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Studies on the molecular mechanism of sex determination in the cobaltcap silverside Hypoatherina tsurugae

メタデータ	言語: eng		
	出版者:		
	公開日: 2018-01-05		
	キーワード (Ja):		
	キーワード (En):		
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URL	https://oacis.repo.nii.ac.jp/records/1488		

## [課程博士・論文博士共通]

## 博士学位論文内容要旨 Abstract

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論文題目 Title	Studies on the molecular mechanism of sex determination in the cobaltcap			
	silverside Hypoatherina tsurugae			
	(ギンイソイワシ Hypoatherina tsurugae の性決定の分子機構に関する研究)			

The sex-determining gene *amhy* (Y chromosome-linked Anti-Müllerian hormone) has been confirmed in Atherinopsid species of the genus *Odontesthes* (*O. hatcheri* and *O. bonariensis*) which occur naturally in inland waters of Argentina, Brazil and Uruguay. The presence of *amhy* in other families of Atheriniforms besides Atherinopsidae is unknown. In order to examine the evolution of *amhy* in this order, I selected the cobaltacap silverside *Hypoatherina tsurugae*, which inhabits the coastal waters of Japan and the Korean peninsula, as a model. The cobaltcap silverside belongs to the family Atherinidae, which is phylogenetically and geographically the most distant family from the Atherinopsidae where *amhy* has been discovered. In addition to the insight on the evolution of *amhy*, this study aims to understand the genetic mechanism of sex determination in *H. tsurugae*, in particular to clarify the role(s) of *amhy* in gonadal sex determination/differentiation and its relation to other sex related key genes (*sox9*, *cyp19a1a*, etc) in these processes.

In the 1<sup>st</sup> chapter, I searched the genome of *H. tsurugae* for the presence of *amhy* gene. The complete gene structure of *amhy* and its somatic homologue *amha* were obtained and compared. The *amha* gene is composed of 2,015 nucleotide bases and seven exons. The TGF-β domain is present in Exon 7 as in other species. The *amhy* gene is composed of 1,838 nucleotide bases and has only 4 exons. Exons 2 and 3 are completely lacking in the *amhy* gene structure. A specific insertion of 195 nucleotide bases is present at the place of exons 2 and 3. The exon 5 sequence is found in genomic sequence but it is not translated. The nucleotide identity between exons of *amha* and *amhy* was more than 80%. The deduced amino acid sequence of Amha (511 aa) and Amhy (340 aa) shared 91% identity. Exons 1, 4, 6 and 7 of *amhy* showed identity to those of *amha* as follows: 71%, 99%, 95% and 81% respectively. The exon 7, which contains the TGF-β domain, shared 93% amino acid identity with the similar domain of *amha* and contains 7 canonical cysteine residues that form disulfide bonds to make cysteine knots during dimer formation.

In the 2<sup>nd</sup> chapter, the role of *amha* and *amhy* gene in gonadal sex differentiation of *H. tsurugae* was examined. The linkage between *amhy*+ genotype and sex phenotype was analyzed in wild samples as well as in laboratory reared fish that were raised from hatching at the average temperature of the spawning season of *H. tsurugae*. The PCR analysis yielded a linkage with male sex in 95% and 85% of the wild adults and the laboratory-reared progeny, respectively. The temporal expression of *amha* and *amhy* gene was studied by qRT-PCR. In *amhy*+ fish, the *amhy* gene is highly expressed during early sex differentiation period while the *amha* gene expression is very low until the early juvenile stage. The spatial expression of *amha* and *amhy* was studied by ISH. This analysis showed that at 4 wah (undifferentiated period), *amhy* is expressed in somatic cells surrounding germ cell. In contrast, signals of *amha* could not be detected at this time. These results indicate that the *amhy* gene in *H. tsurugae* is tightly linked to the phenotypic sex and that it is highly expressed in the gonads of *amhy*+ individuals during the early development of the gonads. Thus, *amhy* can be considered as a strong candidate for sex determining gene in this species.

In the 3<sup>rd</sup> chapter, the objective was to study the gene expression profile of sex-related genes that could be adjuvants to *amhy* gene in sex determination/differentiation of *H. tsurugae*. The expression of six key sex differentiation genes (sox9, dmrt1, gsdf, foxl2, cyp19a1a and scp3) in amhy+ and amhy- individuals of *H. tsurugae* were studied during the early stages of gonadal development. Most of the genes showed a dimorphic expression related to sex genotype (amhy+ / amhy-) with exception of sox9. The reason for the lack of sex dimorphism in sox9 expression during this period may be that this gene is necessary for proliferation of germ cells in both sexes, as shown for other species.

In conclusion, I successfully isolated, cloned and sequenced the *amhy* gene in *H. tsurugae*. This gene is tightly linked with the male phenotype and highly expressed during early gonadal sex determination /differentiation. The sex related key genes *dmrt1*, *foxl2*, *cyp19a1a* and *scp3* showed dimorphic expression and an apparent synchronization with *amhy* gene expression. Future studies should look in more detail about their relations with as well as their regulation by *amhy* in order to corroborate the status of sex determining gene for *amhy* in *H. tsurugae*.