Inverted migration of rare whisker sheatfish in Nong-Han Lake, northeastern Thailand: Implications for conservation

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ABSTRACT. Nong-Han Lake, Thailand, sustains the whisker sheatfish (Micronema bleekeri Günther, 1864), which is a rare species of freshwater catfish. Wild-caught whisker sheatfish has been intensively harvested to meet market demand; yet, genetic information about this species remains unknown. To assist with the *in situ* conservation of whisker sheatfish populations in Nong-Han Lake, 35 and 34 individuals from the middle (MN) and lower (LN) areas of the lake, respectively, were studied using 7 microsatellite loci. Low genetic variation was detected in the MN ($H_O = 0.338, A_R = 2.710$) and LN ($H_O = 0.394, A_R = 2.714$) populations. Genetic differentiation between the 2 populations was significant ($F_{ST} = 0.063, P < 0.05$). The size of recent populations ($N_e < 50$) was found to be 9- to 29-times smaller compared to the estimated historical populations, even though no bottleneck signal was observed. Low genetic diversity was observed, implying that the
populations are at risk of being lost from this site. Of note, migration among the populations inhabiting the middle and lower parts of the lake exhibited opposing trends in changes to the genetic structure. This phenomenon might be due to the operation of a regional irrigation gate over the last decade. The information collected here indicates that the whisker sheatfish populations in Nong-Han Lake require consistent fisheries monitoring and management. Further research about the whisker sheatfish populations from the Mekong and Chao Phraya River basins is required to assist national-scale conservation efforts.

Key words: Effective population size; Microsatellite; Migration; Nong-Han Lake; Whisker sheatfish

INTRODUCTION

The whisker sheatfish (Micronema bleekeri Günther, 1864) is a rare freshwater catfish. The meat of this species is appreciated by local people, making it an economically important fisheries resource. In Thailand, the whisker sheatfish occupies a wide variety of habitats, including rivers, tributaries, floodplains, and swamps. This species is almost completely harvested from natural sources; however, attempts to develop aquaculture programs have been made to meet the high market demand for this species, but with limited success. Consequently, whisker sheatfish populations have been subject to an inevitable decline, primarily caused by over-fishing, habitat degradation, and reduced natural habitat water levels due to high water use for agricultural irrigation. Moreover, this species was recorded as rare at this site in the report of the Sakon Nakhon Inland Fisheries Research and Development Center (Ngoichansri et al., 2002; Daungsawat et al., 2003).

Nong-Han Lake (17°06ꞌ-17°15ꞌN, 104°07ꞌ-104°20ꞌE) is the largest inland water resource of northeastern Thailand. This lake has a high diversity of freshwater fish species, with at least 44 species of fish being taxonomically recorded to date (Jintanugool and Round, 2011). Nong-Han Lake is also one of the sites that have been selected for the implementation of the Large Swamp Inland Fisheries Project, which aims to increase fish productivity through intensive stocking. The major outflow of the lake enters the Nam Kum basin, which further drains southeast to the world’s great Mekong River. The water level of Nong-Han Lake is regulated by the Nam Kum Gate, which is composed of 1 spillway and 2 sluice gates that have been operated for over 15 years. The average depth of the lake is 1.9 m, with a maximum depth of 4.3 m. The water level of Nong-Han Lake fluctuates considerably. At the peak of summer, some marginal areas are completely dry, while other areas are very shallow; thus, creating geographical barriers among fish populations (Jintanugool and Round, 2011).

Microsatellites represent a well-known DNA marker. A microsatellite consists of simple sequences of a unit with 1-6 bases that may be iterated to lengths of up to 100 base pairs (Ellegren, 2004). Changes in the length of microsatellite DNA are generally thought to arise from slippage during the replication process, leading to insertions or deletions of repeat units relative to the template strand (O’Connell and Wright, 1997). Replication slippage also occurs during the polymerase chain reaction amplification of microsatellite sequences in vitro. This type of marker represents a promising tool for a wide range of genetic studies, including genetic linkage map-
ping, conservation genetics, aquaculture, and population genetics (Quan et al., 2006). Microsatellite markers have advantages over other molecular markers because of their locus-specific and co-dominant characteristics. These markers are generally assumed to evolve neutrally (Ellegren, 2004), to be highly polymorphic, and to be widely distributed throughout the fish genome. Only small amounts of DNA are required for microsatellite amplification using PCR and, importantly, the genotypes of microsatellites are highly reproducible (O’Connell and Wright, 1997). Microsatellites have been successfully used to estimate genetic variation in wild and hatchery stocks of many fish species (Hogan and May, 2002; Na-Nakorn et al., 2006; So et al., 2006). In Thailand, several freshwater catfish species have been subject to genetic and population structure studies, including the walking catfish (Na-Nakorn et al., 2004), the striped catfish (Na-Nakorn and Moeikum, 2009), and the giant catfish (Na-Nakorn et al., 2006; Ngamsiri et al., 2006); however, such studies have not been conducted on wild populations of whisker sheatfish. Knowledge about the genetic variation and population characteristics of the whisker sheatfish is thus currently unavailable; yet, it is important to obtain genetic information about this species to assist effective fishery management programs and to safeguard the genetic diversity of the species.

This study aimed to investigate the genetic diversity and population characteristics of the whisker sheatfish in Nong-Han Lake, northeastern Thailand.

MATERIAL AND METHODS

Sample collection and DNA extraction

Thirty-five and 34 whisker sheatfish specimens were collected from the middle (MN) and lower (LN) areas of Nong-Han Lake, Sakon Nakhon Province, Thailand during October-December, 2008. The collection sites were based on Ngoichansri et al. (2002) (Figure 1). Information about sample collection is presented in Table 1. Species identification was based on Rainboth (1996). Tissue samples (about 10 mg) were collected from the distal part of the caudal fin of each fish specimen, and were then preserved in absolute ethanol until DNA extraction. DNA was extracted from samples using an Aqua Pure Genomic DNA Isolation kit (Bio-Rad, California, USA), and analyzed by gel electrophoresis on 1.0% agarose gel stained with ethidium bromide, and compared with the Gene Ruler™ DNA ladder mix (Fermentas, Vilnius, Lithuania).

Microsatellite genotyping

The whisker sheatfish samples were genotyped at 7 microsatellite loci (MB79, MB81, MB153, MB320, MB401, MB456, and MB645), as described by Phongkaew et al. (2011). PCR reactions were carried out in 12.5 μl volumes containing 50 ng of genomic DNA, 20 mM Tris-HCl, pH 8.4, 50 mM KCl, 1.5 mM MgCl₂, 2.0 mM dNTP, 2 pmol of each primer, and 1U Taq DNA polymerase (Invitrogen, Oslo, Norway) on an XP cycler (BIOER, Hangzhou, China). The thermal profile for all loci involved initial denaturation at 94°C for 5 min, followed by 35 cycles at 94°C for 45 s, annealing at 53°C-61°C for 30-45 s (depending on the locus), and at 72°C for 30-45 s, and a final extension at 72°C for 5 min. The PCR products were subsequently separated on 6% (w/v) denaturing polyacrylamide gel (Bio-Rad, California, USA), and visualized after silver-staining. The alleles were sized based on the 50 bp standard marker (Fermentas).
Inverted migration of the Nong-Han Lake whisker sheatfish

<table>
<thead>
<tr>
<th>Collection site</th>
<th>Latitude/ Longitude</th>
<th>Sampling year</th>
<th>Number of samples</th>
<th>Microsatellite diversity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Middle area of Nong-Han Lake (MN)</td>
<td>17°11’20.05’’N, 104°12’46.19’’E</td>
<td>2008</td>
<td>35</td>
<td>2.714</td>
</tr>
<tr>
<td>Lower area of Nong-Han Lake (LN)</td>
<td>17°09’26.92’’N, 104°12’37.20’’E</td>
<td>2008</td>
<td>34</td>
<td>2.714</td>
</tr>
<tr>
<td>Mean</td>
<td></td>
<td></td>
<td></td>
<td>2.714</td>
</tr>
</tbody>
</table>

A = Number of alleles per locus; A_e = effective number of alleles per locus; A_r = allelic richness; H_o = observed and H_e = expected heterozygosity; F_is = degree of inbreeding.

Table 1. Summary of genetic variability across microsatellite loci of whisker sheatfish populations in the middle (MN) and lower (LN) area populations of Nong-Han Lake.

Figure 1. Collection sites of whisker sheatfish samples in the Nong-Han Lake, Sakon Nakhon Province, Thailand. Middle area (MN) (17°11’20.05’’N, 104°12’46.19’’E) and lower area (LN) (17°09’26.92’’N, 104°12’37.20’’E). Map re-drawn from Ngoichansri et al. (2002).

Statistical analyses

**Microsatellite diversity, Hardy-Weinberg, and linkage disequilibrium**

Evidence for genotyping error and large allele dropout for each locus within each
population was assessed using MICRO-CHECKER 2.2.3 (van Oosterhout et al., 2004). Genotype data of whisker sheatfish from 7 microsatellite loci were used to calculate the effective number of alleles ($N_e$), observed heterozygosity ($H_o$), and expected heterozygosity ($H_e$) by the POPGEN 32 software package (Yeh and Yang, 1999). Allelic richness ($A_r$) was calculated using FSTAT 2.9.3.2 (Goudet, 2001). Deviations from Hardy-Weinberg and linkage equilibrium between pairs of loci were tested using Genepop 3.4 (Raymond and Rousset, 1995). Sequential Bonferroni correction (Holm, 1979) was employed to account for multiple testing.

**Genetic differentiation within and between populations**

To evaluate the extent of differentiation, the degree of inbreeding among individuals relative to the rest of their sub-population was calculated for loci based on $F_{IS}$, while inter-population differentiation was calculated based on the multilocus $F_{ST}$ estimator of Weir and Cockerham (1984) using FSTAT 2.9.3.2 (Goudet, 2001).

**Estimation of effective population size and migration rate**

The maximum likelihood-based (ML) coalescent Markov Chain Monte Carlo (MCMC) approach was used to infer the long term (i.e., ‘historical’) effective population size, $\theta (\theta = 4N_e\mu$; where $N_e$ is the effective population size; $\mu$ is the mutation rate per site) and the historical migration pattern, $M (M = m/\mu$; where $m$ is the immigration rate per generation) among whisker sheatfish populations using the program MIGRATE 3.2.6 (Beerli, 2008). For each locus in the dataset, the ML was run for 10 short and 1 long chains, with 50,000 and 500,000 recorded genealogies, respectively, after discarding the first 10,000 genealogies (burn-in) for each chain. SV AR 1.3 (Beaumont, 2004) was used to predict the mutation rate of whisker sheatfish that is currently unknown, but required for the calculation of $m$ (from $M\mu$ in MIGRATE 3.2.6). MSVAR simulations were run for $2 \times 10^8$ iterations, and the first 10% of the output was discarded as the burn-in period. Hence, the remaining output was used to estimate the mutation rate of whisker sheatfish.

The ‘recent’ effective population size was assessed from a temporal sample using an approximate Bayesian computation approach implemented in the program ONeSAMP 1.2 (Tallmon et al., 2008). Prior lower and upper bounds of $N_e$ were set at 2 and 1000, respectively, based on a preliminary survey using the linkage disequilibrium method (Hill, 1981) implemented in NeEstimator 1.3 (Peel et al., 2004).

To estimate recent migration and its direction, a Bayesian method based on multilocus genotypes implemented in BAYESASS was used (Wilson and Rannala, 2003). It is assumed that the loci in the source population were in linkage equilibrium. The method to estimate the posterior probabilities of the migration matrix among neighboring populations is based on MCMC.

**Population bottleneck**

Potential demographic changes or substantial reduction in population size might cause bottleneck effects on the populations of this species. To detect any substantial reduction in population size that commonly occurs in populations of highly exploited species, genetic signatures of the reduction was investigated using the $M$ ratio (Garza and Williamson, 2001). Wilcoxon’s
sign-rank test (Luikart et al., 1998a), and mode-shift test (Luikart et al., 1998b). $M$ statistic values were calculated by $M = k / r$, which simply measures the ratio of the number of alleles, $k$, to the range in allele size, $r$, across microsatellite loci. In addition, the Wilcoxon’s sign-rank test (Luikart et al., 1998a) and a mode-shift indicator test (Luikart et al., 1998b) were performed using BOTTLENECK 1.2.02 (Piry et al., 1999). In the Wilcoxon’s sign-rank test, simulations were completed with 95% confidence intervals (re-sampling for 10,000 replications). The heterozygosity that was expected in a population at mutation drift equilibrium, $H_{eq}$, was calculated under both a strict single step mutation model (SMM) and a realistic (Di Rienzo et al., 1994) two-phase mutation model (TPM), that included a 95% SMM and 5% multistep mutations model (IAM), with 12% variance in TPM being employed, following the recommendation of Piry et al. (1999).

RESULTS

Genetic diversity and genetic differentiation of whisker sheatfish

In total, 38 alleles were observed from the 2 sheatfish populations genotyped at 7 microsatellite loci. The number of alleles per locus ranged from 2 to 4. There were no null alleles, nor were any other genotyping errors detected among loci. No Hardy-Weinberg deviation and no linkage disequilibrium were found in the 2 populations after sequential Bonferroni correction ($P > 0.007$). For the MN population, the observed heterozygosity ($H_o$) was 0.338 and the expected heterozygosity ($H_e$) was 0.360. For the LN population, the $H_o$ was 0.394 and the $H_e$ was 0.424. The inbreeding coefficient of both the MN and LN populations was observed across loci (MN: $F_{is} = 0.060$, 95% confidence limits -0.181-0.285; LN: $F_{is} = 0.071$, 95% confidence limits -0.103-0.448). Means of allelic richness ($A_r$) were 2.710 and 2.714 for the MN and LN populations, respectively. A summary of genetic variability across microsatellite loci is shown in Table 1 and Table S1. The estimate of $F_{ST}$ between the 2 whisker sheatfish populations was significant with moderate genetic differentiation ($F_{ST} = 0.063$, $P < 0.05$).

Population bottleneck observation

As the whisker sheatfish is considered to be a rare, but exploited species, it is likely that there has been a reduction in population size. The $M$ ratio across loci was 1 (no bottleneck signal) for both MN and LN populations. Under strict SMM and TPM (95% SMM and 5% IAM), the Wilcoxon’s sign-rank test did not reveal any recent bottleneck signal in whisker sheatfish populations. The mode-shift test on allele frequency distributions did not display any evidence of bottlenecks.

Effective population size and migration rate: historical versus recent

The maximum likelihood-based coalescent Markov Chain Monte Carlo (MCMC) approach inferred that the mutation scaled population size parameter was higher in the LN population ($\theta = 0.510$) compared to the MN population ($\theta = 0.401$). Between the 2 populations, the mutation rate across loci was similar, at $2.7-2.8 \times 10^{-4}$ per generation. These estimates were translated to a historical effective population size ($N_e$) of 358 and 462 in the MN and LN populations, respectively (Figure 2).
The estimated recent effective population size using the approximate Bayesian computation method for the MN population was 41 (95% confidence interval: 26.56-76.59), whereas it was 16 (95% confidence interval: 11.31-27.84) for the LN population (Figure 2).

Using the maximum-likelihood approach, the historical migration rate of the MN to the LN population was 9.955, while movement in the opposite direction was 4.245. In contrast, the Bayesian method revealed a reduction in recent migration rates. The recent migration rate of the MN population to the LN population was 0.0387, which was 257 times lower compared to the historical migration rate. The recent migration rate of the LN population to the MN population was 0.2128, which was 19 times lower compared to the historical migration rate (Figure 2).

DISCUSSION

Whisker sheatfish is considered a rare species, with low genetic diversity being expected. In our study, 2 populations of whisker sheatfish in Nong-Han Lake exhibited low allele numbers (average $A_R = 2.714$), in addition to low observed and expected heterozygosity (average $H_O = 0.366$; $H_E = 0.392$), when compared to populations of other freshwater fishes described by De Woody and Avise (2000). When using 75 microsatellite loci, the authors obtained an average $A$ and $H_E$ of 9.1 ± 6.1 and 0.54 ± 0.25, respectively, for 13 species of freshwater fish. Moreover, So et al. (2006) obtained an average allelic richness of 9.9 for the migratory sutchi catfish. In addition, Na-Nakorn and Moeikum (2009) reported an allelic richness of 5.6 across the loci of striped catfish in Vietnam. Foulley and Ollivier (2006) suggested that allelic richness is particularly important over the long-term, because it better reflects past fluctuations in population size. In terms
of heterozygosity, Quan et al. (2006) obtained an average $H_o$ and $H_e$ of 0.43 and 0.75, respectively, for 22 wild individuals of northern sheatfish (*Silurus soldatovi*). Surprisingly, the genetic diversity of whisker sheatfish was lower compared to the critically endangered Mekong giant catfish, *Pangasianodon gigas*, observed by Na-Nakorn et al. (2006) and Ngamsiri et al. (2006). These authors obtained averages of 0.55 and 0.37 for $H_o$ and 0.49 and 0.44 for $H_e$, respectively, for these 2 critically endangered species. In addition, the overall $F_{IS}$ value among all loci of the whisker sheatfish was higher than zero, indicating a certain level of heterozygote deficit. This heterozygote deficiency might also be explained by selection, inbreeding, population substructure, and null alleles (Crow and Kimura, 1970). Because of small population size or small numbers of families, the positive $F_{IS}$ value indicated that inbreeding was one of the main causes for the shortage of heterozygotes in the whisker sheatfish population. This suggestion was concordant with many existing reports about other freshwater fishes. For instance, So et al. (2006) found a significant $F_{IS}$ value (0.028) at a spawning area of migratory sutchi catfish. Parra-Bracamonte et al. (2011) found that the degree of inbreeding increased from 0.23 to 0.27 over a 4-year period in a traditional catfish hatchery channel. In this study, we found that the whisker sheatfish in Nong-Han Lake had low genetic diversity, and had a high tendency of inbreeding.

Whisker sheatfish undertake lateral migrations. For example, the geography, landscape, water level of the water body, and the structures that are built to impound water might affect gene flow (Hanfling and Weetman, 2006). There are no previous reports about the movement pattern of whisker sheatfish populations within Nong Han Lake and between other watercourses. However, the lack of fish movement barriers in the past allowed whisker sheatfish populations in Nong-Han Lake to freely migrate to Nam Kum basin, and finally to the Mekong River. In our results, major historical migrations occurred from MN to LN populations (m$_{MN\rightarrow LN}$ = 9.955; m$_{LN\rightarrow MN}$ = 4.245). This result implied that gene flow among whisker sheatfish populations in Nong-Han Lake naturally moves southwards, which is the migratory route to the outlet of Nong-Han Lake leading to Nam Kum Basin and the Mekong River. Today, the migratory route from Nong-Han Lake to the Mekong River is obstructed by the Nam Kum gate (Jintanugool and Round, 2011), which only opened during the flood season (June to September). From year to year, environmental characteristics might change and influence migratory route or direction (Bunt, 2001). Therefore, whisker sheatfish were prevented from moving to areas downstream of the lake. The LN whisker sheatfish population has limited opportunities for downstream migration, resulting in it migrating in the opposite direction, i.e., upstream. The migration pattern showed that the recent migration rate from the LN to the MN population was more than 5 times greater compared to the opposite direction. During sample collection, it was observed that whisker sheatfish in Nong-Han Lake only moved between the lower and middle areas of the lake, and did not enter the upper area. The LN population was more influenced by the sluice gates compared to the MN population. Thus, the LN population of whisker sheatfish attempted to migrate back to the middle area of the lake, rather than migrating to the Nam Kum gate. This observation indicates that an anthropogenic barrier might dramatically alter migratory behavior, and cause a clear inversion in the direction of gene flow among populations of whisker sheatfish in Nong-Han Lake, as reported for other species of *Cottus gobio* (Hanfling and Weetman, 2006).

When the population size is small, as in the study populations, bottleneck signatures are expected. However, in our study, evidence of a recent genetic bottleneck was indicated by 3 different statistical methods. Hundertmark and van Daele (2010) suggested that populations with low levels of diversity might have undergone a bottleneck within a short period of time, and that
such populations might recover following a process described by Garza and Williamson (2001). The authors showed that, for a small population, $M$ continues to decline for approximately 50 generations before it begins to recover. However, the authors also found that even when the value of $M$ has partially recovered, allelic diversity does not recover in a small population. In addition, a significant reduction in whisker sheatfish population size was observed based on estimates of effective population size. The current study showed that whisker sheatfish populations had small historical effective population size ($MN = 358; LN = 462$), whereas the recent population size was reduced ($MN = 41; LN = 16$). In general, a population of $N_e = 500$ is considered to be large enough to maintain genetic diversity for key life-history traits; hence, genetic diversity would only appear to be depleted if $N_e < 500$ (Frankham, 1995). In addition, Franklin (1980) found that inbreeding depression is prevented when $N_e > 50$, whereas the viability or reproductive fitness of a population will decrease when $N_e < 50$. Therefore, at present, the whisker sheatfish population in Nong-Han Lake is vulnerable and at risk of population extinction within short period of time.

Various management strategies should be implemented to safeguard the rare whisker sheatfish populations of Nong-Han Lake, which have been subject to a genetic bottleneck and migratory obstruction. First, lake restoration should be conducted; for example, the gate should be opened for suitable time periods at times of the year relevant to the reproductive strategies of this fish population. Second, fish sanctuaries should be created for in situ conservation. Alternatively, stock enhancement practices should be taken, using offspring from wild spawners for artificial breeding programs. The relevant authorities should advise the reduction or cessation of fish harvests to allow population recovery. Routine genetic monitoring of remaining populations is highly recommended to assist with conservation management of the natural gene pool. Further genetic study involving large numbers of whisker sheatfish populations including both the Mekong River and the Chao Phraya River basins is also required. The whisker sheatfish outside Nong-Han Lake might be considered as a viable alternative genetic resource providing additional genetic diversity when Nong-Han whisker sheatfish are no longer self-sustainable. The conservation and management of the whisker sheatfish population should be conducted at the country scale, within which it might have the potential to become an important aquaculture species. It is therefore important to obtain data about catches for the management and conservation of fish stocks in Nong-Han Lake.

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Supplementary material

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Inverted migration of the Nong-Han Lake whisker sheatfish

2010.


