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-RESEARCH ARTICLE-

Assessing DNA Barcodes for Identification of Pufferfish Species (Tetraodontidae) in Turkish Marine Waters

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Abstract

In Turkish marine waters, pufferfish belongs to Tetraondontidae family are represented with 8 species, *Lagocephalus lagocephalus*, *L. sceleratus*, *L. spadiceus*, *L. suezensis*, *L. guentheri*, *Sphoeroides pachygaster*, *Torquigener flavimaculosus* and *Tylerius spinosissimus*. DNA barcoding can be useful in the assessment of cryptic or morphologically similar species of identification which is widespread in marine environment. The DNA barcode identification of the eight puffer species of the Tetraodontidae family in Turkish marine waters were examined by using mtDNA sequencing of the amplified partial mitochondrial cytochrome c oxidase I (COI) gene. COI contained 189 variable and 337 conservative nucleotides of which 183 were parsimony informative over 526 bp. Mean genetic diversity of all species was found to be 0.18164. The highest (0.26127) and lowest (0.00305) nucleotide divergence was observed between *L.spadiceus* and *T. flavimaculosus* andd between *L. spadiceus* and *L. guentheri*, respectively. The number of different haplotypes were 12 out of 23 sequences, and there was no shared haplotypes between pufferfish species.

Keywords:

Pufferfish, molecular identification, DNA Barcoding, COI

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Introduction

Pufferfishes are marine fish species that are distributed in tropical and subtropical areas of the Atlantic, Indian and Pacific Ocean. Puffers include 28 genera and approximately 184 species in all over the world marine waters within the Tetraodontidae family (Matsuura, 2015; Farrag et al., 2016), among which at least ten are found in the eastern Mediterranean (Farrag, 2014). This indegenous invasive species has established large populations along the coasts of many countries of the eastern Mediterranean basin such as Israel, Lebanon, Turkey (Mediterranean and Aegean coasts), Cyprus and Greece (Aegean and Ionian coasts), while still rapidly expanding westwards along the coasts of Egypt, Libya, and along the entire Tunisian coastline (Soussi et al. 2014). Apart from several large species used for human consumption as a delicious food in few countries, particularly in China, Korea, Japan and Taiwan (Oyaizu et al. 2000), most pufferfish species have not commercial value. Besides the small size of most species, the family is renowned for the occurrence of a powerful toxin in their skin and organs called tetrodotoxin (TTX). Tetrodotoxin is a very potent neurotoxin and one of the strongest marine paralytic toxins (El-Sayed et al., 2003; Sato et al., 2008).

In Turkish marine waters, pufferfishes are represented with 8 species, *Lagocephalus lagocephalus* (Linnaeus, 1758), *Lagocephalus sceleratus* (Gmelin, 1789), *Lagocephalus spadiceus* (Richardson, 1845), *Lagocephalus suezensis* Clark & Gohar, 1953, *Lagocephalus guentheri* Miranda Ribeiro, 1915, *Sphoeroides pachygaster* (Müller & Troschel, 1848), *Torquigener flavimaculosus* Hardy & Randall, 1983, *Tylerius spinosissimus* (Regan, 1908) (Turan et al., 2007).

Molecular genetic studies on mtDNA have proven benefits for examining the phylogeny and phylogeography of marine species (Meyer, 1993; Avise, 1994; Turan et al. 2015a). Sequence analysis of mtDNA regions is a quick tool to reveal phylogenetic relationships of marine species (Avise, 1994; Turan et al. 2008; Tabata & Taniguchi, 2000). Ever since different regions of mtDNA evolve at different rates, specific mtDNA regions have been targeted for inter and intra specific variation (Hauser et al. 2001; Mohindra et al., 2007; Turan et al., 2015b). DNA barcoding is a global venture that provides a standardized and effective genetic marker to marine and freshwater biodiversity, with significant conservation applications. The DNA barcoding approach is concentrated on a single part of the mitochondrial genome, because it presents portions conserved across taxa that are appropriate for primer design, while including polymorphism between and within species (Hebert et al., 2003; Kress & Erickson, 2008). The cytochrome oxidase subunit I (COI) region of the mitochondrial genome is sufficiently diverse so as to let the specific identification of a great majority of fish species (Kochzius et al., 2008; Kochzius et al., 2010).

Simple identification of pufferfishes by DNA barcoding and current level of interspecific and intraspecific genetic variation at pufferfish species which distributed in Turkish waters are very important to know. In spite of the wide scientific interest given to this family because of their ecological impact and having tetrodotoxin in their tissue, there have been no remarkabke study which investigated genetic identification and structure of these species in Turkish waters.

The goal of this study is to evaluate the practicability of DNA barcoding in the monitoring species biodiversity distributed along the Turkish marine waters at two levels by confirming the taxonomic identification and specifying intraspecific and interspecific variations for eight pufferfish species found in Turkish marine waters.

Material and Methods

Species, Lagocephalus lagocephalus, L. sceleratus, L. spadiceus, L. suezensis, L. guentheri and Torquigener flavimaculosus, were collected from the Antalya and Iskenderun Bay. The picture of pufferfish species collected from the Antalya and Iskenderun Bay are shown at Figure.1. All the samples were put in plastic bags individually and frozen at -20 °C till they were transported to the laboratory. All tissue samples were stored at -20 °C and 95 % ethanol till the analysis.

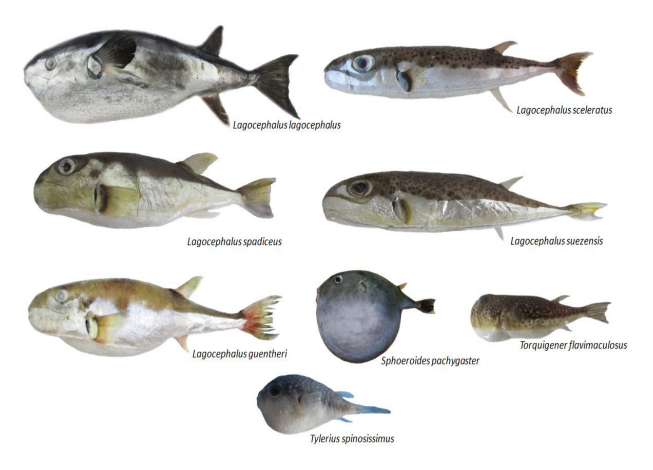


Figure 1. Pufferfish species in Turkish Marine Waters.

Total genomic DNA was extracted from the muscle and fin samples using the DNeasy Blood and Tissue Kit (Qiagen, USA). Manufacturer's protocols were used during all steps. Polymerase chain reaction (PCR) amplification was performed with following selective primers especially designed for this experiment:

COI-Forward 5'-TCAACCAACCACAAAGACATTGGCAC-3'

COI-Reserved 5'- ACTTCAGGGTGACCGAAGAATCAGAA-3'

The PCRs were conducted in a 50 ml total volume with 0.4 uM of each primer, 0.2 mM of dNTP and 1.25U of Taq DNA polymerase in a PCR buffer that included 20mM of Tris–HCl (pH

8.0), 1.5mM of MgCl2, 15 mM of KCl and 1-2 µl template DNA. Denaturation step were at 94°C for 30 s, 50 °C for 30 s, and 72 °C for 45 s for 30 cycles and followed by a final extension for 7 min at 72 °C. The PCR products were visualized using electrophoresis on 1.5 % agarose gel. The DNA sequencing was attempted to determine the order of the nucleotides of mtDNA COI region. The chain termination method by Sanger et al. (1977) was applied with Bigdye Cycle Sequencing Kit V3.1 and ABI 3130 XL genetic analyzer. Sequences for *Sphoeroides pachygaster* and *Tylerius spinosissimus* were taken from GenBank (S. *pachygaster*: JQ681814.1, JF494545.1, KJ709636.1-*T. spinosissimus*: JQ681847.1, KP266781.1, JQ681456.1). The initial alignments of partial COI sequences were performed with Clustal W program (Thompson et al., 1994) and final alignment was completed manually with BioEdit (Hall, 1999).

The molecular phylogenetic tree was constructed using Mega7 (Kumar et al. 2016). A distance-based method as neighbour joining (NJ) (Nei & Kumar, 2000) and a cladistics phylogenetic tree as maximum parsimony (MP) criterion were used. The reliability of the inferred phylogenies was evaluated using the bootstrap method (Felsenstein, 1985) with 1000 replicates.

Results

There were 189 variable and 337 conservative nucleotides of which 183 were parsimony informative over 526 bp sequences. The average nucleotide composition was 21.3% A, 28.8% T, 18% G and 31.9% C. Twelve haplotypes were found out of 27 sequences, and there was not shared haplotypes between species. *T. spinosissimus* had highest number of diferent haplotypes (Table 1). Minimum spanning tree that shows the relationships among the haplotypes (Figure 2.) Variable nucleotide positions of COI DNA barcode in pufferfish species show that Table 2.

Table 1. The number of haplotype and its distribution among species

	L. lagocephalus	L. sceleratus	L. spadiceus	L. suezensis	L. guentheri	S. pachygaster	T. flavimaculosus	T. spinosissimus	Total
Hap1	-	5	-		-	-	-	-	5
Hap2	-	-	4	-	-	-	-	-	4
Hap3	-	-	1	-	-	-	-	-	1
Hap4	-	-	-	5	-	-	-	-	5
Hap5	2	-	-	-	-	-	-	-	2
Hap6	-	-	-	-	2	-	-	-	2
Hap7	-	-	-	-	-	-	1	-	1
Hap8	-	-	-	-	-	-	1	-	1
Нар9	-	-	-	-	-	-	-	1	1

Hap10	-	-	-	-	-	-	-	1	1
Hap11	-	-	-	-	-	-	-	1	1
Han13						3			2
Hap12	-	-	-	-	-	3	-	-	3
Total	2	5	5	5	2	3	2	3	27

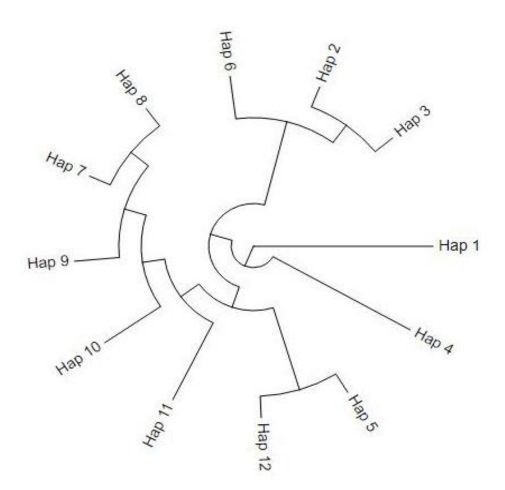


Figure 2. Minimum spanning tree that shows the relationships among the haplotypes.

Table 2. Variable nucleotide positions of COI DNA barcode in pufferfish species. The DNA barcode variable nucleotides are indicated, while identity is shown by dashes.

	10	20	30	40	50	60	70	80	90	100
Hap_1	CCCCACAATAATG	TCTATACAGACA	AGTGATCGAG	CCACATCCTCC	CTTTGCTTCC	CACTCGAGCT	AAGCCTCCA	GCCACTATA	CCTCGCTCGTT	ACCACC
Hap_2	TTA.GG.C.	CTC.C.TGAT	TAC.G.T.CA	GT.C.TC.T	A.CCTT.	G.G.AT.	GCATG	TCCA.	.TC.A.A	г
Hap_3	TTA.GG.C.	CTC.C.TGAT	TAC.G.T.CA	GT.C.TC.T	A.CCTT.	G.G.AT.	GCATG	TCCA.	.TC.A.A	г
Hap_4	c.	c.GTTG	T.CA.C		.сст	гA	.CTCA.C	AG.CC	TATG	GT.
Hap_5	GTAC.	.TC.CAT	TACAACA	C.T	A.CCTTA	AGTG.AT.	GCTT.CG	TCG.	.AATC.ACG	Г.А
Нар_6	GTA.GG.C.	CTC.C.TGAT	TAC.G.T.CA	GT.C.TC.T	A.CCTT.	G.G.AT.	GCATG	TCCA.	.TC.A.A	Γ
Hap_7	TTGCA	CAC.TTTT	TACCCA	TTC.T	AC.CCAGT	G.GGTGTTC	.CTCAT.	AA.T.C.CC	TAC.C.CTACA	TA.C.T
Нар_8	TTGCA	CAC.TTTT	TACCCA	TTC.T	AC.CCAGT	G.GGTGTTC	.CT.GCAT.	AA.T.C.CC	TAC.C.CTACA	TA.C.T
Нар_9	.T.T.TG.CA	c.c.TTTT	TACCCA	TG.TGC	.CGCAAA	A.TG.TG.A.	.cccg.c	AA.TGT	TTC.C.C.T.A	пт.т
Hap_10	.T.T.TG.CA	c.c.TTTT	TACCCA	TG.TGC	.CGCAAA	A.TG.TG.A.	.cccg.c	AA.TGT	TTC.C.C.T.A	гт
Hap_11	.T.T.TG.CA	c.c.T.ATTT	TACCCA	TG.TGC	.CGCAAA	A.TG.TG.A.	.cccg.c	AA.TGT	TTC.C.C.T.A	гт
Hap_12	T.TC.CC.	.TCT.ATAT	TACACC.A.A	.TGCT.CT.	TCATGA	A.TATTG.T.	.C.TTG	CATGAC.GT	ACTATT.G	T.GG
Conti	nued									
	110 1	120	130	140	150	160	1	70	180	
TTCCC	CATCCCCGACTAC	CTTAAACACT	CTCGACACG	стстстсс	CCCGCGCC	GCCCACTC	CACAACC	CGAGAGCG	CTCACCACC	TGTT
CG.T.	T.C.A.TA.TCC	rc.c.g	.GT.G.C	т	TTAGC	.тст	т.ттт	.ACTA	AC <mark>G.</mark> .	
CG.T.	T.C.A.TA.TCC	гс.с.g	.GT.G.C	T	TTAGC	.T.ACT	тттт	.ACTA	AC.TG	
AG.T.	T.CTG1	Γ	AG.G.A	TCT.TT	IGT.AT.T.	.TTTA.		.AG.TA	T	
	T.C.ATTA.TCT									
CG.T.	T.C.A.TA.TCC	rc.c.g	.GT.G.C	Т	TTAGC	.TCT	т.тт	.ACTA	ACG	• • • •
AGG	.T.AATTA.AA.	rc.cga.	A.TC.A	T.TAG.	TATAA.TA	A.C.A.	.TT.T	TAGAGA	.CAG.AG	GTCC
AGG	.T.AATTA.AA.1	r.ccggA.	A.TC.A	T.TAG.	TATAA.TA	A.C.A.	т.т.т	TAGAGA	.CAG	GTCC
CGAT.	AAAG.1	ΓCGAA	.GTGTA	T.TCT.GT.	A.ATC.T	ATC.C.	.CT.C.T	.AGA	тст.т	.A
CGAT.	AAAG.1	ГСАА	.GTGTA	T.TCT.GT.	A.ATC.T	ATC	.CT.T.T	.AG	тст.т	.Α
CGAT.	AAAG.1	ГАА	.GTGTA	T.TCT.GT.	A.ATC.T	ATC	.CT.T.T	.AG	TCT.T	.A
GGT	TTATT.GT	rc.gT.A.	TATA.GCTA	TCTCT.	A.ATAG.TA	.тА.	.c.cct.	т.тт.	ACTT.	.Α

Species special DNA barcode were detected whereas common DNA barcode was not detected for all species. Kimura 2 parameter method (Kimura, 1980) was selected as a best method for intra and interspecific variations. Mean genetic diversity for all species was found to be 0.18164. The matrix of pairwise distances within species is presented in Table 3. The intraspecific genetic diversity within *L. sceleratus*, *L. suezensis*, *L. lagocephalus*, *L. guentheri* and *S. pachygaster* was observed to be zero while it was highest within *T. flavimucolosus* (0.01149). The lowest genetic distance is observed between *L. guentheri* and *L. spadiceus* (0.00305) whereas the highest one is observed between *T. flavimucolosus* and *L. spadiceus* (0.26127).

Table 3. The matrix of intraspecific genetic distances between species (below diagonal) and genetic diversity (transversal diagonal given in bold).

	1	2	3	4	5	6	7	8
L. sceleratus (1)	0.00000							
L.spadiceus (2)	0.19850	0.00229						
L. suezensis (3)	0.13133	0.22827	0.00000					
L. lagocephalus (4)	0.21867	0.10202	0.19204	0.00000				
L. guentheri (5)	0.19673	0.00305	0.22639	0.09859	0.00000			
T. flavimucolosus (6)	0.25622	0.26127	0.24380	0.24518	0.26042	0.01149		
T. spinosissimus (7)	0.21345	0.21469	0.20354	0.20343	0.21338	0.19274	0.00896	
S. pachygaster (8)	0.25101	0.24912	0.23208	0.21013	0.24759	0.25409	0.21512	0.00000

The Neighbour Joining and Maximum Parsimony phylogenetic approaches resulted in similar tree topologies. In Neighbour joining phylogenetic tree, two main phylogenetic nodes were detected; in the first main node, T. *flavimucolosus* and T. *spinosissimus* grouped together. In second main node, 3 branches were detected; S. *pachygaster* was in the first branch, L. *spadiceus*, L. *guentheri* and L. *lagocephalus* were grouped in the second branch on which L. *guentheri* and L. spadiceus were grouped together as a sister group, and L. *sceleratus* and L. *suezensis* were grouped as a third branch (Figure 3).

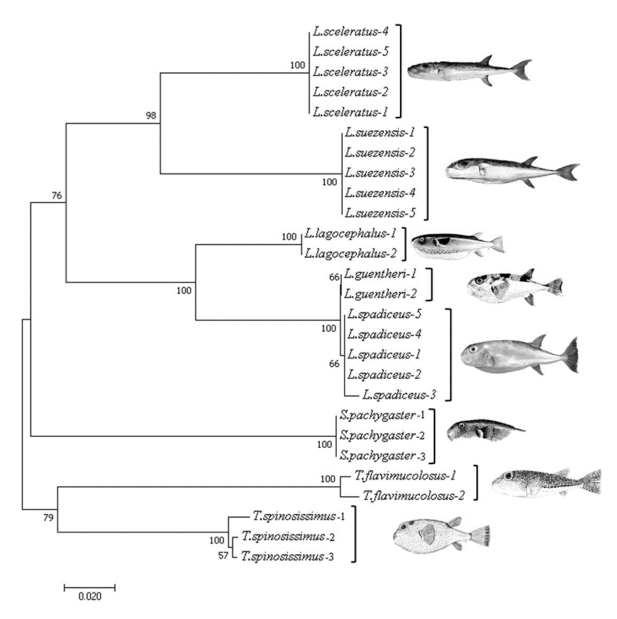


Figure 3. Neighbour joining phylogenetic tree based on COI sequences. Fish drawings from Froese & Pauly (2016).

In the Maximum Parsimony phylogenetic tree, two phylogenetic nodes were detected; in the first node, T. *flavimucolosus* and *T. spinosissimus* were grouped together. In second node 3 branches were detected; *S. pachygaster* was in the first branch, *L. spadiceus*, *L. guentheri* and *L. lagocephalus* were grouped as a second branch in which *L. guentheri* and *L. spadiceus* were grouped together as a sister group, and *L. sceleratus* and *L. suezensis* were grouped as a third branch (Figure 4).

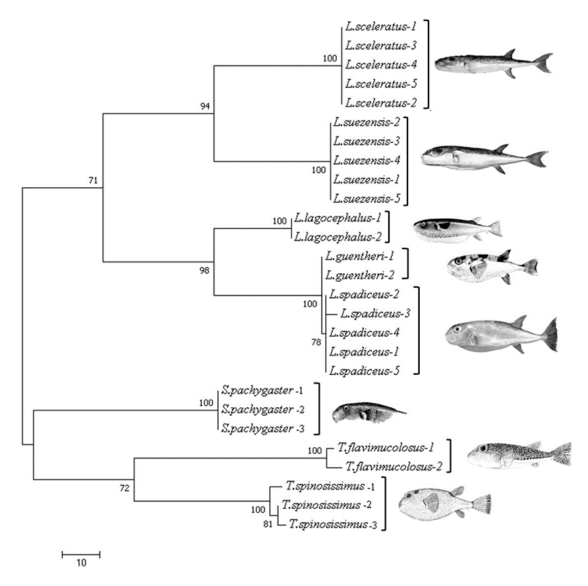


Figure 4. Maximum Parsimony phylogenetic tree based on COI sequences. Fish drawings from Froese & Pauly (2016).

Discussion

In the present study, the DNA barcoding of the eight pufferfish species which are distributed in the Turkish marine waters were investigated. All the species under the three genera were clearly separated in the NJ and MP trees with a high bootstrap value. The universal primers successfully amplified the target region in all species, generating 27 COI barcodes of 526 bp. Common haplotypes was not detected between species, and the DNA barcode sequences clearly discriminated taxonomic status of all pufferfish species examined.

Genetic diversity within species were calculated as zero for *L. sceleratus*, *L. suezensis*, *L. lagocephalus*, *L. guentheri* and *S. pachygaster*. This low genetic diversity may be explained low number of samples sequenced in the present study. On the other hand, the detected low genetic diversity may be due to founder effect that is usually observed in the alien species invation which

form established populations starting from meager individuals. A similar result is reported by Keskin & Atar (2013) using DNA barcoding to identify 89 commercially important freshwater and marine fish species found in Turkish ichthyofauna. Vinas & Tudela (2009) studied genetic identification of eight Scombrid species using mtDNA control region, mtDNA COI gene and nuclear DNA ITS1 region, and reported that credibility of COI gene is questionable that also reported that COI gene is not a good marker for inferring evolutionary relationships in Thunnus species.

The present study support the power of COI for species identification which is in accordance with other similar studies. Lakra et al. (2011) investigated DNA barcoding of fish and marine life representing 79 Genera and 37 Families from the Indian Ocean using cytochrome c oxidase I gene (COI) of the mtDNA and concluded that morphological characters were strongly authenticated the efficacy of COI in identifying the fish species with designated DNA barcodes that make DNA barcoding approach successful. Kochzius et al. (2010) aimed to evaluate the applicability of the three mitochondrial genes 16S rRNA (16S), Cytochrome b (Cyt b), and cytochrome oxidase subunit I (COI) for the identification of 50 European marine fish species by combining techniques of DNA barcoding and microarrays. As a result, while Cyt b and COI are equally well suited for DNA barcoding of fishes. On the other hand, 16S has drawbacks in discriminating closely related species. All these studies have shown that genetic identification by COI barcodes can provide a useful tool to identify species and to detect possibly cryptic species, and even to describe new species.

In conclusion, in this study has strongly authenticated the efficacy of COI in identifying the pufferfish species with designated barcodes. The present results also suggest that COI barcoding can be used as pragmatic method for resolving unambiguous identification of pufferfish species in marine waters of Turkey with applications in its management and conservation.

Acknowledgement

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