



Population genetic structure and demographic history of *Aphanius fasciatus* (Cyprinodontidae: Cyprinodontiformes) from hypersaline habitats in the eastern Adriatic

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Summary: In order to investigate the phylogeography and population genetic structure of the South European toothcarp (*Aphanius fasciatus*), we analysed gene sequences of two mitochondrial markers (cytochrome b and mtDNA control region) in samples from eight localities along the eastern Adriatic coast and combined them with sequences from other Mediterranean localities. Since the South European toothcarp primarily inhabits hypersaline water bodies, it is a good model species for understanding patterns of colonization and dispersal of species adapted to variable conditions. The eastern Adriatic populations are separated into two groups of genetically related populations. The Northern group contains the populations from Sečovlje, Pag, Dinjiška, Nin and Pantan, whereas the Southern group contains the populations from Ston, Ulcinj and Narta. The majority of divergence events date back to the Pleistocene epoch and it is likely that sea level changes during glacial cycles played a significant role in shaping the recent genetic structure of this species. Our results imply pronounced intraspecific structuring of this species, whereas great environmental variations resulted in a smaller intrapopulation genetic diversity of *A. fasciatus* than seen in other Mediterranean fishes.

Keywords: colonization; conservation priorities; divergence; evolution; genetic diversity; South European toothcarp.

Estructura genética poblacional e historia demográfica de *Aphanius fasciatus* (Cyprinodontidae: Cyprinodontiformes) en hábitats hipersalinos del Adriático oriental

Resumen: Con el objetivo de analizar la estructura genética y filogeografía de las poblaciones del fartet oriental (*Aphanius fasciatus*), se analizaron secuencias de genes de dos regiones mitocondriales (citocromo b y región de control del ADNmt) en individuos de ocho localidades a lo largo de la costa oriental del Adriático y de otras localidades mediterráneas. El fartet oriental habita principalmente masas de agua hipersalinas, y es una buena especie modelo para el estudio de los patrones de colonización y dispersión de especies adaptadas a condiciones variables. Los resultados de nuestro estudio muestran que las poblaciones del Adriático oriental se separan en dos grupos genéticamente relacionados. El grupo del Norte contiene las poblaciones de Sečovlje, Pag, Dinjiška, Nin y Pantan, mientras que el grupo del Sur contiene las poblaciones de Ston, Ulcinj y Narta. La mayoría de los eventos de divergencia pudiera ser que se remontaran a la época del Pleistoceno, y es probable que los cambios del nivel del mar durante los ciclos glaciales jugaran un papel importante en la estructura genética reciente de esta especie. Nuestros resultados revelan diferencias intraespecíficas pronunciadas en el fartet oriental, y por otro lado menor diversidad genética intrapoblacional como resultado de grandes variaciones ambientales en comparación con otros estudios de peces del Mediterráneo.

Palabras clave: colonización; prioridades de conservación; divergencia; evolución; diversidad genética; fartet oriental.

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INTRODUCTION

The fish community of the Adriatic Sea consists of some 440 species (Lipej and Dulčić 2010). Its diversity and structure were influenced by the specificities of the Adriatic Sea (Maggio et al. 2009), as a semi-closed area with its own characteristics in terms of salinity, temperature and depth (Astraldi et al. 1999), but also by evolutionary events that have shaped the configuration of the Mediterranean ichthyofauna. Though the Adriatic fish fauna is usually considered to be well studied (Dulčić et al. 2005), information on intraspecific structure and diversity is scarce and insufficient. This is particularly true for species inhabiting specific habitats, such as hypersaline water bodies. Investigations of demographic history and population genetic structure of species adapted to extreme conditions can help to reveal all components of Mediterranean biodiversity, and also to understand the peculiarities of evolution in specific environments.

A suitable model species for investigating evolution in such habitats is the South European toothcarp, *Aphanius fasciatus* (Valenciennes, 1821), a cyprinodont distributed throughout the central and eastern coastal zone of the Mediterranean. It is especially interesting since it is endemic to the Mediterranean, but inhabits habitats different from those of typical marine species.

Aphanius fasciatus usually inhabits brackish environments, such as coastal ponds and lagoons, though there are records of inland water populations (Parenti and Tigano 1993). Along the eastern Adriatic coast, it is found almost exclusively in hypersaline water bodies. It is able to tolerate a wide range of physicochemical parameters, in particular temperature (5–39°C) and salinity (0–180 psu) (Triantafyllidis et al. 2007). Due to its life history traits, such as large adhesive eggs, short generation time, high reproductive rate and, especially, the absence of larval dispersal stages, this species is considered to be an ideal organism for investigating the effects of genetic drift, selection and gene flow on population divergence (Triantafyllidis et al. 2007). Furthermore, *A. fasciatus* has no commercial value, so the distribution of genotypes among populations has likely not been affected by human translocations.

The genetic diversity of several populations of *A. fasciatus* has been investigated, though with opposing results. Hrbek and Meyer (2003) found low genetic variation among *A. fasciatus* populations, despite its wide distribution and potential subjection to Mediterranean vicariant events, whereas other authors documented marked genetic structuring among populations of this species (Rocco et al. 2007, Triantafyllidis et al. 2007). Hrbek and Meyer (2003) explained this low genetic divergence of *A. fasciatus* by occasional migration events that prevented differentiation. Triantafyllidis et al. (2007) analysed samples from 13 localities in Greece and Turkey and concluded that the Messinian Salinity Crisis was the most important event in shaping their recent population genetic structure. Bardakci et al. (2004) investigated Anatolian *Aphanius* populations and revealed limited gene flow among populations. They assumed extreme habitat conditions

to have caused occasional drastic size reductions of *Aphanius* populations that reduced intrapopulational diversity and increased interpopulational differentiation due to genetic drift. Furthermore, they stated that the Quaternary paleoclimate changes had modified the distribution range and intraspecific genetic structure of *Aphanius* species. The most recent study (Ferrito et al. 2013) found strong population genetic structuring within *A. fasciatus* that reflects the historical geography of the Mediterranean basin from the Late Miocene to the Pliocene. They recognized the presence of multiple glacial refugia and interglacial phylogeographic breaks for *A. fasciatus*.

The aims of this study were to investigate the genetic structure and demographic history of several *A. fasciatus* populations mostly inhabiting hypersaline water bodies along the eastern Adriatic coast, and to gain insight into the evolutionary history of this species. These populations have not previously been analysed, and therefore we aimed to ascertain their phylogenetic relationships, degree of intrapopulational diversity, as well as interpopulational differentiation and possible gene flow. Furthermore, we intended to verify the evolutionary hypotheses proposed in previous papers and to estimate time frame of diversification of the Adriatic *Aphanius* populations. Insight into the past history, together with a better understanding of the present status of a species, enables more precise prediction of its future alterations and is thus of great importance for conservation purposes.

MATERIALS AND METHODS

Sample collection

A total of 112 samples of *A. fasciatus* collected from eight localities along the eastern Adriatic coast were included in the study (Fig. 1, Table 1). Only one locality, Pantan, is an inshore brackish lagoon, whereas the remaining locations are hypersaline water bodies, entirely or partially altered into salt pans. Each population/locality was represented by 11–15 samples. Sequence sets for phylogenetic reconstructions were supplemented with available GenBank sequences (Supplementary material Table S1).

DNA extraction and PCR amplification

DNA extraction was performed using a standard extraction kit (DNeasy tissue kit, Qiagen). Two mitochondrial genetic markers with different mutation rates were analysed: the gene for cytochrome b (cyt b) and mitochondrial control region (mtDNA CR). The reaction volume for polymerase chain reaction (PCR) amplifications contained 25 µl AmpliTaq Gold Master Mix (Applied Biosystems Inc.), 2 µl of each primer (10 pmol), 4 µl template DNA and 17 µl water. The primers used for the amplification of the cyt b gene were H15915 and L14724 (Schmidt and Gold 1993), whereas the primers A and E from Lee et al. (1995) were used for the mtDNA CR. The thermal profile for PCR amplification of cyt b was 10 min at 95°C, 35



Fig. 1. – Map of the investigated area with marked sampling localities. The locations on the eastern Adriatic coast that were the main subject of the present study are marked in capital letters. From the remaining localities included in the map, mtDNA CR sequences were incorporated in the second data set and analysed to reveal the intraspecific phylogeographic structure of *A. fasciatus*.

cycles of 60 s at 92°C, 90 s at 48°C, 180 s at 72°C and 4 min at 72°C. PCR conditions for the amplification of mtDNA CR were 10 min at 95°C, 35 cycles of 45 s at 94°C, 45 s at 48°C, 60 s at 72°C and 7 min at 72°C. Sequencing with both forward and reverse primers was

carried out by the Macrogen Service Center using the same primers as for the PCR amplifications.

DATA ANALYSES

Sequence alignment

Homologous regions of *cyt b* and mtDNA CR were aligned manually against previously published sequences using BioEdit Sequence Alignment Editor (Hall 1999). Chromatograms and alignments were checked visually.

Analyses of population genetic polymorphism and diversity

The measures of DNA polymorphism obtained to describe the intrapopulation genetic diversity of the eastern Adriatic populations were haplotype diversity (H_d) and nucleotide diversity (Π). The level of genetic differentiation between populations was assumed by calculating statistics based on haplotypes (H_{ST}) and the one based on nucleotide sequences (K_{ST}). The decision to accept or reject the null hypothesis (there is no genetic differentiation between populations) was made based on the permutation test. Calculations were made using DNAsp software (Librado and Rozas 2009) and were conducted on both data sets (*cyt b* and mtDNA CR).

Genetic structure and historical demography analyses

In order to obtain insight into the intraspecific and population genetic structure, *cyt b* and mtDNA CR sequences from the eastern Adriatic populations were analysed separately using the statistical parsimony method (Templeton et al. 1992) under a 95% parsimony connection limit, using TCS software (version 1.2.1; Clement et al. 2000). With the aim of identifying

Table 1. – Intrapopulation genetic diversity and tests of selective neutrality for east Adriatic *Aphanius* populations, based on two molecular markers. N, number of sequences; h, number of haplotypes; S, number of polymorphic sites; H_d , haplotype diversity; Π , nucleotide diversity; asterisks mark statistically significant Fu's F_s and Tajima's D values; NG and SG explain populations belonging to the Northern and Southern group, respectively.

Population	Genetic marker	N	h	S	H_d	Π	Fu's F_s	Tajima's D
Sečovlje (NG)	<i>cyt b</i>	11	2	1	0.182	0.00016	-0.410	-1.129
	mtDNA CR	10	2	1	0.200	0.00051	-0.339	-1.112
Pag (NG)	<i>cyt b</i>	15	6	11	0.571	0.00129	-1.672	-2.180*
	mtDNA CR	12	4	3	0.455	0.00161	-1.590*	-1.179
Dinjiška (NG)	<i>cyt b</i>	14	4	3	0.571	0.00083	-0.544	0.006
	mtDNA CR	12	2	1	0.167	0.00042	-0.476	-1.141
Nin (NG)	<i>cyt b</i>	15	5	4	0.476	0.00047	-3.228*	-1.816*
	mtDNA CR	12	4	4	0.455	0.00168	-1.489*	-1.747
Pantan (NG)	<i>cyt b</i>	15	2	1	0.133	0.00012	-0.649	-1.155
	mtDNA CR	11	2	1	0.182	0.00046	-0.410	-1.129
Ston (SG)	<i>cyt b</i>	12	4	3	0.455	0.00044	-2.124*	-1.629
	mtDNA CR	11	3	4	0.345	0.00183	-0.019	-1.712
Ulcinj (SG)	<i>cyt b</i>	15	2	1	0.133	0.00012	-0.649	-1.155
	mtDNA CR	11	1	0	0	0		
Narta (SG)	<i>cyt b</i>	15	5	6	0.476	0.0007	-2.103*	-1.983*
	mtDNA CR	12	7	7	0.773	0.00294	-4.446*	-1.944*
Northern group	<i>cyt b</i>	70	13	17	0.551	0.00073	-10.425*	-1.270*
	mtDNA CR	57	10	8	0.293	0.00097	-11.605*	-2.078*
Southern group	<i>cyt b</i>	42	10	11	0.642	0.00083	-6.201*	-1.890*
	mtDNA CR	34	9	11	0.421	0.00163	-7.464*	-2.38*

most likely evolutionary processes responsible for recent phylogeographic structure, nested clade analyses (NCAs; Templeton 1998) were conducted on statistical parsimony networks based on both genetic markers. NCAs were performed using ANeCa software (Panchal 2007), which includes both TCS v1.18 (Clement et al. 2000) and GeoDis v2.2 (Posada et al. 2000), thus automating the nesting algorithm and the inference key that interprets the population structure and history at each nesting level.

Neutrality tests and mismatch distribution

To ascertain whether the observed pattern of genetic diversity is in accordance with the neutral evolution model, tests of neutrality were performed: Fu and Li's tests, and Tajima's test as implemented in DNAsp ver. 5.10.01. Since significant departures from the null hypothesis suggest either selective pressure on the locus under study or changes in population size (Ramos-Onsins and Rozas 2002), the results of the neutrality tests were used to detect traces of population growth. In particular, Fu's F_S statistic is expected to have negative values in expanded populations and it is considered one of the strongest tests in detecting population enlargements (Ramos-Onsins and Rozas 2002). For populations where genetic structures expressed indications of population expansion, their onsets were calculated on the basis of the peaks of theoretical mismatch curves, which were estimated using the same software. The mismatch distribution (the distribution of the number of pairwise mutation differences between sequences) is expected to be unimodal in recently expanded populations and irregular in populations of constant size (Rogers and Harpending 1992). The time of the population growth measured in the units of mutational time is expressed as the parameter τ ($\tau = 2ut$; u is the mutation rate per sequence, t is the absolute time) (Rogers and Harpending 1992). Thereafter, the time of population expansion was calculated from the parameter τ .

Phylogeographic analyses

Phylogeographic analyses were conducted on two data sets. The cyt b haplotypes of *A. fasciatus* obtained in this study were pooled with cyt b sequences of several other *Aphanius* species (*A. sophiae* [Heckel, 1847], *A. mesopotamicus* Coad, 2009, *A. farsicus* Teimori, Esmaili and Reichenbacher, 2011, *A. isfahanensis* Hrbek, Keivany and Coad, 2006, *A. vladkovi* Coad, 1988, *A. iberus* [Valenciennes, 1846], *A. baeticus* Doadrio, Carmona and Fernandez-Delgado, 2002, *A. saourensis* Blanco, Hrbek and Doadrio, 2006 and *A. dispar* [Rüppell, 1829]) retrieved from GenBank (S1) to assemble the first data set. The purpose of phylogenetic reconstruction of this data set was to ascertain the position of eastern Adriatic *Aphanius* populations and their relationships with other *Aphanius* species, since they were not included in any of the previous analyses. The phylogenetic reconstruction was conducted using two methods of phylogenetic inference: maximum parsimony (MP), as implemented in PAUP

(version 4.0b10; Swofford 2002), and Bayesian inference (BAY), implemented in MrBayes (version 3.1.2, Ronquist and Huelsenbeck 2003). For MP analyses, the heuristic search mode with 100 replicates was used, with randomized input orders of taxa, and tree bisection-reconnection branch swapping with all codon sites and nucleotide substitutions types weighted equally. Nonparametric bootstrapping (100 pseudoreplicates, 10 addition-sequence replicates) was used to assess the branch support. Each BAY analysis consisted of two simultaneous runs. For each, four Markov Chain Monte Carlo chains were run for three million generations with trees sampled every 100 generations. The first 20% of the sampled trees were discarded and Bayesian posterior probabilities were estimated from the 50% majority-rule consensus tree of the retained trees. The sequence of *Poeciliopsis monacha* Miller, 1960 (GenBank accession number AF047346), belonging to the same order but a different family than *Aphanius*, was used as an outgroup.

The second data set was composed of the mitochondrial control region sequences obtained in this study, and mtDNA CR haplotypes available from the GenBank (S1, localities marked in Fig. 1). This data set contained haplotypes of *A. fasciatus* only; however, more localities were included to enable more detailed analysis of the intraspecific phylogeographic structure. Since the mitochondrial control region accumulates mutations up to ten times faster than the rest of the mitochondrial genome, it is considered suitable for investigations of rapidly evolving populations (Meyer 1993) and is therefore the genetic marker of choice for investigating intraspecific relationships. In order to visualize genealogical relationships among closely related haplotypes (belonging to the same species) and to enable the possibility of presence of ancestral haplotypes in the recent genetic structure, a phylogenetic network was constructed. Network construction was based on the median-joining algorithm using Network 4.5.1.6. software (Fluxus Technology Ltd.). In addition to the sequences of *A. fasciatus*, the sequence of *A. dispar* (GenBank accession number UO6052) was included to enable identification of ancestral haplotypes.

Divergence time estimation

The molecular clock calibration, namely the estimation of divergence per pairwise comparisons per million years for both genetic markers, was conducted using the Bayesian MCMC coalescent method implemented in BEAST v1.7.0. (Drummond et al. 2012). Since only one evolutionary event has been proposed with a time frame (separation of *A. baeticus* from *A. iberus* was associated with the opening of the Strait of Gibraltar 5.5 million years ago; Perdices et al. 2001), this was used to calibrate the phylogenetic tree based on cyt b haplotypes. A relaxed clock model was employed (since general clock-like behaviour was rejected) with branch rates drawn from an uncorrelated lognormal distribution and a Yule speciation prior. The time of divergence of *A. fasciatus* estimated from this analysis was used to infer the divergence rate of *A. fasciatus*

Table 2. – Results of the genetic differentiation tests (H_{ST} , K_{ST}) based on cyt b data set (below diagonal) and mtDNA CR data set (above diagonal). Numbers in brackets are p-values based on permutation test. P-values lower than 0.02 are considered statistically significant after false discovery rate correction. Values in bold denote populations belonging to the Northern Adriatic group, plain text correspond to the Southern group.

H_{ST}	Sečovlje	Pag	Dinjiška	Nin	Pantan	Ston	Ulcinj	Narta
Sečovlje								
Pag	0.01 (0.35)							
Dinjiška	0.41 (0.00)	0.26 (0.00)						
Nin	-0.01 (0.61)	-0.01 (0.97)	0.27 (0.00)					
Pantan	-0.02 (1.00)	0.03 (0.19)	0.47 (0.00)	0.02 (0.33)				
Ston	0.52 (0.00)	0.30 (0.00)	0.33 (0.00)	0.37 (0.00)	0.57 (0.00)			
Ulcinj	0.74 (0.00)	0.46 (0.00)	0.50 (0.00)	0.54 (0.00)	0.77 (0.00)	0.02 (0.27)		
Narta	0.49 (0.00)	0.32 (0.00)	0.32 (0.00)	0.36 (0.00)	0.54 (0.00)	0.37 (0.00)	0.54 (0.00)	
K_{ST}								
Sečovlje								
Pag	0.00 (0.28)							
Dinjiška	0.31 (0.00)	0.16 (0.00)						
Nin	0.00 (0.94)	-0.01 (1.00)	0.23 (0.00)					
Pantan	0.00 (1.00)	-0.01 (0.75)	0.34 (0.00)	0.00 (0.35)				
Ston	0.91 (0.00)	0.75 (0.00)	0.81 (0.00)	0.87 (0.00)	0.92 (0.00)			
Ulcinj	0.96 (0.00)	0.80 (0.00)	0.86 (0.00)	0.91 (0.00)	0.97 (0.00)	0.00 (1.00)		
Narta	0.85 (0.00)	0.70 (0.00)	0.75 (0.00)	0.82 (0.00)	0.87 (0.00)	0.43 (0.00)	0.53 (0.00)	

mtDNA CR, in addition to the number of mutations per million years. This was subsequently used to calculate divergence times of different populations using the Time estimates sub-program of the Network 4.5.1.6. software.

RESULTS

The phylogenetic reconstruction and analyses of population genetics included 112 newly obtained cyt b sequences that were 1140 base pairs long (GenBank accession numbers: KT280012-KT280033), and 91 new sequences of the mitochondrial control region, with lengths varying between 396 and 398 base pairs (GenBank accession numbers: KT280034-KT280057). For cyt b, 22 different haplotypes were found among the specimens from the eastern Adriatic coast, while for mtDNA CR 24 unique haplotypes were obtained in this study. Population structure and haplotype frequencies are presented in Supplementary Table S2.

Population genetic structure and diversity

The two genetic markers showed similar results regarding intrapopulation genetic polymorphism and revealed differences among the populations (Table 1). Based on the cyt b marker, the range of the number of haplotypes detected per population was 2-6, the haplotype diversity 0.133-0.571, and nucleotide diversity 0.00012-0.00129, whereas for mtDNA CR the range of haplotype number per population was 1-7, for haplotype diversity 0-0.773 and for nucleotide diversity

0-0.00294. Both analysed genetic markers indicated the highest measures of genetic polymorphism in the Pag, Nin and Narta populations, while populations from Pantan and Ulcinj displayed the smallest amount of genetic diversity.

Both statistics calculated to infer genetic divergence among populations (H_{ST} , K_{ST} ; Table 2) revealed the same result: the Adriatic populations form two genetically distinct groups. The Northern group consists of the populations from Sečovlje (Slovenia), Pag, Dinjiška, Nin and Pantan (Croatia). The Southern group includes the populations from Ston (Croatia), Ulcinj (Montenegro) and Narta (Albania). Based on control region sequences, no significant genetic divergence was detected among the populations belonging to the same group, whereas populations from different groups were genetically different. However, the cyt b sequences showed genetic similarity for the majority of populations from the same group, with a single exception in each group (Dinjiška and Narta). When the two groups were compared, genetic polymorphism was higher in the Southern group.

Demographic history

The demographic history of the two population groups from the eastern Adriatic coast is based on different events (Table 3), as inferred by the NCA based on cyt b sequences (Fig. 2b). The NCA performed on mtDNA CR haplotypes (Fig. 2a) implied that allopatric fragmentation was responsible for the observed structure of the whole cladogram, while no

Table 3. – Interpretation of the results of the nested clade analysis of cyt b and mtDNA CR sequences of eastern Adriatic populations, based on the inference key of Templeton (2005). Only clades with significant chi-square statistics (χ^2) are included.

	Clade	χ^2	Chain of inference	Demographic event inferred
cyt b data set	1-1	67.79	1-2-3-4-NO	Restricted gene flow with isolation by distance
	2-1	40.71	1-2-11-12-NO	Contiguous range expansion
	2-2	43.00	1-2-11-12-13-14-NO	Long distance colonization and/or past fragmentation (not necessarily mutually exclusive)
mtDNA CR data set	Total cladogram	108.05	1-2 IOI-T	Inconclusive outcome
	Total cladogram	91.00	1-19-NO	Allopatric fragmentation

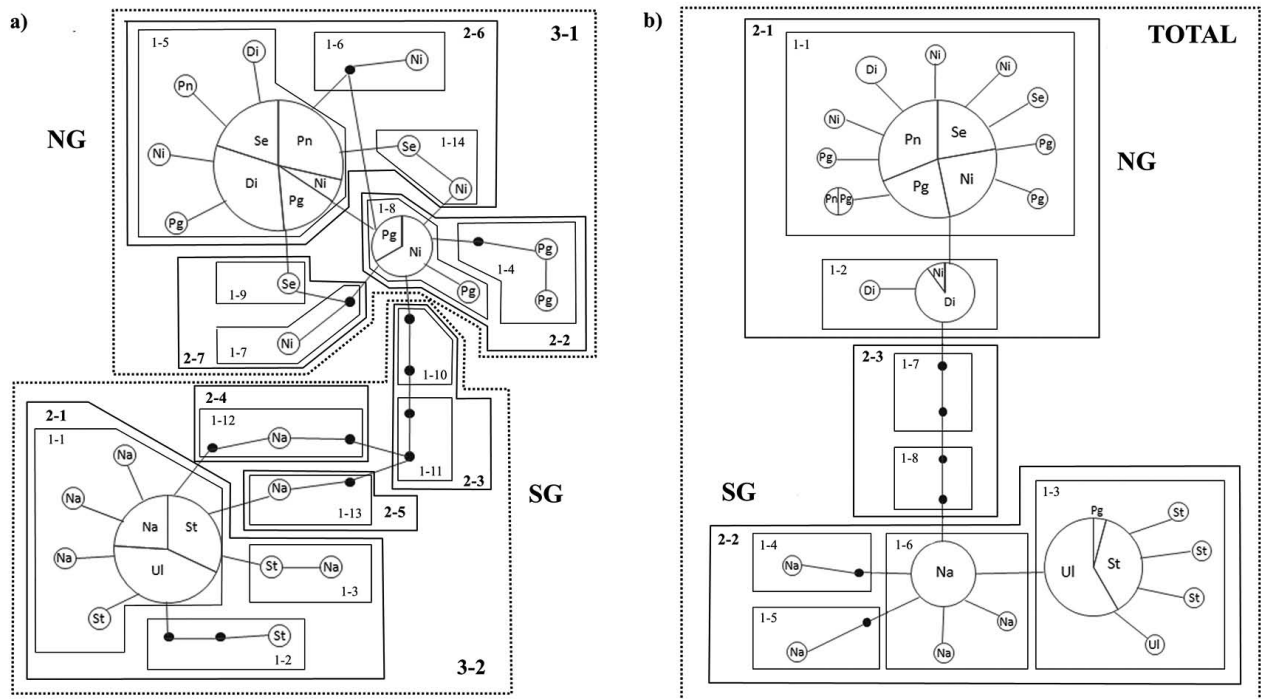


Fig. 2. – Statistical parsimony networks of the eastern Adriatic *Aphanius* populations based on mtDNA CR sequences (a) and nested clade design for the network containing cyt b haplotypes (b). Each line in the network represents a single mutational change; circle size of haplotypes is proportional to their number; small black circles represent unobserved haplotypes. Se-Sečovlje, Pg-Pag, Ni-Nin, Di-Dinjiška, Pn-Pantan, St-Ston, Ul-Ulcinj, Na-Narta. NG and SG explain populations belonging to Northern and Southern group, respectively.

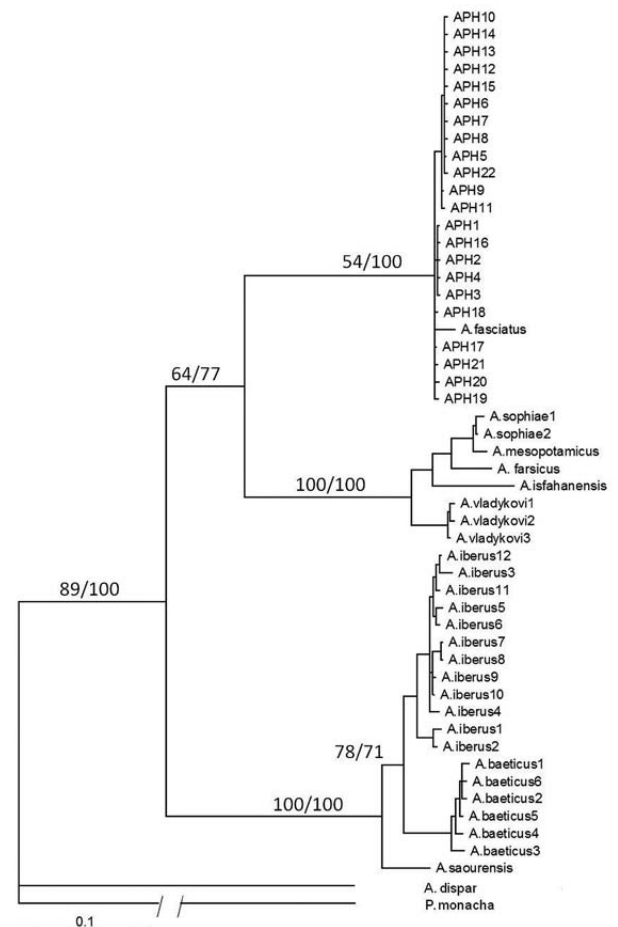
conclusive outcomes could be reached for any of the internal clades.

Fu's F_S and Tajima's D (Table 2) had significantly negative values for both genetic markers only for the Narta population, implying its demographic expansion and also suggesting very little overall deviation from the mutation-drift equilibrium. Nevertheless, traces of population growth were also noted for the Nin population (Fu's F_S statistic was significantly negative for both markers and Tajima's D was significantly negative for cyt b sequences) and probably for the Pag population (Fu's F_S was significantly negative for mtDNA CR and Tajima's D for cyt b sequences) (Table 2). Based on the parameter τ , the time of the onset of population growth dates back to the Pleistocene (324000-344000 years ago (YA) for the Narta population; 122000-444000 YA for the Nin population based on cyt b and control region sequences, respectively; and 177000 YA for the Pag population).

Phylogenetic position of *A. fasciatus*

Both phylogenetic reconstruction methods corroborated the monophyletic position of *A. fasciatus* and the inclusion of samples from the eastern Adriatic coast into the *A. fasciatus* phylogenetic lineage (Fig. 3). This is a sister lineage to the lineage comprising *A. sophiae*,

Fig. 3. – BAY phylogram of the cyt b haplotypes obtained in this study (symbols in capital letters), together with those retrieved from GenBank (first data set). Numbers at nodes represent Bayesian posterior probabilities and maximum parsimony branch support values, respectively.



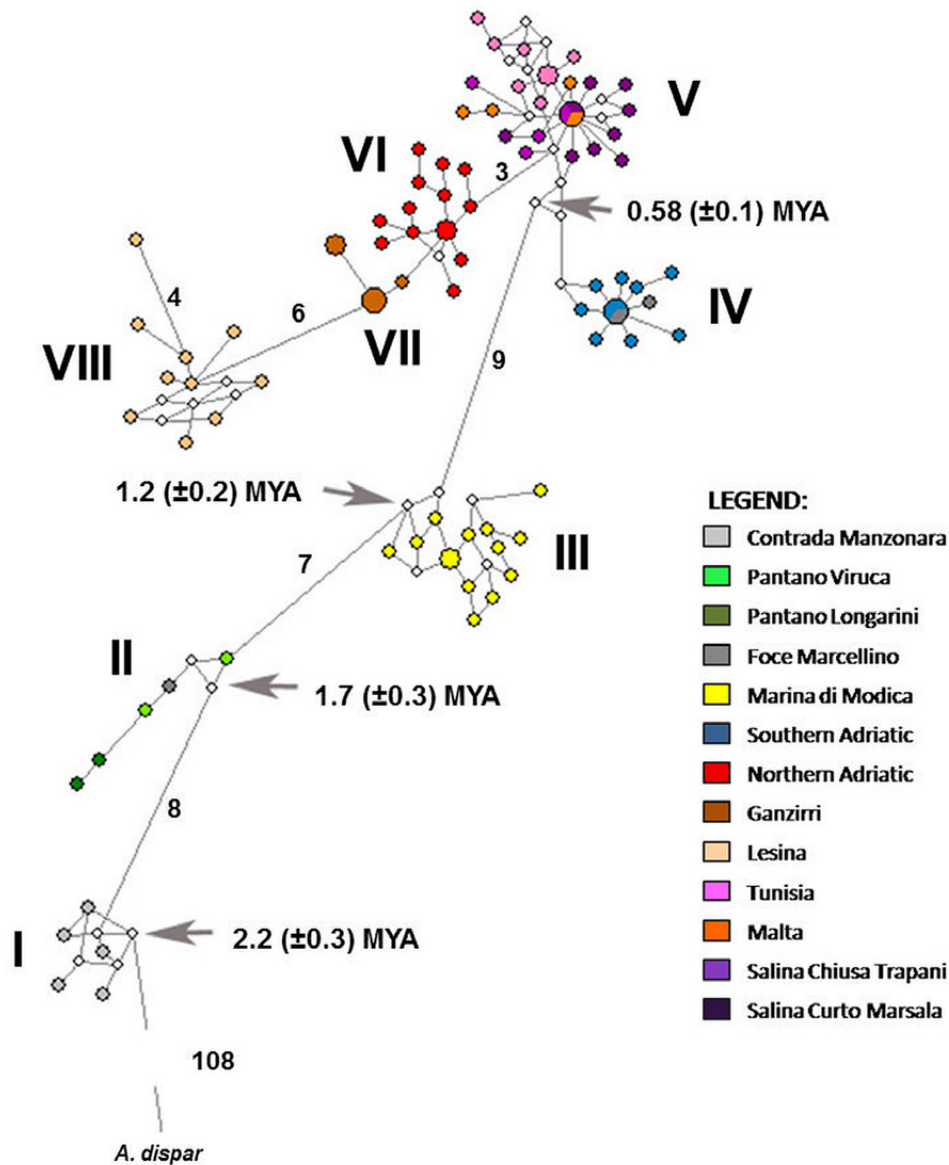


Fig. 4. – Median-joining network of mitochondrial control region haplotypes. Circle size indicates haplotype frequency, whereas branch lengths are proportional to the number of mutations. Where there are more than two mutational steps, the number of mutations is displayed along that branch. Roman numerals indicate clusters with haplotypes from Contrada Manzonara (ITA) (I); Pantano Viruca, Pantano Longarini and Foce Marcellino (ITA) (II); Marina di Modica (ITA) (III); Ston (CRO), Ulcinj (MNE), Narta (ALB) (e.g. Southern Adriatic group) and Foce Marcellino (ITA) (IV); Salina Curto Marsala, Salina Chiusa Trapani (ITA), Malta and Tunisia (V); Sečovlje (SLO), Nin, Pag, Dinjiška and Pantano (CRO) (e.g. Northern Adriatic group) (VI); Ganzirri (ITA) (VII); and Lesina (ITA) (VIII). Numbers by arrows represent estimated divergence time of population or cluster.

A. mesopotamicus, *A. farsicus*, *A. isfahanensis* and *A. vladkovi*. The third phylogenetic group is composed of *A. iberus*, *A. baeticus* and *A. saourensis*, whereas *A. dispar* is more distantly related to the remaining investigated *Aphanius* species.

Phylogeographic structure of *A. fasciatus* and divergence time estimates

The median-joining network comprising 45g the newly obtained mtDNA CR haplotypes and haplotypes from several localities in Italy, Malta and Tunisia (Fig. 4) shows the interesting intraspecific genetic configuration of *A. fasciatus*. Structuring correlated with geographic position is clearly visible, with haplotypes belonging to the same population almost al-

ways clustering together. However, populations with haplotypes separated by a reduced number of mutational steps are not necessarily geographically close and vice versa. Separation of the eastern Adriatic populations into two evolutionary groups (Northern and Southern) is clearly visible in the median-joining network (Fig. 4) and in the statistical parsimony networks (Fig. 2).

Based on molecular clock calibration, the evolutionary rate of *cyt b* was estimated as 1.2% per million years, whereas for the mitochondrial control region, the estimate was 3.6%. The divergence times of different *A. fasciatus* populations are marked on the median-joining network (Fig. 4). The last common ancestor of northern and southern Adriatic populations is dated back to around 0.58 million years ago.

DISCUSSION

The *A. fasciatus* populations along the eastern Adriatic coast form two groups of genetically related populations: Northern and Southern. These groups are genetically distinct, whereas populations within each group are genetically similar. Observed pattern is concordant with the pattern found by Hrbek and Meyer (2003) in their sample, and explained by occasional migrations that prevent differentiation. Moreover, in this study, all haplotypes were specific to the group, with only one exception (haplotype APH1 of the cyt b gene). Based on phylogeographic analyses and the fact that one observed haplotype was shared by both groups (cyt b haplotype APH1, which was found in multiple individuals from Ston and Ulcinj, and also in one fish from Pag), it could be concluded that though the two groups are separated, sporadic migrations of individuals between them are possible. Differences observed in the present genetic structure and polymorphism of the two groups probably better reflect the differences in their demographic history rather than differences in recent habitats. Previous studies also mostly explained that interspecific differences are due to evolutionary events (Rocco et al. 2007, Triantafyllidis et al. 2007). Given the geographic position of the Northern and Southern groups, it is possible that the historical Neretva River presented a barrier to *A. fasciatus* dispersal and promoted the diversification of these two groups. Namely, geological evidence has demonstrated that the Neretva R. was significantly longer during the Pliocene, extending along the Pelješac Peninsula (Alfirević 1965). It is well known that the Neretva R. represented a barrier to the dispersal of several terrestrial taxa (Podnar et al. 2004, Kryštufek et al. 2007, Ljubisavljević et al. 2007), though this is the first indication that it also represented an obstacle to the dispersal of marine organisms.

Although the Northern group is distributed over a greater number of localities, the Southern group is genetically more diverse. Furthermore, both investigated genetic markers indicated a greater variability of the populations from Pag and Nin than the remaining populations within the Northern group, and the greatest genetic polymorphism of Narta among southern populations. Environmental factors are highly variable at all localities, particularly salinity, which can often reach extreme values (37-330 psu), while temperature fluctuations (10-32°C) are also pronounced (Bakran-Petricioli and Jakl 2007). However, fish diversity is the richest at Pantan (Matić-Skoko et al. 2005) and Ulcinj (Schneider-Jacoby et al. 2006), localities where the smallest level of intrapopulational genetic polymorphism of *A. fasciatus* was recorded. Pantan is the only location that has not been changed anthropologically, and it is still an inshore lagoon located at the mouth of the small Pantana River. All other localities have completely or partially been transformed into salt pans where the toothcarp is the only fish species present. Therefore, it is possible that the higher genetic diversity observed in the Pag, Nin and Narta populations is a consequence of lower competitive pressure, whereas

extreme conditions do not limit the expansion of *Aphanius* populations.

The intrapopulational genetic diversity of all investigated *A. fasciatus* populations, even those expressing the highest levels of genetic polymorphism, was lower than the diversity recorded for other Mediterranean fish species (e.g. haplotype diversity of mtDNA CR of *Chromis chromis* [L., 1758] ranged between 0.792 and 1 [Domingues et al. 2005], while haplotype diversity of cyt b of *Pomatoschistus minutus* [Pallas, 1770] in the Mediterranean ranged between 0.822 and 0.844 [Gysels et al. 2004]). Less polymorphism was also observed for *A. iberus* (Perdices et al. 2001) and it has been hypothesized that rapid and pronounced changes of physicochemical and biological features may reduce population size, which, in combination with genetic drift, inbreeding and founder effects, could result in a loss of genetic diversity (Cognetti and Maltagliati 2000).

The phylogenetic position of *A. fasciatus* indicates its closer relatedness to several Iranian *Aphanius* species (*A. sophiae*, *A. mesopotamicus*, *A. farsicus*, *A. isfahanensis* and *A. vladkovi*) than to other Mediterranean species (*A. iberus*, *A. baeticus* and *A. saourensis*). That conclusion is in accordance with the results of previous studies (Hrbek and Meyer 2003), though samples were not composed of the same species and populations. Populations from the eastern Adriatic coast, analysed here for the first time, clustered into the *A. fasciatus* lineage.

Better resolution of intraspecific phylogenetic relationships among *A. fasciatus* populations was obtained through the phylogenetic reconstruction of the mitochondrial control region. The long-term isolation and abruption of gene flow among geographically close populations on the one hand, and the homogeneity of some distinct populations with clearly evidence of gene flow among them on the other, are the most prominent though contradictory features of the genetic structure of *A. fasciatus*. For a more precise explanation of the observed phenomenon, further investigations of the evolutionary history of this species are needed throughout its entire distribution range. Nevertheless, the results of this study also point out significant intraspecific genetic structuring of *A. fasciatus* (Rocco et al. 2007, Triantafyllidis et al. 2007, Ferrito et al. 2013).

Our results imply the importance of the Pleistocene epoch in shaping the recent population genetic structure and diversity of eastern Adriatic toothcarp populations. During the glacial maxima (in particular the Last Glacial Maximum, 25000-15000 YA), the entire northern part of the Adriatic Basin, down to the Zadar-Pescara line, was a wide fluvio-lacustrine plain with a coastline located at the northern edge of the Meso-Adriatic depression (Pirazzoli 2000). The northern border of the Adriatic Sea during the Last Glacial Maximum was located just north of the line connecting the Pantan and Lesina lagoons. The intrapopulational genetic divergence of the majority of the investigated Adriatic populations is of Pleistocene origin, implying that the colonization of the northern and southern parts of the eastern Adriatic localities did not occur

from a single locality after glaciations, but rather that populations remained in isolated patches of residual sea water or in temporary lakes that formed in the northern part of the Adriatic Basin. When connections between those localities were reestablished, gene flow among populations was enabled. Rocco et al. (2007) indicated that paleoclimatic fluctuations in the Quaternary influenced the genetic structure of *A. fasciatus*, though the detailed timing of population divergence has not been determined before. The conclusion that these two population groups have distinct evolutionary courses, despite inhabiting similar habitats located in geographic proximity, is corroborated by the NCP analysis results (see Table 3). For the populations of the Northern group, traces of restricted gene flow with isolation by distance were noted, which is concordant with the previously mentioned hypothesis that these populations survived long periods of the Pleistocene in small pools. On the other hand, the southern populations inhabit localities that were on the sea shore even during the glacial maxima. The importance of the Pleistocene epoch for the demographic history of the investigated species is highlighted by dating the onset of presumable population expansion of the Narta, Nin and Pag populations to that epoch.

Implications for conservation

Based on the results obtained, it can be concluded that the populations from Pag and Nin within the Northern group and the Ston and Narta within the Southern group are the most important populations for the further survival of *Aphanius fasciatus* in the Adriatic Basin. Although Pantan is under a special protection regime precisely because of the toothcarp population, it seems that salt pans have greater significance for the survival of this species, though inshore salt marshes are very important for overall biodiversity. Though those localities (especially Pag, Nin, Ston and Narta) are under substantial human impact, they nevertheless support populations of *Aphanius fasciatus* with high genetic diversity and that seem to be in expansion. For the existence of a species, maintenance of the highest genetic diversity is extremely important because it will allow adaptation to possible environmental changes. Therefore, populations from Pag, Nin, Ston and Narta, in addition to the Pantan population, should receive special conservation measures. Salt pans, as artificial anthropogenic formations and extreme habitats, are unfavourable places for most fish species. On the other hand, reduced competitive pressure makes them a convenient place for the toothcarp. Despite their recorded lower genetic diversity, natural, brackish wetland habitats are also very important for the conservation of populations of this species.

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SUPPLEMENTARY MATERIAL

The following material is available through the online version of this article and at the following link:
<http://www.icm.csic.es/scimar/supplm/sm04198esm.pdf>

Table S1. – The list of localities of the origin of sequences retrieved from GenBank (N, number of sequences). Country codes: SLO, Slovenia; CRO, Croatia; MNE, Montenegro; ALB, Albania; ITA, Italy; MLT, Malta; TUN, Tunisia; GRE, Greece; ESP, Spain; ALG, Algeria; IRN, Iran.

Table S2. – Haplotype frequencies for cyt B and mt DNA CR from the Adriatic localities. NG and SG explain populations belonging to the Northern and Southern groups, respectively.

**Population genetic structure and demographic history
of *Aphanius fasciatus* (Cyprinodontidae:
Cyprinodontiformes) from hypersaline habitats
in the eastern Adriatic**

Ivana Buj, Jure Miočić-Stošić, Zoran Marčić, Perica Mustafić, Davor Zanella,
Milorad Mrakovčić, Tanja Mihinjač, Marko Čaleta

Supplementary material

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Species	Locality	sample size	GenBank accession numbers	source	
<i>A. fasciatus</i>	Ganzirri (ITA)	6	EF440653 -55, 56-59	Rocco et al. 2007	
	Lesina (ITA)	10	EF440660 -69	Rocco et al. 2007	
	Marina di Modica (ITA)	14	EF440670 -83	Rocco et al. 2007	
	Contrada Manzonara (ITA)	5	EF440684 -88	Rocco et al. 2007	
	Pantano Longarini (ITA)	2	AJ605327, 29	Tigano et al. 2006	
	Pantano Viruca (ITA)	2	AJ605325, 28	Tigano et al. 2006	
	Foce Marcellino (ITA)	3	AJ605322, 23, 26	Tigano et al. 2006	
	Salina Curto Marsala (ITA)	5	AM184188, 198-201	Tigano et al. 2006	
		4	AM884565 -68	Pappalardo et al. 2008	
	Salina Chiusa Trapani (ITA)	2	AM184188, 97	Tigano et al. 2006	
		2	AM884569, 70	Pappalardo et al. 2008	
	Salina (MLT)	2	AM184186, 87	Tigano et al. 2006	
	Ghadira Bird Sanctuary (MLT)	3	AM184186, 88, 89,	Tigano et al. 2006	
	Lake Tunisi South (TUN)	6	AM184190 -95	Tigano et al. 2006	
	Ghar El Milh (TUN)	2	AM184190, 96	Tigano et al. 2006	
	Lusa (TUN)	2	JX406312, 26	Annabi et al. 2013	
	Oued Hamdoun (TUN)	2	JX406322, 24	Annabi et al. 2013	
	Sfax (TUN)	2	JX406313, 17	Annabi et al. 2013	
	Klisova marshes (GRE)	1	AF299273	Perdices et al. 2001	
	<i>A. iberus</i>	Villena lagoon (ESP)	2	AF299288 -89	Perdices et al. 2001
		San Pedro Pinatar (ESP)	2	AF299278 -79	Perdices et al. 2001
Albuixec (ESP)		2	AF299286 -87	Perdices et al. 2001	
Santa Pola (ESP)		1	DQ367528	Blanco et al. 2006	
Aiguamolls (ESP)		2	AF299276 -77	Perdices et al. 2001	
Adra (ESP)		2	AF299274 -75	Perdices et al. 2001	
Marchamalo (ESP)		1	DQ367529	Blanco et al. 2006	
<i>A. baeticus</i>	Salado Iro (ESP)	2	AF299284 -85	Perdices et al. 2001	
	Lucio del Hondón (ESP)	2	AF299282 -83	Perdices et al. 2001	
	Salado Conil (ESP)	2	AF299280 -81	Perdices et al. 2001	
<i>A. sophiae</i>	Kor River (IRN)	2	JN547800-01	Alavi-Yeganeh et al. 2014	
<i>A. mesopotamicus</i>	Karkkeh River (IRN)	1	JN565968	Alavi-Yeganeh et al. 2014	
<i>A. farsicus</i>	Kaftarak Spring (IRN)	1	JN547805	Alavi-Yeganeh et al. 2014	
<i>A. isfahanensis</i>	Gavkhoodi Lagoon (IRN)	1	JN565969	Alavi-Yeganeh et al. 2014	
<i>A. saourensis</i>	Mazzer (ALG)	1	DQ367527	Blanco et al. 2006	
<i>A. vladkovi</i>	Zagros (IRN)	1	DQ367526	Blanco et al. 2006	
	Koran River (IRN)	1	JN547802	Alavi-Yeganeh et al. 2014	
	Shalamzar Spring (IRN)	1	JN547803	Alavi-Yeganeh et al. 2014	
<i>A. dispar</i>	Mehran River (IRN)	1	JN547798	Alavi-Yeganeh et al. 2014	

Table S2. – Haplotype frequencies for cyt B and mt DNA CR from the Adriatic localities. NG and SG explain populations belonging to the Northern and Southern groups, respectively

Localities	haplotype frequencies	
	cyt b	mt DNA CR
Sečovlje (NG)	APH5=90.9%; APH22=9.1%	APHCR11=80%; APHCR23=10%; APHCR24=10%
Pag (NG)	APH1=6.7%; APH5=66.7%; APH12=6.7%; APH13=6.7%; APH14=6.7%; APH15=6.7%	APHCR11=41.7%; APHCR15=25.0%; APHCR18=8.3%; APHCR19=8.3%; APHCR20=8.3%; APHCR21=8.3%
Dinjiška (NG)	APH5=7.1%; APH9=64.3%; APH10=21.4%; APH11=7.1%	APHCR11=91.7%; APHCR22=8.3%
Nin (NG)	APH5=73.3%; APH6=6.7%; APH7=6.7%; APH8=6.7%; APH9=6.7%	APHCR11=16.7%; APHCR13=8.3%; APHCR14=8.3%; APHCR15=50.0%; APHCR16=8.3%; APHCR17=8.3%
Pantan (NG)	APH5=93.3%; APH14=6.7%	APHCR11=90.9%; APHCR12=9.1%
Ston (SG)	APH1=75.0%; APH2=8.3%; APH3=8.3%; APH4=8.3%	APHCR1=72.7%; APHCR2=9.1%; APHCR3=9.1%; APHCR4=9.1%
Ulcinj (SG)	APH1=93.3%; APH16=6.7%	APHCR1=100%
Narta (SG)	APH17=73.3%; APH18=6.7%; APH19=6.7%; APH20=6.7%; APH21=6.7%	APHCR1=50%; APHCR5=8.3%; APHCR6=8.3%; APHCR7=8.3%; APHCR8=8.3%; APHCR9=8.3%; APHCR10=8.3%

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