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Published in:
Molecular Ecology

DOI:
10.1046/j.1365-294X.2003.02005.x

2003

Citation for published version (APA):
SHORT COMMUNICATION

MHC diversity in two Acrocephalus species: the outbred Great reed warbler and the inbred Seychelles warbler

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Abstract

The Great reed warbler (GRW) and the Seychelles warbler (SW) are congeners with markedly different demographic histories. The GRW is a normal outbred bird species while the SW population remains isolated and inbred after undergoing a severe population bottleneck. We examined variation at Major Histocompatibility Complex (MHC) class I exon 3 using restriction fragment length polymorphism, denaturing gradient gel electrophoresis and DNA sequencing. Although genetic variation was higher in the GRW, considerable variation has been maintained in the SW. The ten exon 3 sequences found in the SW were as diverged from each other as were a random sub-sample of the 67 sequences from the GRW. There was evidence for balancing selection in both species, and the phylogenetic analysis showing that the exon 3 sequences did not separate according to species, was consistent with transspecies evolution of the MHC.

Keywords: Balancing selection, Genetic variation; MHC; Passerines; Population bottleneck; Transpecies evolution

Received 25 July 2003; revision received 9 September 2003; accepted 9 September 2003

Introduction

The major histocompatibility complex (MHC) is an important component of the vertebrate immune system where it determines which antigens trigger an immune response (Hughes & Yeager 1998). Extraordinary levels of genetic variation occur within the MHC, and how this polymorphism is maintained is still debated, but the selection pressure exerted by pathogens is considered a major factor (Jeffery & Bangham 2000; Hess & Edwards 2002). The MHC is thought to play a role in aspects such as disease resistance, kin recognition, inbreeding avoidance and mate choice (Grob et al. 1998; Penn & Potts 1999).

Studies have characterized and investigated the ecological consequences of MHC variation in mammal and fish species (e.g. Paterson et al. 1998; Langefors et al. 2000; Ditchkoff et al. 2001; Reusch et al. 2001). But in birds, used extensively in studies of natural and sexual selection, little work has focused on the role of the MHC so far (von Schantz et al. 1996, 1997). Recent developments in molecular methods have facilitated the study of the avian MHC and various studies on birds are now underway (Zelano & Edwards 2002). However, before attempting to investigate the ecological consequences of MHC variation in a wild population, it is important to assess MHC variation to determine the sense and feasibility of such a study.

The Great reed warbler (GRW, Acrocephalus arundinaceus) and the Seychelles warbler (SW, Acrocephalus sechellensis), two of the worlds most intensively studied wild bird species, are congeners (Leisler et al. 1997) with markedly different demographic histories. Founded in 1978, the population of GRWs at Lake Kvismaren now consists of approximately 60 breeding individuals (Bensch & Hasselquist 1991; Hasselquist et al. 1995) and, with 12–25 immigrants each year, can be considered part of the normal distribution of this outbred migrant species (Hansson et al. 2000, 2003). In contrast, the SW, a rare endemic of the Seychelles islands that was pushed to the verge of extinction, is an inbred, isolated species. Between 1959 and 68, this species went through a severe genetic bottleneck with only 26–29 birds remaining on the island of Cousin (Crook 1960). The population has since recovered and now remains stable at a carrying capacity of 320–350 birds (Komdeur 1992; Richardson et al. 2002).

In the SW the low levels of variation seen at neutral microsatellite loci (Richardson et al. 2000) are probably the result of the bottleneck this species has been through.
However, balancing selection is thought to be able to maintain variation in MHC genes even in very restricted populations (Hughes & Yeager 1998). Here we measure the MHC class I exon 3 variation in the inbred SW compared to the GRW. We assess both the number of, and genetic variability within, MHC class I alleles and determine if balancing selection has been able to maintain MHC variation in the SW. Finally, we construct a phylogenetic tree to investigate the relationship among MHC alleles between the GRW and the SW.

Materials and methods

Study populations and samples

The SW is a small, sedentary, passerine endemic to the Seychelles Islands. The isolated Cousin Island population has been studied intensively since 1985 (Komdeur 1992; Komdeur et al. 1997; Richardson et al. 2003). In contrast, the GRW is a medium sized passerine bird which breeds in reed beds in the central and northern Palaearctic and migrates to overwinter in sub-Saharan Africa (Cramp 1992). In 1978 the first GRWs were breeding at the study site, 15 km east of Örebro (59°10′ N, 15°25′ E) in Sweden, and since 1983 the population has been studied intensively (e.g. Bensch & Hasselquist 1991; Hasselquist et al. 1995). During the years of study, nearly all birds within both populations have been individually colour-ringed, monitored, and blood sampled (Hasselquist et al. 1995; Richardson et al. 2001). The present study includes samples from the 485 SWs (c. 96%), present on Cousin Island between 2000 and 2002, and the 354 GRW’s (> 96%), breeding in Lake Kvismaren between 1985 and 1996.

Restriction fragment length polymorphism (RFLP) analysis

Total genomic DNA was extracted using either a phenol extraction technique (following Bruford et al. 1998), or a salt extraction method (Richardson et al. 2001). Ten µg of DNA was digested with the restriction enzyme PvuII (Boehringer Mannheim), separated on an agarose gel, transferred to a nylon membrane and hybridized overnight with the probe 21P labelled with [α-32P]dCTP (Westerdahl et al. 1999). The membranes were then washed and exposed to an X-ray film. For the SW, RFLP analysis followed the methods outlined in Westerdahl et al. (1999) and the GRW results are from Westerdahl et al. (1999).

Motif specific polymerase chain reaction (PCR) and denaturing gradient gel electrophoresis (DGGE)

The protocol for motif specific amplification using PCR, followed by the separation of exon 3 sequence using DGGE developed by (Westerdahl et al. 2003) was used to screen efficiently MHC class I exon 3 sequence variation. Exon 3 encodes the peptide-binding region (PBR) of the MHC molecule; hence the region that is critical for the binding of peptides and for generating specific immune responses when the peptide bound is nonself (Hughes & Yeager 1998).

In the GRW, we have evidence of at least four transcribed MHC class I (cDNA) genes (Westerdahl et al. 1999) and an additional four genomic MHC class I genes (Westerdahl et al. 2003). Our sequence specific amplification detects a limited number of alleles from several class I loci, and gives a broad estimate of the MHC class I exon 3 variation within each individual. It is a repeatable method that is more sensitive in detecting genetic variation than the RFLP method and enables us to avoid screening pseudogenes.

Two primer combinations (HN36-GC46 and HN38-GC46) were used in motif specific PCRs; one motif specific primer within exon 3 and one general primer in intron 3 (Westerdahl et al. 2003). The two primer sets amplify different sets of sequences since the primers HN36 and HN38 have different motifs in the 3’-end (AGT and ACG, respectively). Each primer combination amplified 260 bp from the variable exon 3 (the complete exon 3 is 274 bp). Sequences were separated using gels containing 7% 19:1 acrylamide/bisacrylamide, 1 × TAE buffer and a denaturing gradient of urea and formamide (Myers et al. 1987). Gels were run at 60 °C in 1 × TAE buffer (C.B.C. Scientific Company, Inc.) with PCR products from both primer combinations run together in 40–70% (or 40–65% for the GRW primer combination HN36-GC46) denaturant at 190 V for 16.5 h. Standardized markers were used to enable comparisons between gels. Gels were stained using SYBR gold (Molecular Probes), and the DNA was visualized in a FluorImage SI (Molecular Dynamics Inc.).

Sequencing

Ten different DGGE bands occurred in the SW, while 67 bands were detected in the GRW population. Ten, randomly chosen, MHC class I exon 3 sequences amplified with the DGGE primers, were selected in the GRW’s to constitute an equal subsample of DGGE bands for between species comparisons. DGGE bands were excised from gels and dissolved in 150 µL of ddH2O. This solution was frozen (−80 °C) and melted (4 °C) repeatedly, diluted 1:50 and then reamplified with the original primers. The PCR product was purified and directly sequenced on an ABI PRISM 310 Genetic Analyser (Perkin Elmer). Each band was sequenced in 2–5 unrelated individuals. There was good evidence that none of the sequences were from pseudo-genes; there were no deletions in the sequences, particular features for functional MHC genes were found,
e.g. conserved cysteine codons, and several motives were identical to chicken, *Gallus domesticus*, class I genes.

When running DGGE’s based on a PCR where several alleles have been amplified, heteroduplexes may be formed. In this study, all the DGGE bands used gave clear, unambiguous sequences and were not the result of heteroduplexes.

**Phylogenetic analysis**

All 10 SW warbler exon 3 sequences, and a total of 32 GRW exon 3 sequences, including cDNA-sequences (Westerdahl *et al.* 1999), genomic sequences and sequences likely to be from pseudogenes (Westerdahl *et al.* Submitted) were used in the phylogenetic tree. The tree was constructed in **MEGA** 2.1 software (Nei & Gojobori 1986) using the minimum evolutionary method and the entire exon 3 sequences (273 bp). Bootstrap tests verified the probability of the branches. All sequences have been submitted to GenBank (accession numbers; Acar-UA, AY306008–AY306009; Ase-UA, AJ557874–AJ557883).

**Statistical analysis**

The evolutionary distance (*d*) between exon 3 sequences was computed using the **kIMURA** 2-parameter model, while the number of synonymous and nonsynonymous substitutions per site in the PBR of exon 3 was calculated by Nei and Gojobori’s method of pair-wise comparisons, using **MEGA** version 2.1 (Nei & Gojobori 1986). The PBR was superimposed from the human sequence (Bjorkman *et al.* 1987). All tests are two-tailed and means are given ± one standard deviation.

**Results and Discussion**

The level of variation at the MHC class I exon 3 loci is high in the GRW but low in the SW. Both species have roughly an equal number of RFLP fragments (21–25 GRW vs. 23–25 SW), which suggests they have an equal number of MHC class I genes. However, 89% (49/55) of GRW’s had unique class I RFLP genotypes — similar to the 88% (42/48) of GRW’s that are closely related. There was no significant difference between the SW and the GRW in the number of nucleotide differences, overall amino acid variation per site in the PBR than in the non-PBR (Table 1).

The patterns of genetic variation found in the GRW and the SW are consistent with their demographic history. The GRW population is relatively outbred, with individuals immigrating in from the pan-European population (Hansson *et al.* 2000; Hansson *et al.* 2003), while the SW has recently been through a bottleneck and remains totally isolated (Komdeur 1994). The number of MHC alleles in the SW does, however, appear to be high considering the recent population bottleneck, especially when compared to the extremely low levels of variation found with neutral markers (Richardson *et al.* 2000). For example, significantly more microsatellites characterized in the SW were monomorphic (32/63 = 51%, Richardson *et al.* 2000) than were microsatellites characterized in the GRW (1/11 = 9%, Hansson *et al.* 2000) (Fisher exact *P* = 0.018).

Balancing selection appears to play a determinant role in MHC evolution (Bernatchez & Landry 2003) and one indication of this is a higher number of nonsynonymous 

\[ d_n \] than synonymous 

\[ d_s \] substitutions in the PBR. In the present study the ratio of 

\[ d_n \] to 

\[ d_s \] tended to be greater than one in the PBR, but less than one in the non-PBR, for both the SW [PBR; 

\[ d_n/d_s = (0.34 \pm 0.109)/(0.20 \pm 0.11) = 1.65 \]

Non-PBR 

\[ d_n/d_s = (0.05 \pm 0.01)/(0.11 \pm 0.03) = 0.43 \] and the GRW [PBR; 

\[ d_n/d_s = (0.38 \pm 0.12)/(0.33 \pm 0.17) = 1.17 \]

Non-PBR 

\[ d_n/d_s = (0.05 \pm 0.01)/(0.09 \pm 0.03) = 0.52 \]. Although these differences were not significant (SW, *t*-test = 0.28, *P* > 0.05; GRW, *t*-test = 0.28, *P* > 0.05), this may be because we are comparing exon 3 sequences across loci, in which case the numbers of synonymous substitutions are likely to be higher than when comparing alleles within a locus (Hughes & Nei 1989; Westerdahl *et al.* 1999). Evidence for balancing selection also comes from the fact that both species have significantly higher than average amino acid variation per site in the PBR than in the non-PBR (Table 1).

Theory suggests that heterozygous individuals with diverged MHC alleles will be at an advantage, since they will be able to respond to a wider range of pathogens compared with homozygous individuals, or even heterozygous individuals with two similar alleles (Hughes & Yeager 1998). Within a population, a high level of divergence between MHC alleles would provide further evidence that selection is acting to maintain MHC variation. In both the SW and the GRW populations, a high level of divergence was apparent within exon 3 sequences. The average number of nucleotide differences between any two exon 3 sequences was high (Table 1), and only a few exon 3 sequences were closely related. There was no significant difference between the SW and the GRW in the number of nucleotide differences, overall amino acid variation, or average amino acid variation per site in the PBR or in the non-PBR (Table 1). Furthermore, within species variation was not significantly different from between species variation for each measure (Table 1). Interestingly,
although fewer DGGE-alleles (exon 3 sequences) were observed in the SW (10 vs. 67), there was an equally high level of divergence between exon 3 sequences in the SW as there was in the GRW population. This indicates that individuals with diverged MHC alleles have been selected for during and/or after the severe bottleneck the SW population went through. Such a pattern of a low number of MHC alleles, with a high degree of divergence between alleles, has been found in studies of other bottlenecked species (e.g. Mikko et al. 1997; Hedrick et al. 1999; Hoelzel et al. 1999; Hedrick et al. 2000).

In the present study, we have investigated divergence between sequences from several different (unspecified) loci. This could lead to higher estimations of divergence than in intra locus comparisons, since synonymous substitutions are likely to accumulate between loci over time. However, the GRW and SW seem to have an equal number of class I loci and therefore the between species divergence may be similar.
The MHC system in passerine birds is more complex than that observed in chickens (Kaufman et al. 1999; Westerdahl et al. 2000; Freeman-Gallant et al. 2002). This, combined with the extensive MHC class I polymorphism seen in the GRW, has made it difficult to identify the number of class I genes present, and thus to characterize the MHC in detail (Westerdahl et al. 1999). The SW may be an excellent model organism in which to do this, as its limited genetic variation means that the loci involved are more likely to be homozygous, thereby making characterization simpler. The high levels of polymorphism in the GRW have also complicated attempts to investigate the ecological consequences of MHC characteristics in a wild avian population. The SW provides a simplified system in which to investigate such questions: the population is isolated and, while it contains limited genetic variation at the MHC (thus making statistical analysis more tractable), this appears to be maintained by selection. The SW has also been the focus of extensive study and many important factors required for an in-depth investigation are already available or known. On the other hand, the GRW population is more likely to be representative of a normal passerine system than is the SW. Ultimately, using both species in complimentary and comparative studies should provide the most productive research strategy.

Acknowledgements

Nature Seychelles kindly allowed David S. Richardson (DSR) to work on Cousin Island. The Department of Environment and the Seychelles Bureau of Standards gave permission for fieldwork and sampling. We thank K. Pernson for help in the laboratory, and T. von Schantz (TvS), J.F. Dallas and two anonymous reviewers for comments on drafts of this MS. This research has been supported by a Marie Curie Fellowship to DSR (HPMF-CT-2000–01074), and from The Swedish Research Council to TvS.

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