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Mitochondrial Gene Introgression between Spined Loaches via Hybridogenesis

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ABSTRACT—This report deals with an unusual mode of mitochondrial gene introgression between *Cobitis* hankugensis (*C. sinensis*) and *C. longicorpus* which is mediated by a unisexual hybridogenetic system of diploid-triploid *C. hankugensis-longicorpus* complex. Mitochondrial DNA sequences of 3329-3330bp encompassing from upstream ND6 to 12S rDNA indicated that mitochondrial genomes from the diploid hybrids, triploid hybrids, and their parental species are almost identical. Because triploid hybrids produce haploid ova with *C. hankugensis* chromosome set, normal diploid *C. hankugensis* regenerates upon insemination with *C. hankugensis* sperm. If the hybrid carries *C. longicorpus* mitochondrial genome, the regenerated *C. hankugensis* is a nucleo-cytoplasmic hybrid, thus accomplishing the unusual mode of mitochondrial gene introgression.

Key words: polyploidy, hybridization, unisexual reproduction, Cobitis

INTRODUCTION

Genetic introgression between closely related fish species is widely recognized (Smith, 1992; Mukai, 2001 and references therein). The process of gene introgression has been represented by production of fertile hybrid and backcross gradually incorporating genes into recipient populations upon genetic recombination. In this report we show an unusual, probably non-recombinant, and leaping mode of mitochondrial gene introgression which is mediated by a unisexual hybridogenetic system.

Some loaches (family Cobitidae, Osteichthyes) contain diploid-polyploid complexes (Kim and Lee 2000; Saitoh *et al.*, 2000 and references therein; Zhang and Arai, 1999). Occurrence of unisexual (all-female) populations of hybrid origin in some of these complexes is emphasized to be a source of establishment of gonochoric tetraploid population (Vasil'ev *et al.*, 1989), but no one except Kim and Lee (2000) recognized that normal diploid individual can be born from unisexual hybrids. Establishment of tetraploids via unisexual hybrids in loaches has been thought to be a one-way process.

A diploid-triploid hybrid complex occurs (*Cobitis hanku*gensis-longicorpus [*Cobitis sinensis-longicorpus*] complex)

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Note: Nucleotide sequence data reported are available in the DDBJ/ EMBL/GenBank databases under the accession number(s) AB120176-AB120177. in Nakdong River, Korea (Kim and Lee, 1990; Kim et al., 2003). The hybrid complex contains both diploid and triploid populations with few male occurrences. The diploid hybrid contains haploid genomes from C. hankugensis and C. longicorpus each, and the triploid does two haploid genomes from C. hankugensis and one from C. longicorpus (Kim and Lee, 1990). Artificial crossing experiment showed the diploid hybrid produces unreduced diploid hybrid ova, and the triploid does ova of C. hankugensis haploid genome eliminating C. longicorpus genome and reducing (Fig. 1) (Kim and Lee, 2000). Then, normal diploid C. hankugensis regenerates from the hybridogenetic triploid, crossing with male C. hankugensis. The process of establishment of polyploid populations thus may not be a one-way process. If so, this hybrid complex can mediate mitochondrial gene introgression from C. longicorpus to C. hankugensis.

MATERIALS AND METHODS

We have sequenced a portion of mitochondrial genome (3329– 3330 bp) from two female *C. hankugensis*, three diploid hybrids (one male and two females), three triploid hybrids (one male and two females), and two female *C. longicorpus* individuals. Species and ploidy diagnosis followed Kim and Lee (1990) employing morphological and chromosomal examination. These loaches came from Inwol-myon, Namwon-gun, Chollabuk-do, Korea (127°35'E, 35°27'N) except one *C. hankugensis*. The other *C. hankugensis* being examined as a comparative material was from Seangchomyon, Sanchong-gun, Gyeongsangnam-do, Korea (127°50'E, 35°27'N). Both collecting sites are in the Nakdong River basin. The sequenced region encompasses from upstream NADH dehydroge-



Fig. 1. Reproductive mode of *Cobitis hankugensis-longicorpus* complex. Single letters stand for *C. hankugensis* (H) or *C. longicorpus* (L) haploid genome. Solid arrows indicate experimental hybridization (Kim and Lee 2000), while dotted arrows denote presumed pathways. Large circles indicate eggs or oocytes. Haploid genomes being eliminated or released as polar bodies are set in small circles.

Table 1.	PCR and	sequencing	primers	used ir	ı this	study	1
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Name/ Position*	Seq	uence	(5'->	3')						
L12321**	GGT	CTT	AGG	AAC	CAA	AAA	CTC	TTG	GTG	CAA
L14279	GAA	TAC	ATY	ARA	GCT	ACC	CCA	С		
L14504	GCC	AAW	GCT	GCW	GAA	TAM	GCA	AA		
L14547	AGG	CGC	CGG	GTT	AGA	AGC	AAC			
L14760	AAC	СТС	TAA	TGG	CAA	GCC	TAC	G		
H14834	GAG	CCA	AAG	TTT	CAT	CA				
L15007	AAC	ATA	CAT	GCC	AAC	GGA	GC			
H15149	GGT	GGC	KCC	TCA	GAA	GGA	CAT	TTG	KCC	TCA
L15441	TCA	TAT	AAA	GAC	TTA	TTA	GGC	TT		
H15680	CGG	AAT	GTT	AGT	ССТ	CGT	ΤG			
L15926	TGA	AAG	CGC	TGG	тст	TGT	AAT	СС		
L15936	GGT	CTT	GTA	ATC	CGA	AGA	TCG	GAG	GTT	AAA
H15986	TAG	TTT	AGT	TTA	GAA	TTC	TGG	CTT	TGG	GAG
L16019	GCT	ACC	AAA	GCC	AGA	ATT	СТА	Α		
L16019a	GCT	CCC	AAA	GCT	AGT	ATT	CTA	А		
L220D***	AAT	TAC	TAA	GGT	GTG	CAT	AAG	тс		
L317D***	TCA	TGC	ATG	ATA	GAA	CCA	GGG	AC		
H342D***	AAC	CAG	ATG	CCA	GTA	ATA	GTT	С		
H594D***	ATA	TGC	AAT	GCT	TAA	GTT	ATG	тс		
H732D***	TTT	MGG	GGT	TTG	ACA	AGG	ATA			
L620	AAA	GCK	TAG	TAC	TGA	AGA	TGT	ТА		
H651	ATA	AGG	TCG	GGA	CCA	TGC	СТ			
H690	GCG	GAG	GCT	TGC	ATG	TGT	Α			
H721	CGG	GCA	GGG	GAT	TGA	GGG	CAT			
H884	AAC	CGC	GGT	GGC	TGG	CAC	GAG			
L1083	ACA	AAC	TGG	GAT	TAG	ATA	С			
H1358	CGA	CGG	CGG	TAT	ATA	GGC				
H1631	ACA	GGA	тсс	GGA	TGT	CTT	стс	GGT	GTA	AG
H2990**	TGC	ACC	ATT	RGG	ATG	тсс	TGA	тсс	AAC	ATC

* Primer names begin with strand name which they are designed on (L or H), followed by sequence positions of the 3' ends corresponding to positions on the human mt genome (Anderson *et al.*, 1981) or to those on the carp mt genome (Huang *et al.*, 1994) for primers on the D-loop containing region (***).

** Primers used for long PCR.

nase subunit-6 (ND6) to small subunit ribosomal DNA, corresponding to nucleotide positions from 14260 to 1017 of *C. striata* mitochondrial genome (Saitoh *et al.*, 2003). We employed the two step PCR direct sequencing technique (Miya and Nishida, 1999; Kawaguchi *et al.*, 2001). About 7 kb region was first amplified from genomic DNA with long-PCR primer pair. The long-PCR products then worked as templates for short PCRs with combination of 27 primers (Table 1) for direct sequencing using a commercial kit



10 changes

Fig. 2. A maximum parsimony tree of two equally parsimonious topologies obtained by an exhaustive search without character weighting on PAUP* ver. 4.0b (Swofford 1998). Gaps are treated as the fifth character. See Fig. 1 for genome composition of three biotypes. Letters in parentheses indicate male (M) or female (F). A *C. hankugensis* individual with an asterisk was collected at a different collecting site (Seangcho-myon) from other nine individuals.

Table 2. Sequence variable sites (corresponding to *C. striata* [Saitoh *et al.*, 2003]) in

 Cobitis hankugensis-longicorpus complex

Nucleotide	1	111111111111111111111	1111111111111111	
	4	4444444555555555	55666666666666	
Position	2	67788999911223344	99000112234455	67788
of <i>C. striata</i>	5	72319667912893746	37277561327812	33302
	9	27894035662135811	46701880960669	15768
		* *		
Gene	ND6	6 Cyt-b	CR	12S
HH(F)**,***	С	TGATAACCTTTGTTCAC	TGTGAAGTCTTTAA	ACACC
HH(F)	Т	CACCGTTCCCCATCTGT	CACAGGT-TGTCGT	GTGTT
HL(M)	Т	CACCGTTTCCCATCTGT	CACAGGT-TGTCGT	GTGTT
HL(F)	Т	CACCGTTCCCCATCTGT	CACAGGT-TGTCGT	GTGTT
HL(F)	Т	CACCGTTCCCCATCTGT	CACAGGT-TGTCGT	GTGTT
HHL(M)	Т	CACCGTTCCCCACCTGT	CACAGGTTTGCCGT	GTGTT
HHL(F)	Т	CACCGTTCCCCATCTGT	CACAGGT-TGTCGT	GTGTT
HHL(F)	Т	CACCGTTCCCCATCTGT	CACAGGTTTGCCGT	GTGTT
LL(F)	Т	CACCGTTTCCCATCTGT	CACAGGT-TGTCGT	GTGTT
LL(F)	Т	CACCGTTCCCCATCTGT	CACAGGT-TGTCGT	GTGTT

* Non-synomymous substitution.

** See Fig. 1 for genome composition of three biotypes. Letters in parentheses indicate male (M) or female (F).

*** C. hankugensis from different locality from the others.

(Amersham, Bucks, UK) and an ABI373S automated DNA sequencer (ABI, Norwalk, USA).

RESULTS

Nine individuals of *C. hankugensis*, *C. longicorpus*, and hybrids from Inwol-myon turned out to be very close or identical (none to one nucleotide gap, none to three transitions, and no transversion) in their mitochondrial DNA sequences regardless of sex (Table 2). On the other hand one *C. hankugensis* individual from a different collecting site carried a heterogenic mitochondrial genome with 28–30 transitions, five transversions, and two amino-acid substitutions being observed between this individual and other nine individuals. Number of estimated nucleotide substitutions per site (Kimura, 1980) (transition/transversion ratio=5.71) between the two *C. hankugensis* individuals from different localities was 0.01. On the other hand, the values between individuals from the same locality were 0–0.0009 (average=0.0003) regardless of their biotypes.

We could not find any sequence differences between one *C. hankugensis*, one *C. longicorpus*, two diploid hybrids and one triploid hybrid individual. Similarly, one *C. longicorpus* and one diploid hybrid individual were identical in mitochondrial DNA sequences. A maximum parsimony tree with *C. striata* sequence as an outgroup showed a nested distribution of the three biotypes in the tree (Fig. 2).

DISCUSSION

Sequence divergence between Cobitis species is so far

reported to range between 4.6 to 19.2% (Kim *et al.*, 2000; Perdices and Doadrio, 2001; Kitagawa *et al.*, 2001). Sequence divergence between *C. hankugensis* individuals from different localities actually was 1% indicating loach populations are localized and prone to diverge even within a single basin. From an empirical view taking these reports and our result into account, it is unusual that two morphologically and cytologically (Kim and Lee, 1990) distinct species carry mitochondrial genomes with little sequence divergence. Lineage sorting is unlikely over a geological timescale.

One possible explanation is the diploid-triploid hybrid complex as a vehicle of mitochondrial genome between two parental diploid species. If the mother of the initial diploid hybrid was C. longicorpus, it transferred the C. longicorpus mitochondrial genome to triploid hybrids of the next generation (Fig. 1) (Kim and Lee, 2000). The triploid hybrids produce ova with C. longicorpus mitochondrial genome and C. hankugensis haploid chromosome set which sometimes presumably accept C. hankugensis sperm in the natural habitat. Since hybridogenetic complexes show no genetic recombination between heterospecific genomes (Graf and Pelaz, 1989; Schmidt, 1996, but see Mateos and Vrijenhoek, 2002), next generation individuals would be nucleo-cytoplasmic hybrids between C. hankugensis and C. longicorpus. This pathway can accomplish hereby the unusual, probably non-recombinant, and leaping mode of mitochondrial gene introgression. Ecological study is necessary focusing on a mate recognition system between the hybrid complex and their parental diploid species.

Our study sheds light over unusual mitochondrial

grouping among some diploid fish species. Carmona *et al.* (1997) observed unusual mitochondrial clustering of minnows and postulated a hybrid origin or ancient lineage sorting. Kitagawa *et al.* (2001) postulated mitochondrial genome exchange between two *Cobitis* lineages. Introgression events at the diploid level (hybridization and backcrossing) may be responsible for such mitochondrial clustering or genome exchange, but also hybridogenesis can mediate gene introgression. We should especially consider the latter possibilities in minnows and loaches, because unisexual reproduction and polyploidy of hybrid origin occur frequently in these fish groups.

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