression (Pickup et al., 2019), this ancestral reticulation event suggests that hermaphrodites of the K. hermaphroditus lineage played the maternal role in introgression events with a previously unknown lineage, now evident in the genome of Kryptolebias sp. 'ESP'. Tatarenkov et al. (2021) found bi-directional hybridization between two highly selfing strains of K. marmoratus and K. hermaphroditus 'Central clade', while Berbel-Filho et al. (2021) also found a single backcross between an F1 individual and K. hermaphroditus 'Southern clade' in Southeast Brazil. Taken together, these results suggest that although rare, opportunities to outcross and hybridize with the selfing Kryptolebias may occasionally occur. Backcrossing between F1 individuals and males of the Kryptolebias sp. 'ESP' lineage may have then further contributed to the movement of genomic DNA from K. hermaphroditus 'Southern clade' into Kryptolebias sp. ESP. The fact that we only found individuals with K. hermaphroditus 'Southern clade' mtDNA hints at the possibility of local extinction of individuals with the mtDNA of Kryptolebias sp. 'ESP', however we acknowledge that testing of this hypothesis requires further sampling in the area. The question of whether 'pure' Kryptolebias sp. 'ESP' individuals still exist or evidence for this (possibly extinct) lineage can only be found in extant admixed populations with K. hermaphroditus is open to investigation. Thus far, the closest sampling site around the Espírito Santo locality (only 5 km apart from Coqueiral Beach, where Kryptolebias sp. 'ESP' was found) had mtDNA haplotypes and hermaphrodite appearance commonly found in K. hermaphroditus (Lira et al., 2021).

The single reticulation event found in our phylogenetic networks analysis seems to contradict the finding of multiple introgression events using site patterns counts (ABBA-BABA tests) found between Kryptole*bias* lineages of the mangrove killifishes clade. However, these two types of introgression tests tend to recover introgression events at different time scales. ABBA-BABA test assume that multiple substitutions at a particular site are rare or do not occur, as many substitutions at individual sites could affect the patterns of site discordance. This assumption tends to not hold true for deeply diverged taxa (Hibbins and Hahn, 2022). Therefore, ABBA-BABA tests are usually more suitable for testing more recent introgression events. Phylogenetic networks, on the other hand, use discordance between gene trees and/or concordant factors, being thus less impacted by the multiple substitutions at individual sites, making them more suitable for estimating ancestral introgression events (Hibbins and Hahn, 2022). While we only found evidence for one reticulation event from K. hermaphroditus 'Southern clade' into Kryptolebias sp. 'ESP.' in our phylogenetic network using PhyloNetowrks, our phylogenetic network visualization using SplitsTree revealed many tree discordances in the mangrove killifish clade, which is indicative of potential introgression. Our genome-wide ABBA-BABA tests further confirmed that indication, with evidence of introgression in the hybrid zones recently described between K. marmoratus and K. hermaphroditus 'Central clade' (Tatarenkov et al., 2021), between K. hermaphroditus 'Southern clade' and K. ocellatus (Berbel-Filho et al., 2021), and finally a significant signal of introgression in the hybridization found here between K. hermaphroditus 'Southern clade' and Kryptolebias sp. 'ESP'. In addition to the assumption of low substitutions per site, it is also important to acknowledge here another caveat of ABBA-BABA tests, which is the assumption of no ancestral population structure in the

ancestor between *P*1, *P*2 and *P*3 (Hibbins and Hahn, 2022). If present, ancestral population structure can result to similar deviations of site patterns counts as the ones caused by real introgression events (Eriksson and Manica, 2012). Despite those limitations, overall, our findings suggest that introgressive events in *Kryptolebias* are common, both at ancestral and/or recent time scales Considering the possibility that both lineages of *K. hermaphroditus* (Central and Southern clades) may belong to the same species (Lira et al., 2021), our results indicate that *K. hermaphroditus* has been involved in at least three different introgression events with other *Kryptolebias* species across its range. Although ABBA-BABA tests cannot evaluate the direction of introgression, these findings challenge the idea that highly-selfing taxa provide low opportunities for hybridization and introgression in the long-term (Berbel-Filho et al., 2021; Pickup et al., 2019).

#### 4.2. The mysterious Kryptolebias sp. 'ESP' lineage

Given the unusually high frequency of males and their unique external coloration, we sampled in the same locality (Coqueiral Beach, in Aracruz, Espírito Santo (Supplementary Fig. S1)) described in Costa (2016) to grasp further insights on the taxonomic status of this population. In total, we sampled 46 Kryptolebias individuals from this sampling locality, three of them could be identified based on external coloration as 'K. hermaphroditus' males according to Costa (2016). All other non-male individuals captured in this sampling locality had the typical external appearance of hermaphrodites of K. hermaphroditus, as also reported by Costa (2016). However, our results revealed that in this locality-two clearly differentiated Kryptolebias lineages/species coexist. The fact that the Kryptolebias sp. 'ESP' lineage was only found in a case of mito-nuclear discordance with K. hermaphroditus, calls for attention to the possibility of a cryptic species of Kryptolebias in the region. In fact, Sarmento-Soares et al. (2014) found two Kryptolebias populations in other coastal streams in the state of Espírito Santo. Those individuals were initially identified as K. ocellatus. Later, Costa (2016) collected individuals from Coqueiral beach (the same locality sampled here) and identified them as K. hermaphroditus. Our analyses did not find any evidence that individuals from Coqueiral Beach are K. ocellatus, however they add another syntoptic Kryptolebias lineage currently coexisting with K. hermaphroditus in Coqueiral Beach. In addition, Costa (2016) used the male coloration found on males from this locality (together with another locality in Rio de Janeiro state) to generally describe the coloration of K. hermaphroditus males. While his description of male coloration is detailed and accurate, there are clear differences in coloration between the K. hermaphroditus males found by Berbel-Filho et al. (2016), other males sampled in different localities within the species range (Amorim et al. (2022); Supplementary Fig. S1) and the ones described in Costa (2016). Although we do not have nuclear data for male individuals from this population, the evidence presented here for Kryptolebias sp. 'ESP' prompts for further taxonomical evaluation on the identity of males from this population, especially whether the males described in Costa (2016) represented males of Kryptolebias sp. 'ESP' or K. hermaphroditus (or both).

Another striking feature of the hybrid zone between *Kryptolebias* sp. 'ESP' and *K. hermaphroditus* in Coqueiral Beach is the evidence that despite the two species have exchanged DNA in the past and are currently syntopic, there is no apparent evidence of current gene flow between them (e.g., no F1s or early hybrid generations). Although we acknowledge our small sampling size for this population, this scenario may suggest a strong mechanism of pre and/or postzygotic reproductive isolation between the two sympatric lineages. Alternatively, we cannot fully rule out the possibility that the two individuals of *K. hermaphroditus* in Coqueiral beach found here represent a recent case of secondary contact, given the high heterozygosity found in *Kryptolebias* sp. 'ESP' together with the evidence from recent phylogeographic studies indicating that *K. hermaphroditus* has been dispersing southwards along the mangrove forests in the Brazilian coast recently (Berbel-Filho et al.,

2020; Lira et al., 2021; Tatarenkov et al., 2011; Tatarenkov et al., 2017). Overall, the evolutionary history of *Kryptolebias* sp. 'ESP', its potential distribution, as well its historical and current relationship with *K. hermaphroditus* is now open for investigation, with putative scenarios such as hybrid speciation involving *K. hermaphroditus* and a previously unknown (and possibly extinct) lineage, ancestral introgression involving a yet unknown lineage, as fruitful lines of research to understand the origins of the mysterious *Kryptolebias* sp. 'ESP' lineage.

#### 5. Conclusion

Introgression is the most common cause of mito-nuclear phylogenetic incongruences among taxa (Toews and Brelsford, 2012). Our phylogenetic reconstruction using mtDNA and genome-wide nuclear sites for the genus *Kryptolebias* generally agreed with previous reconstructions but yielded different topologies for the same set of species. Such discordance seems to have been caused by past introgression events in the genus. More importantly, our nuclear reconstruction recovered a cryptic *Kryptolebias* lineage hidden behind the case of mitonuclear discordance. The striking example of mito-nuclear discordance found here (with an unknown lineage having the same mtDNA haplotype of the introgressing lineage) highlights the need of using multiple genomic regions (particularly with different genomic, levels of recombination and inheritance properties) when reconstructing phylogenetic histories and making taxonomic inferences in clades where introgression is relatively common, such as *Kryptolebias*.

#### CRediT authorship contribution statement

Waldir M. Berbel-Filho: Conceptualization, Methodology, Formal analysis, Investigation, Resources, Data curation, Writing – original draft, Writing – review & editing, Funding acquisition, Visualization. George Pacheco: Methodology, Software, Formal analysis, Writing – review & editing. Andrey Tatarenkov: Resources, Writing – review & editing. Mateus G. Lira: Resources, Writing – review & editing. Carlos Garcia de Leaniz: Resources, Funding acquisition, Supervision, Writing – review & editing. Carlos M. Rodríguez López: Methodology, Software, Writing – review & editing. Sergio M.Q. Lima: Resources, Funding acquisition, Supervision, Writing – review & editing. Sofia Consuegra: Resources, Funding acquisition, Supervision, Writing – review & editing, Project administration.

#### **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

#### Data availability

The 21 additional *cox1* sequences generated for this study are available at GenBank (access numbers: OM962875-OM962895). Merged FastaQ files generated for this study can be found at accessed at NCBI (accession PRJNA815481). FastaQ files for the samples of *K. hermaphroditus* and *K. ocellatus* generated in Berbel-Filho et al. (2021) and used here can be accessed at NCBI (accession PRJNA563625). All scripts used in the project are available at: https://github. com/waldirmbf/KryptolebiasGenomics.

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#### Appendix A. Supplementary material

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