

Kin-structured subpopulations in Eurasian perch (*Perca fluviatilis* L.)

GABRIELE GERLACH*, UWE SCHARDT, REINER ECKMANN & AXEL MEYER

Department of Biology, PO Box 5560, University of Konstanz, D-78457 Konstanz, Germany

Based on ecological and behavioural studies it has been assumed that Eurasian perch (*Perca fluviatilis*) within one lake may not represent one panmictic population, but that they are subdivided into subpopulations. In order to investigate the genetic substructuring of populations, we used gene frequencies of five microsatellite loci to compare perch from six different sites from Lake Constance, Germany, and as outgroups perch from the lake Grosser Vätersee, Berlin, and two Swiss lakes, Lake Zurich and Lake Walensee. We examined whether homing behaviour of subadults to the spawning sites of their parents occurs and whether philopatric behaviour of adults results in significant population genetic substructuring. The distribution of genetic variation revealed two major, genetically distinct populations in Lake Constance: one in the eastern part of the lake and another in the western part ($G_{ST}=0.07$). Within each of these two populations, no further genetic substructuring, nor any indication of inbreeding could be detected, either because genetic exchange was sufficiently high or because the time since separation has been too short. Homing behaviour of subadults to parental spawning sites after having spent several weeks of their life cycle in the pelagic zone could not be detected. Instead, subadults stay within either the western or the eastern region of the lake. There is evidence that some shoals contain full- and half-sibs. Despite females spawning in close proximity to each other, some siblings stay together. This might suggest that perch possess kin preferences and kin recognition.

Keywords: genetic substructuring, kin aggregation, microsatellites, *Perca fluviatilis*, Percidae.

Introduction

Recently, analyses of genetic population structure in several aquatic species have generally found that individuals from different parts of a lake or tributary do not form a single panmictic population, but instead are typically genetically substructured into different local populations (Carvalho, 1993; Tessier *et al.*, 1997; Brunner *et al.*, 1998; Chenoweth *et al.*, 1998). It has been assumed that the number of these subpopulations is a function of the number of geographical or physical barriers encountered by a species during its life cycle (Sinclair, 1988). Genetic drift or differential selection regimes in these isolated local populations might play an important role in the evolution of new species (Dieckmann & Doebeli, 1999; Kondrashov & Kondrashov, 1999). Genetic isolation of local populations is caused, among other reasons, by philopatric behaviour and local mating of adults and homing behaviour of subadults which return to parental spawning sites after spending a part of their life cycle in different habitats.

Female perch over three years old spawn from April to June, attaching their eggs in up to 1 m long strands on stones and other structures in the littoral zone. After hatching, larvae stay for approximately two weeks in this area before they drift into the pelagic zone where they live for about one month (Wang & Eckmann, 1994). In July, after the abundance of pelagic zooplankton decreases, juvenile perch return to the shallow littoral zones. If homing behaviour existed, then individuals from a specific locality would be expected to be genetically more similar to each other and to show significant genetic difference from those of other localities.

To date, behavioural and ecological studies have led to the assumption of sedentary behaviour in adult Eurasian perch (*Perca fluviatilis*). For example, in Lake IJssel in the Netherlands, Willemsen (1977) marked and released adult *Perca fluviatilis* away from their capture site; later, he recaptured them at the place of first capture. Similar results were reported for perch from the English Lake Windermere (Kipling & Le Cren, 1984). Analysis of allozyme variation revealed the existence of two distinct populations in the northern and southern

*Correspondence. E-mail: gabi.gerlach@uni-konstanz.de

parts of Lake Windermere (Bodaly *et al.*, 1989). Perch from Lake Constance also carry different intensities of parasite infection according to their capture sites, suggesting that adult fish are resident, possibly philopatric, and have little contact with populations from other localities in the lake (Balling, 1992; Dieterich, 1998).

Homing behaviour spectacularly demonstrates the often astonishing ability of a wide range of animals for orientation and navigation. Atlantic and Pacific species of salmon return to their natal rivers after years at sea from a distance of hundreds or even thousands of kilometres (Hasler & Scholz, 1983; Gyllensten & Wilson, 1987). Similarly, Oceanic loggerhead turtles (*Caretta caretta*) and green turtles (*Chelonia mydas*) return to spawn at the beach where they hatched (Allard *et al.*, 1994; Bowen *et al.*, 1994; Bolten *et al.*, 1998), and many bird species return to their nesting places after crossing whole continents (Dingle, 1996). Breeding in similar environmental conditions as the parental generation did can increase reproductive success and therefore favours selection for homing behaviour. A DNA microsatellite analysis was undertaken to examine the genetic diversity of perch populations of different ages from several

localities in the littoral zone of Lake Constance and from three other lakes in Germany and Switzerland to evaluate genetic substructuring and homing behaviour.

Materials and methods

Sampling

Perch were collected at six different localities in Lake Constance (see Fig. 1) during the winter of 1997/1998. Additionally, perch from three other lakes were included in this study: Grosser Vätersee, near Berlin, Germany, and Lake Zurich and Lake Walensee, Switzerland. At locality 5, animals were sampled by electro-fishing along a reed belt over a distance of 400 m (Fig. 1). A trawl net was used to sample at locality 1; all other localities were sampled with bottom set gill nets. All animals per locality were of the same age; but the age of the perch differed among localities: 0 + (caught before one year): OG, KR, GN; 1 + (caught before 2 years of age): US, B; 2 + (caught before 3 years of age): LA, RO; 3 + (caught before 4 years of age): LW, LZ. DNA microsatellite analysis was performed on 18 individuals per locality.

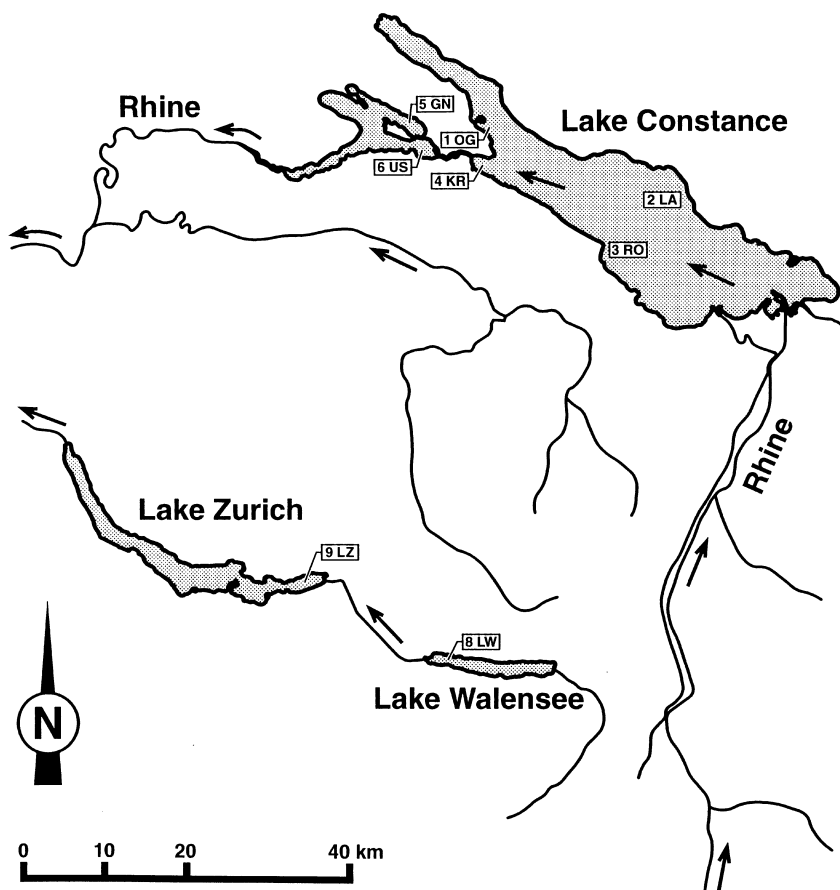


Fig. 1 Map of sampling localities of perch in Lake Constance: 1 OG Obere Güll, 2 LA Langenargen, 3 RO Romanshorn, 4 KR Kreuzlingen, 5 GN Gnadensee, 6 US Untersee, and in two lakes in Switzerland: 8 LW Lake Walensee and 9 LZ Lake Zurich. Arrows indicate the main current in Lake Constance.

Microsatellite analysis

Muscle and liver tissues were stored in 80% ethanol until DNA extraction. Genomic DNA was extracted according to standard phenol–chloroform procedures (Sambrook *et al.*, 1989). Polymerase chain reaction (PCR) amplification and genotyping were conducted for five dinucleotide microsatellite loci (CA and GT repeats), using primers developed for walleye (*Stizostedion vitreum*) (Borer *et al.*, 1999) (Table 1). Amplification products (1.8 μ L) were electrophorized on 6% denaturing polyacrylamide gels for 3.5–6 h and visualized using X-ray-sensitive films. The allele sizes found for each locus were determined and numbered according to their length (e.g. allele 1 being the smallest).

Data analyses

Genetic diversity was quantified in terms of heterozygosity, number of alleles per locus and allele frequencies observed in all local populations. GENEPOP version 1.2 (Raymond & Rousset, 1995) was used for estimating deviations from Hardy–Weinberg equilibrium at each locus and globally over all loci. Using FSTAT version 2.8 (Goudet, 1995) F_{IS} -values (Weir, 1990) and G_{ST} values were assessed by jackknifing procedures (1000 iterations) and used for calculations of relatedness within local populations. According to Wright (1969) who described genetic population structures in terms of the F -statistics, the relationship between F_{ST} , F_{IT} and F_{IS} can be written as $(1 - F_{IT}) = (1 - F_{ST})(1 - F_{IS})$. It is possible to partition the intraclass relatedness 'r' in the same way (Pamilo, 1984). The relatedness values in a subdivided population can be expressed as $(1 - r_{GT}) = (1 - r_{ST})(1 - r_{GS})$ where r_{GS} is the average relatedness

among individuals in social groups within subpopulations, r_{ST} is the average relatedness of individuals among subpopulations, and r_{GT} is the average relatedness of individuals in groups with reference to the total populations (Pamilo, 1989). Therefore, intraclass relatedness, i.e. genotypic correlation among group members, can be written as $r_{GS} = 2F_{GS}/(1 + F_{IS})$, where F_{GS} is the gametic correlation of group members within a subpopulation and F_{IS} measures inbreeding. Pamilo (1984, 1985) further proposed an inbreeding-corrected relatedness, which is estimated as: $Relatc = [r - 2Capf]/(1 + Capf)/[1 - 2Capf/(1 + Capf)]$ in the program FSTAT 2.8, where $Capf$ is an equivalent to F_{IT} . When the population is structured, $Relatc$ adequately removes the increase in relatedness due to this structuring (see Chapuisat *et al.*, 1997).

Although it is straightforward to estimate the average relatedness in social groups, estimating values for specific individuals with, e.g. microsatellite data is still problematic: On average, full-sibs share 50% of their alleles and half-sibs 25%, but it depends on the frequency of alleles within the population how informative sharing alleles at a single locus is. The frequency of each allele within the population has to be taken into account for classifying individuals as being related or not. Therefore, we did not use the value of sharing 50% of the alleles as the criterion but calculated the threshold for being full- or half-sibs according to the allele frequencies of perch in Lake Constance by extending a method described by Blouin *et al.* (1996). We randomly generated 1000 pairs of unrelated individuals and calculated the number of shared alleles using SAS version 6.12 (SAS Institute, 1986). To evaluate the number of shared alleles of full-sibs we randomly chose 1000 pairs of individuals, generated two offspring each and

Table 1 Microsatellite characteristics. H_{obs} (observed heterozygosity), H_{exp} (expected heterozygosity). Perch examined were caught at nine different localities of Lake Constance, Lake Walensee, Lake Zurich (both situated in Switzerland) and Grosser Vätersee (near Berlin)

Locus	No. of individuals	No. of alleles	% of most common allele	H_{obs}	H_{exp}	Primer sequences
<i>Svi6</i>	162	6	37.7	0.39	0.54	F: CATATTATGTAGAGTGCAGACCC R: TGAGCTTCACCTCATATTCC
<i>Svi17</i>	162	7	37.7	0.86	0.74	F: GCGCACTCTCGCATAGGCCCTG R: CGTTAAAGTCCTTGGAAACC
<i>Svi18</i>	162	7	40.1	0.82	0.74	F: GATCTGTAAACTCCAGCGTG R: CTTAAGCTGCTCAGCATCCAGG
<i>Svi26</i>	162	10	82.4	0.15	0.31	F: CAGACAAACAGGTTGGAGAG R: CTAATTATCTTCTGGCGGAC
<i>Svi29</i>	162	4	69.9	0.58	0.45	F: GATCCCCCTCCTCTCCCCTC R: CTTTTTCTGCCCTGTCCCAAC

calculated the matches of alleles between them. We used a similar procedure to generate the mean number of matched alleles for half-sibs. Dividing by the number of loci gives M_{xy} , the average number of matches per locus between individuals x and y . Using a logistic regression analysis (SAS Institute, 1995) we calculated the probabilities for each M_{xy} to belong to either of the three different groups (unrelated, half- or full-sibs) analogous to a likelihood ratio principle. The threshold for being classified as unrelated was $M_{xy} \leq 0.5$, as half-sibs $0.5 > M_{xy} < 0.7$, and as full-sibs $M_{xy} \geq 0.7$. In a subsequent analysis we calculated M_{xy} for each pair of individual perch per sampling site and determined the number of full-sibs and half-sibs according to these thresholds. We determined whether the observed number of full- and half-sibs at each locality was higher than would be expected from combinations of unrelated animals which were misclassified as being related. The analysis was performed by 1000 simulations of constructed matrices which were derived by creating 18 virtual individuals on the basis of the gene frequency of perch in Lake Constance and comparing their alleles pairwise as described above. The performed significance test based on the null hypothesis that there were no full- or half-sibs among the sampled individuals. For each locality it was calculated if the observed numbers of full- and half-sibs were significantly different from this distribution.

Pairwise F_{ST} -values (θ -values) (Weir, 1990) were determined based on the gene frequencies using an analysis of molecular variance (AMOVA) (Michalakis & Excoffier, 1996) available in the program ARLEQUIN version 1.1 (Schneider *et al.*, 1997). Probability values of the pairwise F_{ST} -values were adjusted for multiple simultaneous table-wide tests using the sequential Bonferroni adjustments (Rice, 1989) to minimize type-I errors. Genetic distance values were analysed according to Reynolds *et al.* (1983) with $D = -\ln(F_{ST})$ calculated with the program GENETIX (Belkhir *et al.*, 1997). Mantel

matrix correlations (Mantel, 1967) were used to compare geographical (straight-line distance between different localities) and genetic distance D within Lake Constance.

Results

Population substructuring

Comparing all perch populations from Lake Constance, a F_{ST} -value of 0.054 was calculated. Genetic subdivision between local populations was examined by calculating pairwise F_{ST} -values (Table 2). According to these data, perch sampled from six different localities in Lake Constance can be subdivided into two large populations. The first population is composed of individuals from the east (localities 1, 2 and 3) and the second in the western part includes specimens from localities 4, 5, and 6 (Fig. 1). The G_{ST} value (corrected) for all loci between these two populations is 0.07 ($P < 0.001$) indicating a moderate genetic substructuring. Whereas perch from localities 1 and 2 differed significantly from those from localities 4, 5, and 6 the genetic exchange between localities 3–4 and 3–6 was higher. F_{ST} -values between these localities were moderate (0.051 and 0.048, respectively), but after Bonferroni corrections these values proved not to be statistically significant.

Perch from both Swiss lakes, Lake Walensee and Lake Zurich, were genetically similar ($F_{ST} = 0.022$, $P < 0.30$). Interestingly, perch from Lake Zurich and Lake Walensee were genetically more similar to animals from the eastern part of Lake Constance (localities 1, 2, and 3) ($F_{ST} = 0.054$, $P < 0.001$) than to those from the western part (localities 4, 5, and 6) ($F_{ST} = 0.112$, $P < 0.001$). As expected, the highest and most statistically significant degree of genetic subdivision was observed between perch from the lake near Berlin and all other localities (mean F_{ST} -values = 0.256, $P < 0.001$). A phenogram

Table 2 Pairwise F_{ST} -values between subpopulations of perch in Lake Constance (1–6); Grosser Vätersee, Berlin (7); Lake Walensee, Switzerland; (8); and Lake Zurich, Switzerland (9). Asterisks indicate subpopulations which showed a statistically significant genetic differentiation after sequential Bonferroni adjustments (Rice, 1989)

	1 OG	2 LA	3 RO	4 KR	5 GN	6 US	7 B	8 LW	9 LZ
1 OG	0								
2 LA	0.017	0							
3 RO	0.005	-0.004	0						
4 KR	0.078*	0.092*	0.051	0					
5 GN	0.079*	0.104*	0.073*	0.041	0				
6 US	0.069*	0.089*	0.048*	0.002	0.007	0			
7 B	0.27*	0.236*	0.229*	0.256*	0.278*	0.263*	0		
8 LW	0.067*	0.07*	0.06*	0.137*	0.091*	0.105*	0.251*	0	
9 LZ	0.04*	0.069*	0.054*	0.161*	0.115*	0.12*	0.295*	0.022	0

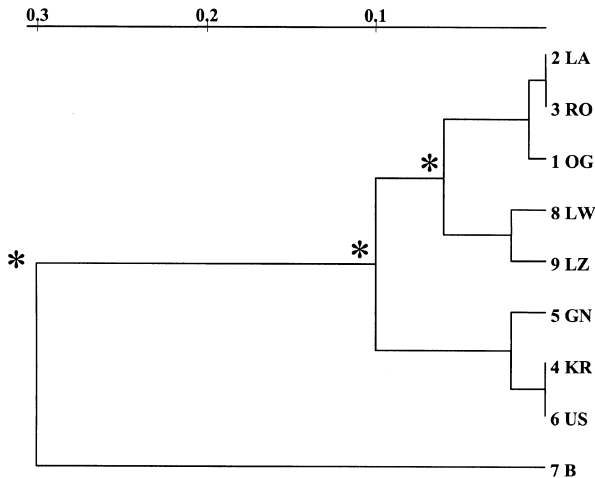


Fig. 2 Phenogram of genetic distances constructed by UPGMA between all investigated populations of perch. *Indicates statistically significant differences between the branches.

(Fig. 2) illustrates the different genetic distances (D -values) among perch from all localities studied.

When genetic distances of populations from Lake Constance were plotted against the logarithm of geographical distances between populations, a significant positive correlation (Mantel test; $R^2 = 0.26$; $P < 0.001$) was found, indicating an isolation by distance effect (Slatkin, 1993).

Genetic structure of local populations

The degree of allelic polymorphism at five microsatellite loci varied between four and 10 alleles per locus

(Table 1). Some alleles were only found in perch from one locality or a specific lake (e.g. alleles 1 and 4 of locus *Svi6* found only in Lake Constance; alleles 3 and 5 of locus *Svi26* found in the Grosser Vätersee near Berlin).

F_{IS} -values varied from 0.068 to -0.276 and showed a statistically significant excess of heterozygotes in six out of nine subpopulations, as illustrated in Table 3. On average, over all loci and local populations of Lake Constance $F_{IS} = -0.117$. The mean degree of relatedness between individuals per locality was $r = 0.115$ using the calculation of Queller & Goodnight (1989) and $Relatc = 0.674$ (relatedness corrected) (Pamilo, 1984, 1989; Goudet, 1995), indicating that animals collected at each locality were closely related. In order to test for kin aggregations we calculated pairwise genetic relatedness between all individuals of each locality using a program based on the SAS routine IML version 6.12 (SAS Institute, 1986) according to Blouin *et al.* (1996). Based on the distribution of allele frequencies of the total population of Lake Constance, individuals were classified as unrelated when $M_{xy} \leq 0.5$, as half-sibs when $0.5 < M_{xy} < 0.7$ or as full-sibs when $M_{xy} \geq 0.7$. In four out of six localities the number of individuals which shared more than 50% of the alleles (classified as full- and half-sibs) or 70% of the alleles (full-sibs) was significantly higher than in randomly generated groups of 18 unrelated individuals. Therefore, we concluded that perch indeed stayed in groups of kin. Only at localities 3 and 4 did the number of observed pairs of animals classified as full-sibs not exceed the expected values.

Table 3 F_{IS} -values and kin structure for each subpopulation. The proportion of randomization that gave a larger F_{IS} -value than the observed was used to find significant deviations from Hardy–Weinberg equilibrium (FSTAT version 2.8) (Goudet, 1995). The number of full-sibs ($M_{xy} \geq 0.7$) was determined by comparing alleles of the 18 individuals sampled at each locality (18 pairwise comparisons = 153, i.e. at locality 2 LA 88 out of 153 pairwise comparisons showed a M_{xy} value ≥ 0.7). The number of full- and half-sibs was determined using the criterion $M_{xy} > 0.5$. Relatedness of perch at localities 7, 8, and 9 was not considered

Local populations	1 OG	2 LA	3 RO	4 KR	5 GN	6 US	7 B	8 LW	9 LZ
F_{IS}	-0.204	-0.276	-0.060	0.068	-0.267	0.000	-0.182	-0.121	-0.275
P -value	0.012	0.001	0.274	0.878	0.002	0.580	0.004	0.035	0.002
	*	***	NS	NS	**	NS	**	*	**
No. of full-sibs	65	88	40	32	84	49	—	—	—
P -value	<0.001	0.036	0.19	0.44	<0.001	0.043	—	—	—
	***	*	NS	NS	***	*	—	—	—
No. full- + half-sibs	97	126	76	57	129	92	—	—	—
P -value	0.02	0.001	NS	NS	<0.001	0.05	—	—	—
	*	***	—	—	***	*	—	—	—

* $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$.

NS, not significant.

We found the same degree of relatedness in animals of different age, which suggests that related animals shoal together even with increasing age.

Discussion

Two major, genetically different populations of the Eurasian perch exist in Lake Constance: one in the eastern part of the lake including localities 1, 2, and 3 and a second one in the western part including localities 4, 5, and 6 (Fig. 2). Genetic differentiation was indicated by a statistically significant G_{ST} value of 0.07. These results correspond to those of Bodaly *et al.* (1989), who could distinguish between two distinct populations of perch in the northern and southern part of the English lake Windermere using allozyme analysis.

Within each of these two populations in Lake Constance, pairwise F_{ST} -values do not differ significantly. Because adult perch are supposed to be sedentary (Willemsen, 1977; Kipling & Le Cren, 1984), the degree of inferred genetic exchange within these populations might depend on distribution patterns of the larvae. According to this scenario, there might be separate areas in the eastern and western part in Lake Constance where larvae spend their pelagic phase before they return to the littoral zone. To which bank of the lake early juveniles return might depend on wind-driven transport of surface waters, but there is a clear separation between the two parts of the lake.

F_{ST} -values between localities 3 (RO) and 4 (KR) and between 4 (KR) and 5 (GN) and 6 (US) were not significantly different from 0. This result indicates genetic exchange between animals of these localities which can be explained by transportation, i.e. of larvae by the main water currents in Lake Constance. The Rhine river flows into the lake at the south-east, near locality 3 (RO) (Fig. 1). Near KR, where the Rhine flows out of the lake, there is a strong, continuous current towards US (Fig. 1).

In this study fish were collected during the winter. Therefore, we do not know yet whether individuals spend some time at different sites but return to their parental spawning sites during the mating period in early summer, resulting in an even stronger genetic substructuring among localities. This might be the case in walleye (*Stizostedion vitreum*), a fish quite closely related to perch with a similar ecology, where tagging studies indicated that animals moved readily between lakes during the year (Nepszy *et al.*, 1991); but significant differences in genotypic frequencies among spawning sites were consistent with the hypothesis of natal homing to spawning sites for both sexes during the mating season (Stepien & Faber, 1998). Experimentally supported natal homing was also found in walleye in

laboratory-reared released broodstock. This investigation found that natal homing in spawning walleye is governed by a genetically based response to environmental cues (Jennings *et al.*, 1996). More genetic data of perch from different localities should be collected during the spawning time. This would provide further evidence whether perch migrate during some periods of their life cycle but spawn at the same sites as their parents.

Perch from both Swiss lakes (Lake Zurich and Lake Walensee) which are interconnected by the Linth channel (Fig. 1) are genetically similar. The observation supports the assumption that animals use the channel to migrate between the lakes. Interestingly, perch from these lakes were genetically more similar to perch trapped in the eastern part of Lake Constance ($F_{ST}=0.054$, $P < 0.001$) than animals of the eastern and western parts of Lake Constance are to each other ($F_{ST}=0.07$, $P < 0.001$). This finding supports geological studies that there was a connection between the Swiss lakes and the eastern part of Lake Constance approximately 15 000 years ago (Kiefer, 1972). It also indicates that the genetic divergence between the two populations within Lake Constance is quite strong.

Our study reveals for the first time that perch were staying in shoals of half-siblings or more closely related individuals, as indicated by a high $Relatc = 0.674$ (Pamilo, 1984, 1989) and a high mean $r=0.115$ using the analysis of Queller & Goodnight (1989). The appearance of kin aggregations was also strongly supported using an extended method according to Blouin *et al.* (1996) revealing full- and half-sibs per locality. Blouin *et al.* (1996) demonstrated the ability of microsatellites to discriminate unrelated individuals from full-sibs with at least 97% accuracy. Attaining this level of accuracy necessitates using a large number of microsatellite loci. However, similar robustness was shown by Piortney *et al.* (1999) using a limited number of microsatellite loci for calculating pairwise relatedness.

The occurrence of shoals of full- and half-sibs can be explained by the mating system, but there have been few observations of mating under natural conditions; first attempts of filming spawning perch showed that multiple paternity may occur: two different males were observed fertilizing eggs of one female (Treasure, 1981).

Perch live in age-structured shoals (Wang & Eckmann, 1994; Imbrock *et al.*, 1996). The same level of relatedness was found in shoals of 0+ and those of 2+ years old fish, which suggests that related animals stayed together not only during the larval phase but continued to do so with increasing age. Several species of fish spend periods of their life cycles in kin aggregations. This result has been also observed in the Atlantic cod (*Gadus morhua*) (Herbinger *et al.*, 1997), coho salmon (*Oncorhynchus kisutch*) (Quinn & Busack, 1985), and in freshwater

species like cichlids (*Sarotherodon melanotheron*) (Pouyaud *et al.*, 1999), common shiners (*Notropis cornutus*) (Ferguson & Noakes, 1981), and three-spine sticklebacks *Gasterosteus aculeatus* (FitzGerald & Morrissette, 1992). Cichlids (*Sarotherodon melanotheron*) originating from riverine environments appeared to be panmictic, but in samples from open waters such as lagoons, showed sibling aggregations (Pouyaud *et al.*, 1999). Fontaine & Dodson (1999) did not reveal kin-biased distribution among juveniles of Atlantic salmon. But this study was performed in a large river which coincides with our findings that animals are less related in areas with high water currents (localities 3 and 4). These highly dispersive environmental conditions are less conducive to the establishment of kin-biased distribution.

Shoaling behaviour is proposed to be an altruistic trait that may have evolved through kin selection (Pitcher, 1986). Individuals might increase their inclusive fitness by providing the benefits of cooperative behaviour to related individuals (Hamilton, 1964). The mechanisms by which these kin groups develop and stay distinct from other such groups is unknown. A kin recognition system might permit larvae to distinguish between kin and non-kin which hatch in close proximity to them. They might use olfactory cues from the eggs, as has been shown in tadpoles, which learn to distinguish between kin and non-kin during their early life phase (Blaustein & Waldman, 1992). Alternatively, female perch might mark the substrate where the eggs are laid, as is assumed for Atlantic salmon (*Salmo salar*) which occupy fixed territories on the stream bed during the summer months (Kalleberg, 1958; Keenleyside & Yamamoto, 1962). The substrate within each territory is probably marked with odours that can be recognized by kin (Stabell, 1987). Active compounds produced in the liver and expelled via the intestinal tracts could mediate this recognition (Quinn & Hara, 1986; Stabell, 1987). Laboratory experiments on recognition of kin vs. non-kin groups of Atlantic Salmon (*Salmo salar*) (Brown & Brown, 1993, 1996) reveal the possible advantage of kin aggregations. Kin groups had fewer aggressive interactions, used a greater proportion of 'threat' behaviour as opposed to fighting, and had improved growth, especially in subordinates.

Kin recognition mechanisms play an important role in assortative mating (Hepper, 1986). By avoiding breeding with close relatives but choosing genetically similar mating partners adaptations to specific environmental conditions are maintained which might be the basis for processes of speciation (Kondrashov, 1998; van Oppen *et al.*, 1998). Additional behavioural studies of perch might be able to identify the behavioural mechanisms of these processes.

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