A microsatellite study in the Łęgucki Młyn/Popielno hybrid zone reveals no genetic differentiation between two chromosome races of the common shrew (Sorex araneus)

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Received: 13 May 2010 / Accepted: 8 November 2010 / Published online: 22 February 2011 © The Author(s) 2011. This article is published with open access at Springerlink.com

Abstract This study investigated a chromosome hybrid zone between two chromosomal races of the common shrew (Sorex araneus). Gene flow and genetic structure of the hybrid zone, located in the northeast of Poland, were studied using seven polymorphic autosomal microsatellite loci (L9, L14, L33, L45, L67, L68, L97) and a Y-linked microsatellite locus (L8Y). Seventy-five animals (46 of the Łęgucki Młyn race and 29 of the Popielno race) from nine different localities were examined and the data were analyzed using hierarchical AMOVA and F-statistic. The studied microsatellite loci and races (divided into nine geographical populations) were characterized by observed heterozygosity ($H_O$), expected heterozygosities within ($H_S$), and between ($H_T$) populations, inbreeding coefficient ($F_{IS}$), fixation index ($F_{ST}$), and average allelic richness ($A$). We found that genetic structuring within and between the two chromosome races were weak and non-significant. This finding and unconstrained gene flow between the races indicates a high level of migration within the Łęgucki Młyn/Popielno hybrid zone, suggesting that evolutionarily important genetic structuring does not occur in interracial zones where races which are not genetically distinct come into contact.

Keywords DNA polymorphism · Gene flow · Genetic structure · Microsatellites · Sorex araneus

Introduction

The common shrew (Sorex araneus), a small insectivore, seems to be an ideal object for the study of evolutionary processes, especially to investigate the role of chromosomes in speciation. It is a unique species, first of all, due to its great chromosome variation (both within and between populations) (Searle and Wójcik 1998). The basis of the quantitative and qualitative variation of the common shrew’s autosomes is chromosome aberrations in which the main cause is believed to be centric fusions (Robertsonian fusions) (Searle and Wójcik 1998). However, the recent study by White et al. (2010) taking into account different types of mutation (fusions, fissions, whole-arm reciprocal translocations, and zonal raciation events) revealed that fusions may be much less important than previously imagined.

As an effect of karyotype variation, S. araneus has been divided into almost 70 chromosome races, ten of which are found in Poland (Wójcik et al. 2003; Fedyk et al. 2008). In many cases, the neighboring races created hybrid zones, and several of them have already been described (Searle and Wójcik 1998; Jadwiszczak 2002;
In hybrid zones, detailed analysis has been done of either modifications enabling gene flow between neighboring races (acrocentric peak, recombinant peak, or elimination of acrocentrics from the zone) or factors responsible for the formation of isolation barriers between races (Searle and Wójcik 1998; Fredga and Narain 2000; Moska 2003).

Different molecular markers have been used in studies of the evolutionary processes of *S. araneus* (allozymes, mtDNA, microsatellites), of which microsatellites have been found a sensitive measure to detect patterns of hybridization. Up to now, the presence of over 40 highly polymorphic autosomal microsatellite loci and one polymorphic Y-chromosome microsatellite (L8Y) of this species have been reported (Wyttenbach et al. 1999; Balloux et al. 2000; Lugon-Moulin et al. 2000; Brügger et al. 2002a, b; Lugon-Moulin and Hausser 2002; Andersson 2004; Basset et al. 2006). L8Y provides an opportunity for investigating the level of male gene flow between populations and comparing it with the gene flow of both sexes (Lugon-Moulin and Hausser 2002).

The aim of the present study was to analyze the level of genetic differentiation in the Łęgucki Młyn/Popielno hybrid zone using microsatellite markers. Moreover, by analyzing the genetic structure within and between chromosome races, we tried to contribute to efforts of elucidating the role of interracial hybrid zones in the evolutionary process of speciation.

### Material and methods

#### Study area

The hybrid zone between the Łęgucki Młyn and Popielno karyotypic races is located in the northeast of Poland (Fig. 1). The Łęgucki Młyn race (Lg) is characterized by chromosomes: \( g/r, h/k, i/o, j/l, m/n, p, q \) while the Popielno race (Po) by: \( g/r, h/q, i/k, j/l, m/n, o, p \) (Wójcik 1986; Moska 2003).

Shrews were collected over the period 1996–1999 from nine sites: Prawdowo (P), Lisunie-Kulinowo (LK), Lipowo (L), Nowy Most (NM), Krutyń I (KI), Krutyń II (KII), Zielony Lasek (ZL), Zakręt (Z), and Karwica Mazurska (KM) (Fig. 1). Sites P, LK, L, NM, and ZL were situated in the area of the Łęgucki Młyn race (southern side of the hybrid zone) and KI, KII, Z, and KM in the area of the Popielno race (northern side of the hybrid zone) (Moska 2003).

#### Sampling and molecular methods

For the microsatellite analysis, 75 individuals of two races were used, 46 of the Łęgucki Młyn race and 29 of the Popielno race. Seven polymorphic autosomal microsatellites (L9, L14, L33, L45, L67, L68, L97) and a Y-linked microsatellite (L8Y) were used for the analysis. Amplification conditions for each locus were carried out according to Balloux et al. (1998) and Lugon-Moulin et al. (2000). In order to improve estimation of the size of the amplification
product, one primer of each pair was labeled with a fluorescent dye (FAM or JOE) on the 5′-end, which allowed analyses with the use of an ABI 3100 Avant automated sequencer (Applied Biosystems).

Statistical analysis

The data were statistically analyzed with the application of two different approaches to the genetic structure of the Łęgucki Młyn/Popielno hybrid zone:

1. First, the samples from the nine sites were considered as nine geographical populations;
2. Second, the nine geographical populations were grouped according to their location in the area occupied. Populations P, LK, L, NM, and ZL situated in the area of the Łęgucki Młyn race formed group I; populations KI, KII, Z, and KM from the area of the Popielno race formed group II.

For the geographical populations, allelic richness (A), the observed (H_0), and expected (H_E) heterozygosity were computed for all loci using SAS v. 8.0 (User’s Guide 2000) and ARLEQUIN v. 3.1 (Excoffier et al. 2005).

Inbreeding coefficients (F_IS) were estimated to indicate the within-population heterozygote deficiency due to non-random mating. F_IS was calculated per locus, overall loci and per population for the geographical populations. Fixation index (F_ST) was estimated per locus and for overall microsatellite loci. Permutations were used for testing the values of F_IS and F_ST for significant departure from zero. These tests were done using 10.000 permutations of alleles within samples (F_IS) and 10.000 permutations of genotypes among samples for F_ST (Arlequin v. 3.1).

To estimate the gene flow between the races, conventional F-statistics were used (Weir and Cockerham 1984). These statistics are recommended rather than R-statistics for studies with small sample sizes (ten individuals or less) and limited numbers of loci (less than 20) (Gaggiotti et al. 1999).

The genetic structure of populations was calculated using the analysis of molecular variance (AMOVA) as implemented in Arlequin v. 3.1. A hierarchical analysis of variance divided the total variance into different parts of covariance which were then used to calculate different fixation indices (Excoffier 2000). Significance levels of the different fixation indices were obtained by permuting genotypes for F_ST, among geographical populations; for F_SC, among geographical populations within the races; and for F_CT, among chromosomal races (Excoffier et al. 1992).

Results

Polymorphism and genetic variability

Most autosomal loci displayed a high level of polymorphism. One hundred and thirty six alleles were found at seven autosomal microsatellite loci. The most variable locus (L97) had 30 alleles, whereas the least variable one (L67) had 12 alleles (Table 1).

Table 1 Number of alleles (n_a), observed heterozygosity (H_0), expected heterozygosities within (H_S) and between (H_T) populations, inbreeding coefficient (F_IS), and fixation index (F_ST) estimated for geographical populations and both races

| Locus | n_a | H_0 | H_S | H_T | F_IS | P | Geographic populations | Race group |
|-------|-----|-----|-----|-----|------|---|------------------------|------------|
| L9    | 27  | 0.86| 0.83| 0.94| 0.08 | 0.0200 | Lg                     | 0.010      |
| L14   | 14  | 0.82| 0.79| 0.89| 0.07 | 0.0986 | 0.023                  |
| L33   | 26  | 0.68| 0.80| 0.93| 0.27*| 0.0000 | 0.020                  |
| L45   | 13  | 0.57| 0.69| 0.78| 0.28*| 0.0000 | 0.033                  |
| L67   | 12  | 0.68| 0.68| 0.73| 0.08 | 0.0105 | 0.011                  |
| L68   | 14  | 0.85| 0.79| 0.87| 0.00 | 0.5489 | 0.008                  |
| L97   | 30  | 0.79| 0.75| 0.94| 0.15*| 0.0009 | 0.038*                 |
| All loci | 136 | 0.75| 0.76| 0.87| 0.13*| 0.000  | 0.020                  |

*P<0.01

Micro-Checker program v. 2.2.3 (Van Oosterhout et al. 2004) was used to identify genotyping errors due to null alleles.

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Table 2 Number of individuals (n) used in the analysis of microsatellites (number of males is given in brackets), average allelic richness (A), observed (H0) and expected (H0) heterozygosity and inbreeding coefficient (Fis) for each of the geographical populations

| Locality | Number (n) | A    | H0   | H0   | FIS  | P    |
|----------|------------|------|------|------|------|------|
| P        | 10 (5)     | 7.11 | 0.76 | 0.86 | 0.11 | 0.0681 |
| LK       | 15 (8)     | 8.55 | 0.69 | 0.80 | 0.14*| 0.0061 |
| L        | 3 (2)      | 3.33 | 0.65 | 0.75 | 0.10*| 0.0000 |
| NM       | 7 (3)      | 6.55 | 0.76 | 0.88 | 0.02 | 0.4851 |
| KI       | 10         | 4.12 | 0.86 | 0.88 | −0.12| 1.0000 |
| KII      | 4 (3)      | 8.33 | 0.79 | 0.88 | 0.00 | 0.5903 |
| ZL       | 11 (3)     | 8.11 | 0.81 | 0.89 | 0.04 | 0.2649 |
| Z        | 13 (8)     | 10.11| 0.78 | 0.87 | 0.03 | 0.1939 |
| KM       | 2          | 2.75 | 0.64 | 0.81 | 0.29 | 0.3461 |
| Total    | 75 (32)    |      |      |      | 0.06*| 0.0027 |

*P<0.01

from 0.57 to 0.86, and over all loci amounted to 0.75. Expected heterozygositys within samples (H0) ranged from 0.68 to 0.83 with an average value of 0.76, whereas over all samples (HT) ranged from 0.73 to 0.94 with the average value of 0.87 (Table 1). The average H0 in different geographical populations ranged from 0.64 to 0.86, whereas H0 fluctuated from 0.75 to 0.89 (Table 2).

Genetic structure of the Łęgucki Mlyn/Popielno hybrid zone

FIS estimated over all loci revealed significant departure from HWE expectations (FIS=0.063, P<0.01). The FIS observed in each geographical population ranged from 0.12 (Kl) to 0.29 (KM) and departed significantly from HWE in two of them: L/K and L (P<0.01; Table 2). All other geographical populations showed no FIS values significantly different from zero. In the analysis of mean FIS over all loci, a very high and significant heterozygote deficit (FIS=0.13) was observed. FIS per locus over all populations ranged from 0.00 (L68) to 0.28 (L45). Results from the permutation procedure indicated that the FIS values were significant for three loci: L33, L45, and L97 (Table 1).

The FST over all geographical populations was small and not significant (FST=0.02; Table 1). Locus-specific FST ranged from 0.008 (L68) to 0.038 (L97), and significant genetic differentiation was found only at locus L97 (Table 1). The mean FST was also calculated for both races separately. The Łęgucki Mlyn race (populations P, L/K, L, NM, ZL) showed FST of 0.027, whereas the Popielno race showed FST of 0.003. For both races values of FST were small and non-significant (Table 1).

The AMOVA analyses based on FST showed a lack of genetic differentiation between chromosomal races, both for autosomal microsatellite loci and for the Y-locus. For the autosomal loci, 0% (−0.26%) of the total variation was caused by variation between races, whereas for L8Y it was 3.03% of the total variation. Most of the variation was found within geographical populations as regards to both autosomal loci and L8Y (99% and 90%, respectively). For all genetic markers low and non-significant variation was found among geographical populations within the races.

Discussion

So far, detailed studies using DNA microsatellites have been carried out in three hybrid zones: two located in Sweden (Uppsala/Hällefors and Abisko/Sidensjö) (Wyttenbach et al. 1999; Andersson 2004), and one located in Poland (Đrnholce/Białowieża) (Jadwiszczak et al. 2006). The Abisko/Sidensjö and Đrnholce/Białowieża hybrid zones are the ones where hybridization of karyotypically divergent chromosome races takes place (races belong to distinct karyotypic types: NEKG and WEKG, WEKG and EEKG, respectively) (Fredga 1996; Jadwiszczak 2002). The analysis of genetic differentiation revealed weak genetic structuring and in consequence unimpeded gene flow between the races. However, in the Abisko/Sidensjö hybrid zone, significant reduction of male gene flow was found when the genetic marker linked to Y-chromosome (L8Y) was studied (Andersson 2004; Jadwiszczak et al. 2006). In the Uppsala/Hällefors hybrid zone, both races are not chromosomally distinct (they belong to the same evolutionary group, WEKG) (Narain and Fredga 1996). Based on genetic differentiation of autosomal microsatellite loci, weak genetic structuring over all localities was found (Wytenbach et al. 1999).

A comparison of the studied hybrid zones leads to the conclusion that the Łęgucki Mlyn/Popielno hybrid zone strongly resembles the Uppsala/Hällefors hybrid zone. These two races (Lg and Po) are chromosomally quite closely related (both races represent EEKG) and in the hybrid zone complex heterozygotes are theoretically expected to form only short chains (CIV or CV) during meiosis.

Molecular analysis of seven autosomal loci did not reveal significant differentiation between the Łęgucki Mlyn and Popielno chromosomal races. The FST values were small and not significant, either over all geographical populations or within the races. Our results were comparable with the weak genetic structuring observed in the similar Uppsala/Hällefors hybrid zone.

When we analyzed the level of genetic structure with regard to the Y-chromosome microsatellite, we found a level ten times higher compared to autosomal loci (FST=0.10). Our findings are comparable to results previously reported by different authors who observed differences in the estimates for autosomal microsatellite loci and for the Y-locus. For
of genetic structure based on the Y-microsatellite variation compared to the autosomal microsatellite (Andersson et al. 2004; Balloux et al. 2000).

In our study, \( F_{IS} \) values over all loci and over all populations were high and significantly different from zero (\( F_{IS}=0.13 \) and \( F_{IS}=0.06 \), respectively). A closer look at \( F_{IS} \) for each geographical population revealed that populations LK and L showed a high and significant heterozygote deficit (\( F_{IS}=0.14 \) and \( F_{IS}=0.10 \), respectively; \( P<0.01 \)). The high \( F_{IS} \) may be partly caused by the Wahlund effect (White and Searle 2007). In one geographical population (LK), shrews were caught on two meadows separated by a busy road, so a deficit of heterozygotes could be caused by pooling independent populations in HWE. Previous studies have shown no indications of null alleles at any of the loci used in this study, indicating that high \( F_{IS} \) was probably not caused by presence of null alleles (Wytenbach and Hauser 1996; Wytenbach et al. 1999; White and Searle 2007).

However, the analysis of the data used in our study with the use of Micro-Checker software identified null alleles in three loci (L33, L43, and L97) where a significant (\( P<0.01 \)) inbreeding coefficient was estimated. This is probably the most reasonable explanation of the lack of heterozygosity in those loci. The deficit of heterozygotes observed in our study may be also explained by the small sample sizes (particularly from populations L, KII, and KM, where samples of fewer than five individuals were taken).

In conclusion, the results described in this paper give insight into the speciation process which may take place in hybrid zones. Our study, with the use of microsatellite markers, did not reveal significant differences between the genetically close Łęgucki Mlyn and Popielno chromosomal races, which was proven by small chromosomal differentiation. It seems that the evolutionarily important genetic structuring does not occur in interracial zones, where races which are not genetically distinct come into contact.

Acknowledgements The field assistance of Professor S. Fedyk and Dr. W. Chętnicki is gratefully acknowledged. We wish to thank M. Kowalska and A. Jonkisz for their laboratory help. Thanks also go to anonymous referees for their useful comments.

Declaration concerning ethical standards Our experiments comply with all current laws in Poland.

Conflict of interest The authors declare that they have no conflict of interest.

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