
ONLINE RESOURCES

Population genetic diversity of marble goby (*Oxyeleotris marmoratus*)

inferred from mitochondrial DNA and microsatellite analysis

CHENG ZHAO^{A,B#}, XIAOPING ZHU^{A,B#}, YICHUN GU^{A,B}, QINTAO WANG^{A,B},
ZECHENG LI^{A,B}, SHAOWU YIN^{A,B*}

^aCollege of Life Sciences, Key Laboratory of Biodiversity and Biotechnology of Jiangsu Province, Nanjing Normal University, Nanjing, Jiangsu 210023, China

^bCo-Innovation Center for Marine Bio-Industry Technology of Jiangsu Province, Lianyungang, Jiangsu 222005, China

[#]These authors contributed equally to this work.

Short title: Population genetic diversity analysis of marble goby

Keywords: Marble goby, Genetic diversity, MtDNA control region, Microsatellite, Population structure

Introduction

The marble goby, (*Oxyeleotris marmoratus*, Bleeker), also well known as the sand goby, is widely distributed in Southeast Asia especially in Malaysia, Singapore, Thailand, and Vietnam. (Mohsin *et al.* 1983; Cheah *et al.* 1994; Inger *et al.* 1963; Loo *et al.* 2015). It is an important commercial fish because of its good taste, rich nutrition, large size and potential in aquaculture. In China, the marble goby was introduced in 1986, and the comprehensive study has been conducted in the artificial propagation and breeding. Nowadays, it is widely bred in southern China such as Hainan, Guangdong and Fujian. In recent years, artificial culture of marble goby has suffered from germplasm degradation, such as, earlier sexual maturity, and reduced disease resistance which leading to a drastic reduction of the production (Lin and

Kaewpaitoon *et al.* 2000). Furthermore, the number of wild marble goby has decreased dramatically due to habitat destruction and overfishing (Cheah *et al.* 1994). As a consequence, the marble goby may have the problem of genetic diversity, especially in Vietnam and China, the major breeding countries of marble goby.

Mitochondrial DNA has been widely used to identify population genetic structure and variability due to its rapid evolutionary rate and complete maternal inheritance (Wilson *et al.* 1985; Sato *et al.* 2004). Most vertebrate mtDNAs contain a single non-coding segment, called displacement-loop region (D-loop region). The D-loop region of the mtDNA is highly polymorphic and it has 5–10 times rate of nucleotide substitution than nuclear DNA (Aquadro and Greenberg 1983). Therefore, it is often used to study the phylogenetic relationship, evaluate the population diversity, and determine the phylogeographic structure in fish species (Zhao *et al.* 2016; Zou *et al.* 2015; Chan *et al.* 2016). As for nuclear molecular markers, simple sequence repeat (SSR) markers are more informative and powerful tools for population genetic study because they are multi-allelic, co-dominant, and abundant in eukaryotic genomes when compared with other molecular marker systems. Thus, both mtDNA control region and SSR markers are favored as genetic tools for assessing genetic variation, genome mapping, population structure and intraspecific phylogenesis (Sharma *et al.* 2015; Romana-Eguia *et al.* 2004). In this study, 14 microsatellite and mitochondrial DNA markers were used to analyze the genetic diversity and population structure among three populations of marble goby. Our study will provide valuable information for facilitating the breeding program and the genetic conservation of marble goby.

Materials and methods

Sample collection and DNA extraction

Fin tissues of marble goby were acquired from a total of 91 individuals at three locations

including Mekong river of Ho Chi Minh City (N 106°63', E 10°76'), Vietnam (n = 28), Tianyuan aquatic seeding field of Zhuhai City (N 113°58', E 22°27'), Guangdong, China (n = 31) and Duanshan fry breeding base of Sanya City, Hainan (N 109°51', E 18°31'), China (n = 32) (Figure 1). Marble goby samples from three locations were designated “YS”, “GS” and “HS”, respectively. The details of sample we collected were listed in Table S1. Total genomic DNA was extracted following standard Proteinase K digestion and phenol-chloroform extraction method described previously (Hayano *et al.* 2003).

Mitochondrial control region sequencing and analysis

The primer pair of non-coding D-loop region was referred to its closely-related species *Eleotris acanthopoma* (GenBank accession EU369677) by using Primer Premier 5.0 software (5'-CTGCCTCAAAGAAGGGAGATT-3', 5'-TCACAGGGGTGCGGATACT-3'). PCR was carried out in 25 μ L reaction mixture containing 2.5 μ L 10 \times PCR buffer, 2 μ L 25 mM MgCl₂, 2 μ L 2 mM dNTP, 1 U Taq DNA polymerase, 0.5 μ L each primer (10 μ M), 16.3 μ L ultrapure water and 1 μ L (200 ng/ μ L) DNA template. PCR reaction conditions were performed by an initial denaturation step at 94°C for 5 min followed by 35 cycles at 94°C for 30 s, 52°C for 45 s, and 72°C for 1 min with a final extension at 72°C for 8 min. The products were sequenced directly using an ABI 3730XL automatic sequenced by BGI-Premier Scientific Partner (Beijing, China). The sequences were identified as mtDNA control region via the comparison with the sequences existed in NCBI (accession numbers: EU177621-EU177628) based on nBLAST. The DNA sequences were aligned with ClustalX 2.0 multiple-alignment program. Selection of the best-fit nucleotide substitution models was performed in jmodeltest 2.0. The most appropriate nucleotide substitution model was GTR+I+G. The phylogenetic analyses were performed using the maximum likelihood (ML), which by the PhyML 3.0 program. Bootstrapping was performed with 1000 replications. Nucleotide diversity (π) and haplotype diversity (h) and Tajima's D were estimated by using DnaSP 5.0. Population structure was

evaluated by F_{ST} , which was calculated by an analysis of pairwise difference (distance method) with the AMOVA in Arlequin Version 3.1.

Microsatellite genotyping analysis

Sixteen microsatellite markers of *Oxyeleotris marmoratus* have been developed by traditional method with constructing microsatellite-enriched library (Luo et al. 2013). Fourteen polymorphic microsatellite loci were selected, which show relative high allele number. These microsatellite loci were used for genotyping the marble goby from three different regions. The primer sequences, microsatellite core sequences, product sizes, and annealing temperature were listed in Table S2. PCR amplification was conducted in a reaction volume of 20 μ L containing 100 ng of genomic DNA, 1.5 mM Tris-HCl, 0.4 mM $(\text{NH}_4)_2\text{SO}_4$, 0.0375 mM MgCl_2 , 0.2 mM each dNTP, 0.2 μ M each of primer, and 0.5 U *Taq* polymerase (Fermentas). The PCR was performed at the following conditions: an initial hot start at 94°C for 5 min, followed by 30 cycles with denaturation at 94°C for 30 s, annealing at locus-specific temperature for 30 s, extension at 72°C for 30 s, and a final extension at 72 °C for 5 min. Fragment size of amplified DNA was determined on an AB 3500XL automatic sequencer. Popgene32 was used to calculate the number of alleles (A), observed heterozygosity (H_o), expected heterozygosity (H_e) and Wright's fixation index (F_{IS}). Hardy-Weinberg equilibrium (HWE), polymorphism information content (PIC), variance in allele frequencies and F_{ST} was achieved by Arlequin 3.1 software. Estimates of gene flow were derived by the equation: $N_m = [(1/F_{ST})-1]/2$.

Results and Discussion

Sequence analysis of the 855 bp mtDNA control region revealed 29 variable sites in 91 marble goby individuals. The numbers of variable sites were 11, 12 and 25 in YS, GS and HS populations, respectively. 91 sequences could be defined as 20 haplotypes based on the

nucleotide variation. Haplotype and nucleotide diversities of the three populations ranged from 43.4% to 80.4% and 0.127% to 0.455% respectively (Table 1). The total nucleotide diversity (0.31%) and haplotype diversity (61.6%) were observed, which is similar to that of some fish species, such as shortnose sturgeon (*Acipenser brevirostrum*) (Grunwald *et al.* 2002) and chum salmon (*Oncorhynchus keta*) (Sato *et al.* 2004). A negative Tajima's *D* value was observed in the YS populations ($P < 0.05$), which might be resulted from group expansion (Han *et al.* 2008). All haplotypes of mtDNA control region are available from GenBank under accession numbers of JN645822-JN645841. H20, comprising 61.5% of all samples, was the most widespread haplotype among three population (Table 2). It might derive from a single ancestor according to the theory of maternal inheritance of mtDNA. In the phylogenetic analyses, the maximum likelihood (ML) tree was constructed using the GTR+I+G model. Three clusters were supported by significant bootstrap proportions (node A: BP = 87%, node B: BP = 84%, and node C: BP = 99%). However, Maximum likelihood (ML) tree could not clearly differentiate haplotypes of specimens from three different locations (Figure 2). It could be explained by the occurrence of same haplotypes in different populations.

A total of 104 alleles at 14 microsatellite loci were detected from three different populations. Genetic heterozygosity which includes observed heterozygosity and expected heterozygosity. The average expected heterozygosity ($H_e = 0.472-0.504$) and observed heterozygosity ($H_o = 0.311-0.400$) at fourteen loci were observed in our study (Table 3), which is similar with Ruzainah's results (Ruzainah *et al.* 2009). YS populations ($H_o = 0.400$, $H_e = 0.504$) exhibited the highest value among all three samples, and this was consistent with the consequence of genetic diversity revealed by PIC value. Such results were ascribed to there is no wild marble goby resource in China. The population of marble goby of China was introduced from Southeast Asia. Through long-term artificial breeding, the genetic diversity of the cultured marble goby populations were relative low compared to Vietnam population.

Reduced genetic diversity in Chinese populations may result from founder effects. Founder effects could occur in artificial propagation when used a small number of broodstock individuals or no wild individuals introduced, which can led to the loss of genetic diversity (Allendorf *et al.* 1987). HEW tests showed that most of fourteen loci deviated significantly from Hardy-Weinberg expectations ($P < 0.05$) and exhibited heterozygote deficiency. This might be caused by the presence of null alleles (Pemberton *et al.* 1995) or the Wahlund effect (Hartl and Clark 1997).

AMOVA was conducted to describe the variance components of marble goby populations. The results of AMOVA from mtDNA control region and microsatellite suggested that most of the variations occurred within populations (Table 4), which is basically accordance with the population structure of marble goby in Southeast Asia (Chew *et al.* 2011). The pairwise genetic distance and genetic similarity were evaluated according to Nei's (1978) formula. GS and HS populations showed the lowest pairwise genetic distance (0.0338) and the highest pairwise genetic similarity (0.9668) (Table 5), which suggested that the genetic distance between Chinese population (GS-HS) was smaller than YS-GS and YS-HS. The result of mtDNA and microsatellite both indicated that higher genetic divergence and lower gene flow were found between Chinese and Vietnamese population (Table 6). F_{ST} value, in present study, was much smaller when compared with marble goby based on the mtDNA (Chew *et al.* 2011). The difference may be related to the sampling locations and sample quantity. In order to further elucidate the genetic diversities of marble goby, more sampling locations, size, and wild populations should be collected.

Our results demonstrated that the genetic diversity and genetic distance of Chinese populations were reduced when compared with the YS counterparts. The study, thus, suggests that the genetic conservation of marble goby populations in China is urgency. Moreover, it is important that the protection of wild stock and more wild resources should be introduced in

artificial breeding to maintain the genetic diversity of cultivated populations. Our research provides valuable information for the management and conservation of the marble goby.

Acknowledgement

This study was supported by the National Spark Program Project.(2011GA690368), Jiangsu Agricultural Science and Technology Independent Innovation Funds Project (CX(11)1037), Scientific foundation of Nanjing Normal University (2011104XGQ0071), and Project Foundation of the Academic Program Development of Jiangsu Higher Education Institutions (PAPD), China. National Undergraduate Training Program for Innovation and Entrepreneurship (201610319006).

References

- Allendorf F. W. and Ryman N. 1987 Genetic management of hatchery stocks. Washington Sea Grant Program, pp. 141-159. University of Washington Press, Seattle, USA.
- Aquadro C F. and Greenberg B D. 1983 Human mitochondrial DNA variation and evolution: analysis of nucleotide sequences from seven individuals. *Genetics*. **103**, 287-312.
- Chan J., Li W., Hu X., Liu Y. and Xu Q. 2016 Genetic diversity and population structure analysis of qinghai-tibetan plateau schizothoracine fish (*gymnocypris dobula*) based on mtDNA d-loop sequences. *Biochem. Syst. Ecol.* **69**, 152-160
- Cheah S. H., Senoo S., Lam S.Y. and Ang K. J. 1994 Aquaculture of a High-Value Freshwater Fish in [Malaysia]: the Marble or Sand Goby (*Oxyeleotris marmoratus*, Bleeker). *Naga*. **17**, 22–25.
- Chew H.H., Senoo, S., Tsunemoto, K., Nakagawa, Y., Miyashita, S., Murata, O. and Kato, K. 2011 Population structure of marble goby *Oxyeleotris marmoratus* (Bleeker) in Southeast Asia inferred from mitochondrial DNA. *Aquaculture Sci.* **59**, 383-391.
- Grunwald C., Stabile J., Waldman J. R., Gross R. and Wirgin I. 2002 Population genetics of shortnose sturgeon *Acipenser brevirostrum* based on mitochondrial DNA control region sequences. *Mol. Ecol.* **11**, 1885–1898.
- Han Z, Gao T, Takashi Y. and Yasunori S. 2008 Genetic population structure of *Nibeal biflora* in Yellow Sea and East China Sea. *Fish. Sci.* **74**, 544-552.
- Hartl D. L., and Clark A. G. 1997 Principle of population genetics. Sinauer Associates, Sunderland.
- Hayano A., Amano M. and Miyazaki N. 2003 Phylogeography and population structure of the Dall's porpoise, *Phocoenoides dalli*, in Japanese waters revealed by mitochondrial DNA. *Genes Genet. Syst.* **78**, 81–91.
- Inger R. F. and Chin P. K. 1963. The freshwater fishes on North Borneo. *Q. Rev. Biol.* **45**:1-268.
- Lin C. K. and Kaewpaitoon K. 2000 An overview of freshwater cage culture in Thailand. In: Lin, C.K., Liao, I.C. Eds. Proceedings of the First International Symposium on Cage

Aquaculture in Asia. pp. 237-242. Manila, Philippines.

Loo P., Chong C., Ibrahim S. and Sabaratnam V. 2015 Manipulating Culture Conditions and Feed Quality to Increase the Survival of Larval Marble Goby *Oxyeleotris marmorata*. *N. Am. J. Aquacult.* **77**, 149–159.

Luo J., Zhu X., Peng Y. and Yin S. 2013 Isolation and characterization of 16 microsatellite loci in marble goby (*Oxyeleotris marmoratus*). *Genet. Mol. Res.* **12**, 2020-2023.

Mohsin A. K. M. and Ambak M. A. 1983 Freshwater Fishes of Peninsular Malaysia, pp. 282. University Pertanian Publishing House, Serdang.

Pemberton J. M., Slate, J., Bancroft, D. R. and Barrett, J. A. 1995 Nonamplifying alleles at microsatellite loci: a caution for parentage and population studies. *Mol. Ecol.* **4**, 249–252.

Romana-Eguia M.R.R., Ikeda M., Basiao Z U. and Taniguchib N. 2004 Genetic diversity in farmed Asian Nile and red hybrid tilapia stocks evaluated from microsatellite and mitochondrial DNA analysis. *Aquaculture.* **236**, 131–150.

Ruzainah A., Siti Azizah M. N., Patimah I., Othman A. S. and Jamsari, A. F. J. 2009 Isolation and characterization of eight polymorphic microsatellite loci for the marble goby, *Oxyeleotris marmoratus*. *Mol. Ecol. Resour.* **9**, 1375-1429.

Sato S., Kojima H., Ando J., Ando H., Wilmot, R. L., Seeb, C. W. and Efremov, V. 2004 Genetic population structure of chum salmon in the Pacific Rim inferred from mitochondrial DNA sequence variation. *Environ. Biol. Fish.* **69**, 37–50.

Wilson A. C., Cann R. L., Carr S. M., George M., Gyllensten, U. B., Helm-Bychowski, K. M. and Higuchi, R. G. 1985 Mitochondrial DNA and two perspectives on evolutionary genetics. *Biol. J. Linn. Soc.* **26**, 375–400.

Zhao L., Chenoweth E. L., Liu J. and Liu Q. 2016 Effects of dam structures on genetic diversity of freshwater fish *Sinibrama macrops*, in Min River. *Biochem. Syst. Ecol.* **68**, 216-222.

Zou Z., Li D., Zhu J., Han J., Xiao W. and Yang H. 2015 Genetic variation among four bred populations of two tilapia strains, based on mitochondrial d-loop sequences. *Mitochondr. DNA.* **26**, 426.

Received 1 January 2017, in final revised form 26 March 2017; accepted 28 March 2017
Unedited version published online: 31 March 2017

Table 1 Genetic diversity estimation and Tajima's *D*-statistics on mtDNA control region sequences (855 bp)

Sample	N	No. of haplotypes	Haplotype diversity h (%)	Nucleotide diversity π (%)	Mean pairwise difference	No. of variable sites	Tajima's <i>D</i>
YS	28	5	43.4	0.127	1.085	11	-2.014*
GS	31	7	53.5	0.308	2.628	12	-0.405
HS	32	14	80.4	0.455	3.869	25	-1.415
Total	91	20	61.6	0.311	2.647	29	-1.696

An asterisk indicates the significant deviation from beta-distribution at $P < 0.05$

unedited version

Table 2 Number of fish from three populations of marble goby according to haplotype distribution

Haplotypes	H1	H2	H3	H4	H5	H6	H7	H8	H9	H10	H11	H12	H13	H14	H15	H16	H17	H18	H19	H20
YS														1	2			3	1	21
GS												1	3			3	1	1	1	21
HS	1	1	1	2	1	1	1	1	2	1	1					3	2			14
Total	1	1	1	2	1	1	1	1	2	1	1	1	3	1	2	6	3	4	2	56

unedited version

1 **Table 3** Genetic variation in fourteen microsatellite loci in three breeding groups of marble goby

Sample	Locus														Average across loci
	H113	H137	H27	H60	H167	H94	Y12	H117	H138	H191	H56	H142	H97	M351	
YS <i>N</i>	21	28	28	28	28	28	27	28	28	28	28	27	27	27	
<i>A</i>	6(162-186)	3(222-236)	11(272-294)	1(200-200)	2(230-230)	3(146-166)	5(217-229)	3(274-282)	6(154-166)	7(260-288)	4(260-270)	9(93-123)	15(191-263)	4(248-293)	5.642
<i>H_E</i>	0.753	0.260	0.786	0.000	0.036	0.260	0.575	0.519	0.678	0.646	0.260	0.805	0.798	0.687	0.504
<i>H_O</i>	0.762	0.071	0.821	0.000	0.036	0.000	0.148	0.607	0.500	0.250	0.250	0.704	0.519	0.926	0.400
<i>PIC</i>	0.690	0.240	0.747	—	0.035	0.240	0.504	0.397	0.620	0.607	0.242	0.762	0.770	0.611	0.497
<i>F_{is}</i>	-0.037	0.720	-0.064	—	-0.018	1.000	0.738	-0.192	0.249	0.606	0.022	0.109	0.338	-0.373	
<i>P</i>	0.000*	0.000*	0.059	—	1.000	0.000*	0.000*	0.552	0.095	0.000*	0.449	0.000*	0.000*	0.005*	
GS <i>N</i>	31	31	31	31	31	31	31	31	31	31	31	31	31	31	
<i>A</i>	7(148-184)	3(210-236)	8(242-298)	2(180-200)	1(232-232)	5(146-166)	3(207-229)	3(276-282)	5(138-162)	15(252-284)	5(258-270)	10(93-123)	15(203-263)	6(247-293)	6.286
<i>H_E</i>	0.721	0.154	0.711	0.063	0.000	0.268	0.235	0.463	0.552	0.846	0.268	0.754	0.861	0.712	0.472
<i>H_O</i>	0.290	0.097	0.516	0.000	0.000	0.097	0.000	0.258	0.323	0.516	0.290	0.516	0.516	0.936	0.311
<i>PIC</i>	0.671	0.146	0.672	0.061	—	0.256	0.215	0.377	0.441	0.816	0.255	0.710	0.832	0.643	0.469
<i>F_{is}</i>	0.591	0.361	0.262	1.000	—	0.633	1.000	0.434	0.406	0.380	-0.103	0.304	0.391	-0.336	
<i>P</i>	0.000*	0.020*	0.000*	0.017*	—	0.000*	0.000*	0.002*	0.001*	0.000*	1.000	0.000*	0.000*	0.006*	
HS <i>N</i>	32	32	32	32	32	32	32	32	32	32	32	32	32	32	
<i>A</i>	7(162-186)	3(212-236)	11(242-302)	2(160-200)	3(220-232)	3(160-166)	2(227-229)	3(280-286)	7(138-172)	10(260-288)	5(260-272)	9(93-123)	16(181-259)	4(248-293)	6.071
<i>H_E</i>	0.658	0.298	0.789	0.062	0.253	0.301	0.340	0.441	0.549	0.755	0.233	0.732	0.874	0.652	0.493
<i>H_O</i>	0.562	0.281	0.656	0.000	0.156	0.031	0.000	0.344	0.531	0.281	0.188	0.375	0.688	1.000	0.364
<i>PIC</i>	0.605	0.265	0.755	0.059	0.229	0.272	0.258	0.363	0.500	0.715	0.222	0.687	0.849	0.575	0.454
<i>F_{is}</i>	0.277	0.353	0.172	1.000	0.371	0.841	0.882	0.166	0.248	0.542	0.033	0.312	0.346	-0.412	
<i>P</i>	0.001*	0.027*	0.005*	0.016*	0.009*	0.000*	0.000*	0.235	0.121	0.000*	0.050	0.000*	0.000*	0.000*	

2 *N*, sample size for successful PCR analysis; *A*, number of alleles (allele size range); *H_E*, expected heterozygosity; *H_O*, observed heterozygosity; *PIC*, polymorphism
3 information content; *F_{is}*, Wright's (1978) fixation index; *P*, p-value of Hardy-Weinberg equilibrium; “*”, Hardy-Weinberg equilibrium ($P < 0.05$); “—”, there is only one
4 allele at the locus

5 **Table 4** AMOVA analyses of three marble goby populations, AMOVA I obtained from mtDNA control
 6 region and AMOVA II obtained from microsatellite loci

Source of variation	AMOVA I			AMOVA II		
	Sum of squares	Variance components	Percentage of variation	Sum of squares	Variance components	Percentage of variation
Among populations	5.275	0.041Va	2.811	18.835	0.105Va	3.35
Within populations	123.714	1.406Vb	97.189	544.242	3.040Vb	96.65
Total	128.989	1.447		563.077	3.145	

7
 8
 9

unedited version

10 **Table 5** Nei's genetic identity (above diagonal) and genetic distance (below diagonal)

Populations	YS	GS	HS
YS	****	0.9461	0.9464
GS	0.0554	****	0.9668
HS	0.0551	0.0338	****

11

12

unedited version

13 **Table 6** Genetic divergence (below diagonal) and gene flow (above diagonal) among three marble goby
 14 populations

Populations	mtDNA			Microsatellite		
	YS	GS	HS	YS	GS	HS
YS		7.585	10.164		11.605	12.546
GS	0.062*		—	0.041*		22.213
HS	0.047*	-0.004		0.038*	0.022*	

15 An asterisk indicates the significant deviation from beta-distribution at $P < 0.05$
 16

unedited version

17 **Table S1** Information of sample collection

Populations(abbreviation)	Latitude longitude	Sampling date	Sample number
Vietnam(YS)	N106°63' E10°76'	Jan.2011	28
Guangdong(GS)	N113°58' E22°27'	Mar.2011	31
Hainan(HS)	N109°51' E18°31'	Apr.2011	32

18

unedited version

19 **Table S2** Characterization of 14 marble goby (*Oxyeleotris marmoratus*) microsatellite loci

Locus	Repeat motif	Primer sequences (5'-3')	Size range (bp)	Ta (°C)	N_A	H_D	H_E	P(HWE)	Accession NO.
H113	(CA)11	F:GGAAGCTGCTGACCTTGACTC R:CCTATGGTCCGTCAGAGTGT	148-188	60	10	0.6604	0.7128	0.1519	JF264394
H137	(GT)9	F:GATCAGAGGGTTCAGAAAGCAG R:CATTACAGCACCGACAGAGGA	210-236	58	4	0.2075	0.2209	0.5780	JF264402
H27	(TG)21N105(AC)5	F:GATCAACAGTGTTCGCGTTAGG R:TCTCACCTGATGGAAAGATGG	242-302	56	13	0.7358	0.7686	0.2894	JF264376
H60	(GT)13N15(TG)6	F:GTTTGGCTGAAATGGTAGTGTG R:TGGAATGATGCTAGTGGCTGT	160-200	56	3	0.0943	0.1255	0.0854	JF264381
H167	(AC)9N73(AC)6	F:TCCATTACAGCACCGACAGAG R:GATCAGAGGGTTCAGAAAGCAG	220-232	58	3	0.2830	0.3245	0.5791	JF264409
H94	(TG)12	F:GAGGATTCCTCCGCTTCTATG R:GCCGTCCTTCGTGTTGTCTTG	146-166	56	5	0.2452	0.3012	0.1256	JF264391
Y12	(CA)7N4(CA)6	F:ATTATGATCCCCCACCAGCT R:TGTGATTGTCCCCTCTCACAG	207-229	57	3	0.3208	0.3859	0.2631	JF419699
H117	(CA)8N111(CA)5	F:ATAGCTCTGCGACGTGATTGG R:GACTTAGCTTTACCCTGTGGA	276-286	58	4	0.3962	0.4381	0.4815	JF264411

H138	(TC)5(AC)13	F: TAAGCCAGTGCCAGCAGAGT R: GCCCTGATTGTGACTGTGGAG	138-172	58	6	0.4717	0.4963	0.3953	JF264403
H191	(AC)11N51(GA)10N9(TG)4T2(TG)6	F: TGACATCTGTCTGGCTTCG R: GCCTGCGTCTTTGACAACCTC	254-288	58	15	0.7170	0.8033	0.2897	JF264413
H56	(AC)12	F: GCGAATTGCTGCAAGTGAGA R: GGTGTTGGGAGGAAGTGTAGGA	258-272	56	6	0.2453	0.2424	0.5998	JF264379
H142	(CA)6G(AC)8N5(CA)7	F: GAAATTGGAACGGGAGGCA R: ATGGGAGCCACGACTCACA	93-123	58	11	0.7547	0.7175	0.3493	JF264405
H97	(CA)24	F: AATCTGGCTTGACGCACTCT R: TTCCGCACGGTATCCTCTT	191-263	56	20	0.8491	0.8927	0.0445*	JF264392
M351	(TTCAA)5	F: GATCCTTTGCTCTGTTTCAG R: TCCTGGGTCTGTTAGTGTAG	247-293	54	5	0.6981	0.6759	0.1523	JF419693

20 N_A = allele number; H_O = observed heterozygosity; H_E = expected heterozygosity; T_a = annealing temperature; HWE = Hardy-Weinberg equilibrium. Statistically significance

21 at * $P < 0.05$.

Figure 1. Three sampling locations of marble goby: Vietnam, Guangdong and Hainan are represented by square, circle and triangle dot, respectively.

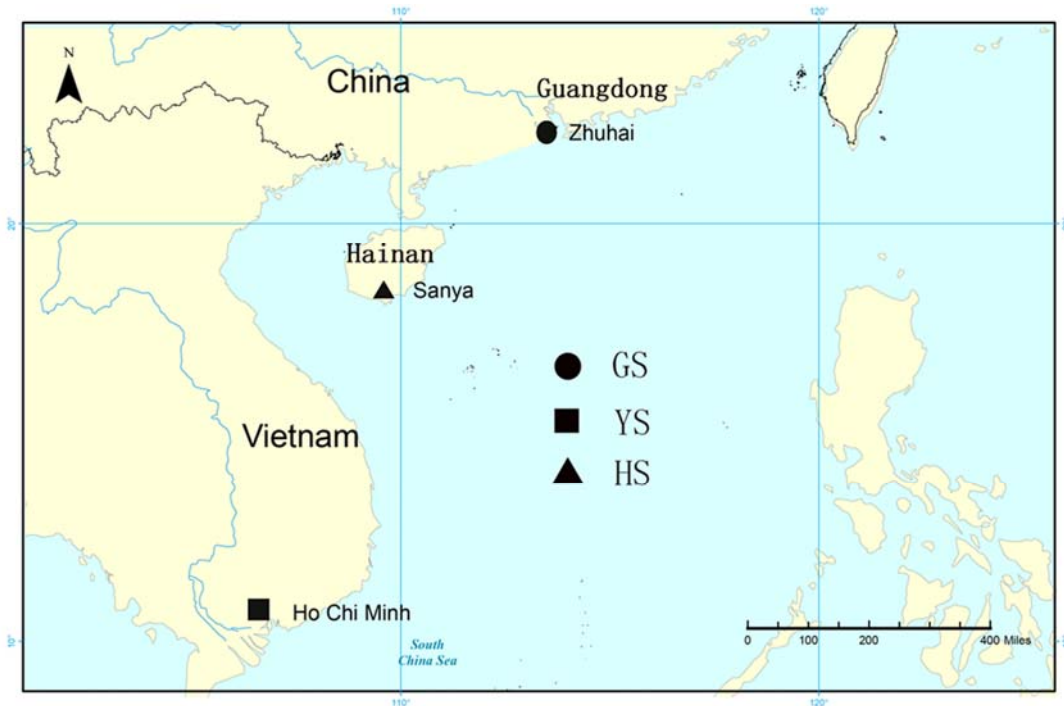
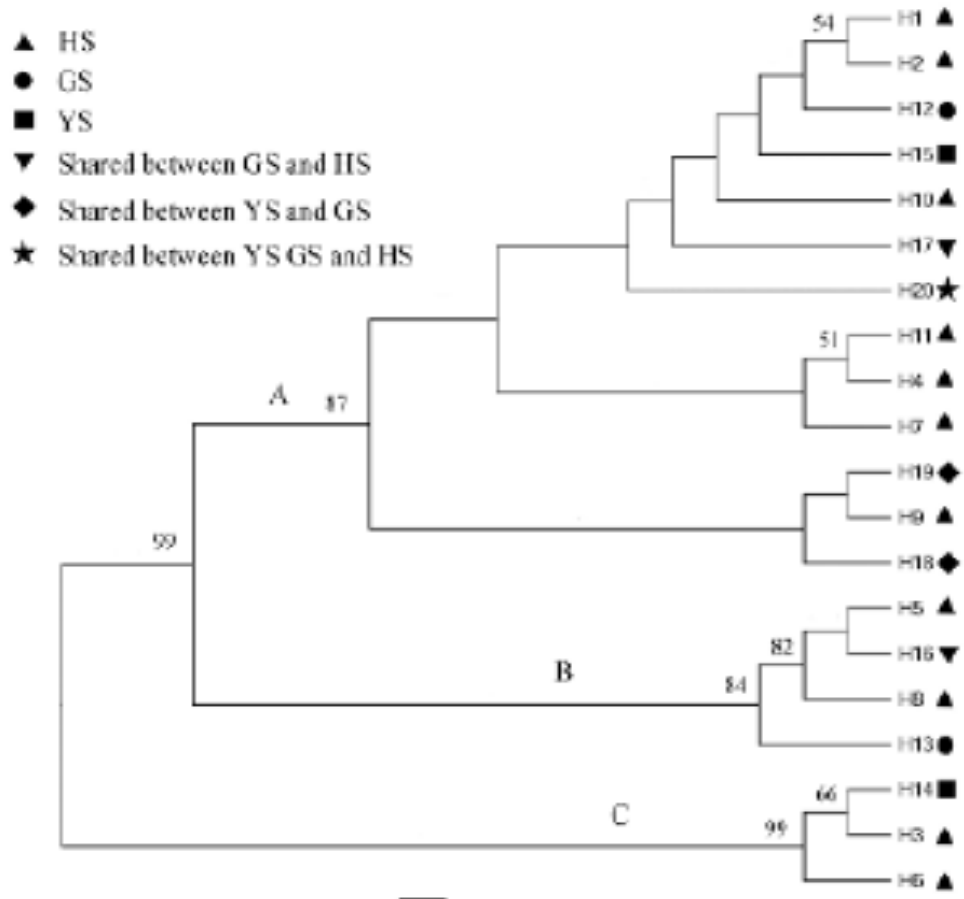


Figure 2. Maximum likelihood (ML) tree of the mtDNA control region haplotypes from *Oxyeleotris marmoratus*. The bootstrap proportions (>50%) in 1,000 replications were shown as nodes



0.001

uneditc