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Limited diversity associated with duplicated class II MHC-\textit{DRB} genes in the red squirrel population in the United Kingdom compared with continental Europe

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Abstract

The red squirrel (*Sciurus vulgaris*) population in the United Kingdom has declined over the last century and is now on the UK endangered species list. This is the result of competition from the eastern grey squirrel (*S. carolinensis*) which was introduced in the 19th century. However, recent evidence suggests that the rate of population decline is enhanced by squirrelpox disease, caused by a viral infection carried asymptomatically by grey squirrels but to which red squirrels are highly susceptible. Population genetic diversity provides some resilience to rapidly evolving or exotic pathogens. There is currently no data on genetic diversity of extant UK squirrel populations with respect to genes involved in disease resistance. Diversity is highest at loci involved in the immune response including genes clustered within the major histocompatibility complex (MHC). Using the class II *DRB* locus as a marker for diversity across the MHC region we genotyped 110 red squirrels from locations in the UK and continental Europe. Twenty four *Scvu-DRB* alleles at two functional loci; *Scvu-DRB1* and *Scvu-DRB2*, were identified. High levels of diversity were identified at both loci in the continental populations. In contrast, no diversity was observed at the *Scvu-DRB2* locus in the mainland UK population while a high level of homozygosity was observed at the *Scvu-DRB1* locus. The red squirrel population in the UK appears to lack the extensive MHC diversity associated with continental populations, a feature which may have contributed to their rapid decline.

**Keywords:** Red squirrel, MHC *DRB*, Population, UK, diversity, Squirrelpox virus, disease
Introduction

The Eurasian red squirrel (*Sciurus vulgaris*) is currently on the endangered species list in the United Kingdom (UK) although not in the rest of its pan-Eurasian range. Within the UK the majority of the population is restricted to Scotland with fragmented populations remaining in England and Wales, while the distribution of the eastern grey squirrel (*S. carolinensis*) has expanded to match that vacated by the red squirrels. As recently described in detail by Signorile et al. (2016), the North American eastern grey squirrel was introduced and subsequently translocated across the UK and Ireland on at least 30 occasions from the 1870’s until the 1920’s (Middleton 1930; Shorten, 1954, Barrett et al. 1999). Grey squirrel numbers have increased ever since and have been estimated at around 2.5 million while red squirrel numbers have declined to approximately 120,000 (Harris et al. 1995). In continental Europe the grey squirrel has also been introduced to Northern Italy on at least three occasions between 1948 and the 1990s, followed by numerous translocations and undocumented releases (Martinoli et al. 2010; Bertolino et al. 2008, 2014). However, no evidence of the SQPV has been reported which may partially explain the slower rate of decline in Northern Italian red squirrels compared with those in the UK.

The principal factors that underlie the rapid decline of the red squirrel and replacement by grey squirrels in the UK include competition from the grey squirrel (Gurnell et al. 2004; Kenward and Holm 1993; Tompkins et al. 2002; Wauters and Gurnell 1999) and disease caused by infection with the squirrelpox virus (SQPV) (Thomas et al. 2003; La-Rose et al. 2010). SQPV, a member of the Poxviridae family (Thomas et al. 2003; McInnes et al. 2006; Darby et al. 2014) is thought to be transmitted by asymptomatic grey squirrels (Sainsbury et al. 2000; Tompkins et al. 2002) to highly susceptible red squirrels. It has been estimated that on average 61% of grey squirrels in the UK are seropositive for SQPV (McInnes et al. 2006), although this fluctuates between 100% and 0% depending on the density of squirrels supported by different types of woodland.

Infection of red squirrels with SQPV generally results in death within 2-3 weeks of infection which is likely to be a result of starvation and dehydration due to the inability to forage for food and water and the combined effect of secondary, mainly bacterial, adventitious infections. In areas where red and grey squirrels coexist the decline of red squirrels is up to twenty five times faster if the grey squirrels are carrying SQPV than if they are free from the virus (Rushton et al. 2006). As a consequence, the red squirrel is unlikely to survive in the UK unless populations are maintained in favourable conifer habitats that reduce competition and immigration by grey squirrels (Gurnell et al. 2002).
In response to the threat posed by SQPV, a number of red squirrel strongholds have been established in the UK which combine measures to control exposure to the grey squirrels with habitat improvement. However, small isolated populations often suffer from reductions in genetic diversity due to inbreeding depression and the effect of genetic drift (Keller and Waller 2002; Charlesworth and Willis 2009). This reduces the ability of such populations to respond to rapidly evolving endemic and exotic pathogens compared with larger more genetically diverse populations (Frankham and Ralls 1998; Bernatchez and Landry 2003). Maintaining existing red squirrel diversity while developing strategies that allow diversity to increase within isolated populations will be important for the long term sustainability of the red squirrel strongholds. Historical evidence indicates that red squirrels may have experienced severe population declines and bottlenecks and there is a complete lack of knowledge on genetic diversity of extant UK populations especially with respect of genes involved in disease resistance. Previous analyses of genetic diversity in the red squirrel have targeted nuclear, neutral microsatellite and mitochondrial markers providing important information on the population structure (Barrett et al. 1999; Grill et al. 2009, Hale et al. 2001) but limited information on the role of diversity in the response to SQPV infection.

The highest levels of genetic diversity within mammalian populations are located within genes involved in the immune response including those clustered together within the major histocompatibility complex (MHC), (Horton et al. 2004; Robinson et al. 2013). As a consequence, MHC loci are frequently used as a source of genetic markers in studies of population diversity and population health (Sommer 2005; Osborne et al. 2015). The MHC is divided into three major clusters of closely linked genes, class I, II and III. MHC class I and II genes encode proteins responsible for the presentation of small fragments of pathogen proteins for recognition by antigen specific receptors on CD8 or CD4+ T cells respectively (Bjorkman 1987; Germain and Margulies 1993). The specificity of the immune response is influenced by the range of pathogen peptides presented by MHC molecules. The majority of MHC diversity associated with the class II MHC loci locates to the second exon which determines part of the peptide binding groove. As a consequence, allelic diversity influences the range of peptides recognised by the immune system (Hughes and Yeager 1988; Hughes and Nei 1989) and many associations with susceptibility to autoimmune and infectious disease have been described (reviewed by Trowsdale 2011).

Earlier analyses of fragmented populations of European ground squirrel (Spermophilus citellus, Ricanova et al. 2011) and spotted suslik, (Spermophilus suslicus, Biedrzycka and Radwan 2008) described high
levels of allelic diversity at the class II MHC-DRB locus. Therefore, this study aims to characterise the DRB locus in red squirrels which will allow a comparison of diversity in fragmented UK red squirrel populations with populations from continental Europe.
**Materials and methods**

**Red squirrel samples**

Genomic DNA was prepared from 42 tissue samples obtained from red squirrels selected from archived material held at the Zoological Society of London (ZSL). These animals were found dead and submitted to the ZSL between 1996 and 2006 and represent three locations within mainland UK; Central Scotland, North West England, North East England and two island populations, the Isle of Wight and Jersey in the Channel Islands.

Twelve road kill samples were obtained from the stronghold population on the Isle of Arran located of the West coast of Scotland, six samples from South West Scotland, six samples from North Central Scotland, thirteen from Northern Scotland and three from Northern Ireland. Eighteen samples of continental European red squirrels were obtained from Belgium and Northern Italy. The location and number of animals sampled at each location is detailed in Figure 1. For comparative purposes, DNA was also prepared from an eastern grey squirrel from the South West of Scotland.

**Preparation of DNA**

Genomic DNA was extracted from muscle or spleen samples using the DNeasy blood and tissue kit (Qiagen) following the manufacturer’s instructions. The quantity and quality of DNA was estimated using a nanodrop spectrophotometer.

**Preparation of RNA**

Pseudogenes and gene fragments are common features of MHC regions in other mammalian species (Kumanovics et al. 2003). To provide evidence that the Scvu-DRB loci are functional, cDNA was prepared from mRNA isolated from the spleen of a red squirrel following euthanasia of a suspected case of squirrelpox from South West Scotland. The spleen was removed, suspended in RNAlater™ and archived at -20°C. Total RNA was prepared from 20 mg of spleen tissue using the Precellis Ribolyser Tissue RNA kit. Genomic DNA was also prepared from the same sample.

**Targeting the red squirrel DRB loci**

PCR primers Scvu351F and Scvu338R which amplify a 243 bp fragment of the second exon of the red squirrel DRB locus were designed using a DRB cDNA sequence from the tassel-eared squirrel (accession number M97616) as the template. Both primers are located within the second exon. The primer sequences are listed in
Table 1. Each PCR reaction was carried out in a final volume of 50 µl containing 200 nM of each primer, 1U Taq polymerase (Promega, Paisley, UK) and 50 ng of DNA template. Amplification reactions were performed under the following cycling conditions; 94°C for 4 minutes followed by 30 cycles of 94°C for 30 s, 60°C for 30 s and 72°C for 30 sec. A final cycle of 72°C for 5 min was added to complete the reaction.

Analysis of PCR products

The products of each PCR reaction were separated on a 1% agarose gel, stained with gel red and visualised under a UV transilluminator. PCR products were purified using the SV Gel and PCR Clean-Up System (Promega), quantified and sequenced in both directions using primers Scvu351F and Scvu338R. The forward and reverse sequences were aligned using the SeqManII™ program of the DNASTAR package and polymorphic positions identified. As the primers amplify the products of two polymorphic DRB1 loci in order to define the allelic diversity at each locus the PCR fragments are cloned.

Cloning and sequence analysis

Scvu-DRB alleles were cloned into the pGEM-T-easy vector (Promega) and individual clones identified by colony PCR. Digestion of the colony PCR product with the restriction enzyme Rsa I followed by resolution of the fragments on an 8% polyacrylamide gel allowed the selection of clones with identical restriction patterns for sequencing. Depending on the complexity of the direct sequence analysis, up to 12 clones were sequenced in both directions. Sequencing or Taq induced errors were eliminated through comparison with the direct sequence of the PCR product. The majority of alleles including those that differ by single nucleotide substitutions were identified multiple times from different DNA samples and in some cases from cDNA as well as genomic DNA. Those alleles identified from single samples were cloned and sequenced independently from two different PCR reactions to eliminate possible artefacts associated with amplification and cloning.

Red squirrel Class II DRB nomenclature

We followed the accepted convention of MHC allelic nomenclature proposed by Klein et al. (1990) - which uses the first two letters of the genus and species (Scvu) followed by the locus (Scvu-DRB1) and then an allele designation (Scvu-DRB1a, 1b, 1c, based on the order of their identification). DRB alleles were assigned to either the DRB1 or DRB2 locus depending on sequence similarity and phylogenetic clustering. The allelic nomenclature shown in Table 2 is used throughout.
Analysis of Scvu-DRB gene transcription

First strand cDNA was prepared using the ImProm-II RT system (Promega) in a 40 µl reaction using 200 ng of Total RNA. Using the full length DRB transcript from the tassel-eared squirrel (Sciurus aberti) as a template, primers Scvu363 and Scvu364 (listed in Table 1) were designed within exons 1 and exon 3 and tested for their capacity to amplify the Scvu-DRB transcripts. Reverse transcription-PCR was carried out in 50 µl reactions using each combination of forward and reverse primer, 3 µl of cDNA template and 200 nM of each primer in GoTaq polymerase master mix (Promega, Paisley, UK). Amplification reactions were performed under the following conditions; 94°C for 4 minutes followed by 30 cycles of 94°C for 1 min, 55°C for 1 min and 72°C for 1 min. Fragments were visualised on 1% agarose gels and those of the expected size were gel purified and cloned into the pGEM-T-easy vector as detailed above.

Sequence analysis

Scvu-DRB gene sequences were assembled from each bi-directional sequence using the SeqManII program. All polymorphic sites were inspected manually. All sequences have been deposited in the European Nucleotide Archive and assigned accession numbers listed in Table 2. Multiple alignments of the nucleic acid and predicted amino acid sequences were produced using Clustal Omega available on the EMBL-EBI website (http://www.ebi.ac.uk/Tools/msa/clustalo/). Multiple alignments of the Scvu-DRB sequences generated here and other published sequences were used to estimate maximum likelihood trees using PhyML-aLRT (Version 2.4.5) (Anisimova and Gascuel 2006) launched from TOPALi v2.5 (Milne et al. 2008). Prior to phylogenetic tree estimation, the model selection feature in the TOPALi v2 package which produces improved estimates of Likelihood values was used to select the nucleotide substitution model JC+G (Jukes and Cantor 1969). To test for positive selection we compared the average number of synonymous substitutions per synonymous site (dS) with the average number of non-synonymous substitutions per non-synonymous site (dN) for codons predicted to determine the antigen-binding sites (ABS), the remaining sites (non-ABS) and all sites. We used the modified Nei and Gojobori method with Jukes–Cantor correction as the substitution models. The codons predicted to determine amino acids associated with the APS were selected according to Reche and Reinherz (2003) and are shown in Figure 2. The average dN and dS and their variances estimated using 10000 bootstrap replicates were used to test the null hypothesis that H₀, dN=dS (test for neutrality) using a Z test. This analysis was carried out in MEGA version 6 (Tamura et al 2013). Rejection of the null hypothesis in favour of the
alternative hypothesis where $dN > dS$ where the probability values $P$ are less than 0.05 is considered evidence for positive selection.
Results

Identification of two Scvu-DRB loci

A 243 bp fragment of the second exon of the Scvu-DRB locus was initially amplified from 10 red squirrel DNA samples from UK population 1 (Figure 1). Sequence analysis of the PCR fragments identified 29 identical polymorphic positions in each of these 10 animals. The presence of two distinct sequences which were identical in all ten animals was confirmed through analysis of individual clones obtained from four of these animals. The two sequences did not appear to segregate as expected for alleles at a single locus as no animal homozygous for either allele was identified. Therefore, rather than alleles at a single locus, we concluded that they are likely to represent two independent DRB loci, inherited together within a single haplotype. All 10 animals genotyped appeared homozygous for this one haplotype. The presence of two independent and polymorphic DRB loci was confirmed through identification of alleles at each locus in animals from populations 11 and 12 from Belgium and Italy respectively. The sequences identified from population 1 were used as reference sequences for each of these loci and termed Scvu-DRB1a and Scvu-DRB2a (Supplementary Figure 1).

Are both Scvu-DRB loci transcribed?

Using primers Scvu363 and Scvu364 located in exons 1 and 3, three correctly spliced transcripts representing two alleles at locus 1, (Scvu-DRB1a and Scvu-DRB1b) and a single allele at locus 2, (Scvu-DRB2a) were identified in sample 15 from population 2 (Supplementary Table 1), confirming that both loci are transcribed and therefore likely to be functional. No polymorphic sites were identified in the genomic DNA primer binding sites within exon 2 suggesting that the genotyping primers are likely to amplify the majority of DRB allelic diversity in red squirrels. The genotyping of a DNA sample from the same squirrel produced an identical result to the cDNA analysis.

Scvu-DRB sequence analysis

Sequence analysis of the PCR products from the remaining 90 samples identified a range of nucleotide substitutions not present in population 1. Where novel and multiple substitutions were identified, individual alleles were resolved through cloning. A total of 19 Scvu-DRB1 alleles and 5 Scvu-DRB2 alleles were identified. The alleles associated with each squirrel sample are shown in supplementary Table 1. The sequences have been assigned ENA database accession numbers LN832043 to LN832063 as shown in Table 2. The nucleotide sequences of the 24 Scvu-DRB variants are shown in supplementary Figure 1 while the predicted amino acid...
sequences are shown in Figure 2. The Scvu-DRB1 locus is the more polymorphic of the two with 19 of the 24 alleles. Twenty seven polymorphic nucleotide positions corresponding to 15 non-synonymous substitutions were identified within the second exon of the Scvu-DRB1 locus compared with 16 polymorphic positions corresponding to 8 non-synonymous substitutions within the second exon of the Scvu-DRB2 locus. Allelic diversity at both DRB1 and DRB2 loci was generally associated with small numbers of nucleotide substitutions with many alleles differing at only one or two positions. Alleles DRB1a and 1b, DRB1m and 1n and DRB2b and 2c differ at single synonymous substitutions. Alleles DRB1e and 1h show the highest level of diversity with 90% identity while the most diverse DRB2 alleles, DRB2a and 2e, show 93% identity in pair-wise comparisons. Inter-locus diversity is greater with 85% identity between DRB1a and DRB2a.

Substantial allelic diversity within and between DRB1 and DRB2 loci is associated with positions predicted to directly interact with peptides bound within the peptide binding domain (Figure 2). Sixteen of the eighteen amino acid positions estimated by Reche and Reinherz (2003) to directly interact with peptides bound within the class II MHC peptide binding domain are shown to be variable or adjacent to a variable amino acid in red squirrels (Figure 2). As positive selection is thought to drive and maintain diversity at MHC loci we tested the hypothesis that dN > dS at codons predicted to determine the antigen-binding sites (ABS), the remaining sites (non-ABS) and all sites. This hypothesis was rejected in the analysis of all sites (dN-dS = 0.96, p = 0.17) and the non ABS (dN-dS = -0.73, p = 1.0) and only at ABS sites was the hypothesis supported (dN-dS = 2.663, p=0.004).

Phylogenetic analysis

The relationship between Scvu-DRB1 and B2 sequences was further explored by phylogenetic analysis using the nucleic acid alignment shown in supplementary Figure 1. The tree topology (Figure 3) generally supports the two locus hypothesis as the two major clusters are formed by the DRB1 and DRB2 allelic lineages, the only exception being Scvu-DRB1l which clusters independently of the other DRB1 alleles despite sharing many of the nucleotide and amino acid motifs characteristic of the DRB1 locus. This may be due to a recombination event between DRB1 and DRB2 loci. The S. aberti (Scab-DRB) and the S. carolinensis (Scca-DRB) sequences all cluster with the Scvu-DRB1 loci.

The distribution of Scvu-DRB1 and Scvu-DRB2 allelic diversity in UK and continental European red squirrels
The distribution and frequency of the 19 *Scvu-DRB1* alleles and 5 *Scvu-DRB2* alleles in UK and continental European red squirrels is shown in Figure 4 and Table 3 respectively. Twelve *Scvu-DRB1* and 4 *Scvu-DRB2* alleles were identified in the 18 animals from continental populations 11 and 12, while only 6 *Scvu-DRB1* and a single *Scvu-DRB2* allele were found in 55 samples obtained from six UK mainland populations. Both DRB loci were homozygous in 78% of animals from the mainland UK compared with 16% of the continental red squirrels.

The highest level of allelic diversity with 12 DRB alleles associated with 9 distinct haplotypes was identified in the population from northern Italy, while the population with least diversity was population 1 from central Scotland with only a single haplotype. These data indicate that the extensive allelic and haplotype diversity associated with continental European red squirrels is not present in UK populations analysed.

With the exception of population 10 from the Isle of Arran, the *Scvu-DRB1a/Scvu-DRB2a* haplotype dominates the UK population. This haplotype was not identified in the continental populations or in the small number of samples from the Channel Islands. Given the proximity of the Channel Islands to the French coast, it is not surprising that they share alleles with continental populations. However, population 10 shares allelic diversity with samples from Belgium rather than with other UK populations. This suggests that this population may have a more recent continental European origin.
In response to selection by rapidly evolving pathogens, genes associated with protective immunity are often highly diverse (Barreiro and Quintana-Murci 2010). Such diversity increases the probability of population survival in the face of novel infections whereas populations with limited diversity are less secure. A major source of immunological diversity is within the MHC where substantial allelic diversity is thought to be maintained by a form of balancing selection (heterozygous advantage and/or frequency dependent selection) arising from the requirement to respond to rapidly evolving or novel pathogens (Hughes and Yeager 1998; Meyer and Thomson 2001). High levels of allelic diversity at MHC loci are often associated with large populations with high levels of genetic exchange whereas low levels are often associated with smaller, more isolated populations (reviewed in Sommer et al. 2005; Radwan et al. 2010).

**Comparison of class II MHC DRB diversity in UK and continental squirrel populations**

While comparing diversity at the class II MHC Scvu-DRB locus in UK red squirrels with populations from continental Europe, we identified a duplication of the Scvu-DRB locus, described Scvu-DRB1 and Scvu-DRB2 transcripts and sequenced families of alleles at each locus. We provide evidence of positive selection at sites associated with the binding of peptide antigens in agreement with orthologous loci in other species (Babik et al. 2005, Cizkova et al. 2011). Limited Scvu-DRB1, Scvu-DRB2 allelic and haplotype diversity was identified in geographically distinct populations of red squirrel in the UK. A single DRB haplotype (DRB1a/DRB2a) appears to dominate the UK population with levels of homozygosity ranging between 68% and 100% depending on the population analysed. In contrast, substantial allelic diversity was identified in samples from continental Europe where Belgian and Italian populations provided 12 Scvu-DRB1 and 4 Scvu-DRB2 alleles from 18 animals compared with only 6 Scvu-DRB1 and a single Scvu-DRB2 allele in 55 samples from 6 populations from the UK mainland. While it is likely that some alleles present at lower frequencies will not have been recorded in both continental European and UK squirrels, it is clear that the extensive MHC diversity in continental European squirrels is not present in UK populations.

**Origin of the Scvu-DRB1a/Scvu-DRB2a haplotype**

The origin of the Scvu-DRB1a/Scvu-DRB2a haplotype which dominates the UK red squirrel populations is unclear. This haplotype may be a remnant from the original population that colonised the British Isles following the end of the last ice age between 7 and 10 thousand years ago when the UK remained connected with Western
Europe. The failure to identify this haplotype in the continental European or Channel Island populations supports this observation; however our analysis is limited to 18 animals from Italy and Belgium and is clearly not representative of the continental population as a whole. There is evidence that the original red squirrel population that colonised the British Isles was almost driven to extinction in the 18th century (summarised in Barratt et al. 1999). The lack of MHC diversity supports this extreme population bottleneck in which all but the most frequent alleles were lost due to inbreeding and drift. Historical records, confirmed by recent genetic analysis, indicate that animals from other parts of the UK and from Western Europe were re-introduced to restore lost or depleted UK populations including some from Scandinavia, re-introduced to secure populations in southern Scotland (Hale et al. 2004). The Scvu-DRB1a/Scvu-DRB2a haplotype may have originated with animals from Scandinavia which subsequently expanded throughout the UK. By extending future analyses to include samples from Scandinavia and other areas of Western Europe, the origin of the Scvu-DRB1a/Scvu-DRB2a haplotype may become clearer.

Consequence of limited DRB diversity in UK red squirrels

Consistent with functional class II MHC-DRB orthologues in other vertebrates, much of the allelic diversity is associated with non-synonymous substitutions at locations predicted to interact with peptides held within the peptide binding groove (Hughes and Nei 1989). Such diversity influences the range of peptides presented to CD4+ve T cells, one of the key regulatory cell types controlling both antibody and cellular responses to viral infection. Any reduction in the range of pathogen antigens available for recognition by the immune system may influence subsequent responses to infection at individual and population levels. However, the diversity between DRB loci suggests that each may present a distinct range of peptides for recognition by the immune system (Brown et al. 1993). Haplotypes with two diverse DRB loci will allow a wider array of peptides to be presented to T cells compared with haplotypes with only a single functional DRB locus. While this study has focused on the Scvu-DRB loci as a marker for MHC diversity, additional class II and class I loci will be included in future analyses, allowing a more complete picture of MHC haplotype diversity in squirrel populations from the UK and continental Europe.

Levels of MHC diversity in continental European red squirrels are consistent with a robust population associated with frequent genetic exchange between populations. This is in contrast to the limited diversity in the UK squirrels which is consistent with a strong founder effect which has led low levels of diversity in the remaining small isolated populations in the UK. Inbred wildlife populations are often susceptible to
environmental change including the introduction of new pathogens and SQPV appears to be responsible for much of the decline of the UK red squirrel population (Rushton et al. 2006). Wildlife populations within a stable environment are generally resilient to the endemic pathogens; a range of which (adenovirus; Sainsbury et al. 2001), (hepatozoon species; Simpson et al. 2006) (mycobacteria; Meredith et al. 2014) have been described in red squirrels in the UK. However, the impact of these infections appears limited compared with the exotic SQPV, although they might have a stronger impact on captive collections (Everest et al. 2014; Shuttleworth et al. 2014).

Providing evidence for a direct link between MHC diversity and squirrelpox disease susceptibility remains challenging as samples from healthy animals with evidence of SQPV exposure for comparison with samples from animals known to have been killed by the virus are required. While limited MHC diversity may contribute directly to the decline of the UK red squirrel population through a failure to present protective antigens for recognition by the immune system, it may also reflect a general decrease in diversity across the genome (reviewed by Sommer et al. 2005). Previous analysis of UK red squirrel population diversity using neutral markers such as the mitochondrial d-loop (Barratt et al. 1999) and a range of microsatellites (Hale et al. 2004; Grill et al. 2009) also identified limited diversity compared with continental populations.

Limited MHC diversity has been described in other species and populations which have gone through population bottlenecks. These include the cheetah, where limited diversity at the MHC has been linked to susceptibility to viral infection (O’Brien et al. 1985) and in the Tasmanian devil, where it has been linked with susceptibility to a transmissible tumour (Siddle et al. 2007). Limited MHC diversity has also been recorded in expanding populations following a population bottleneck, including the European Beaver (Ellergren et al. 1993) the European and North American Moose (Miko and Anderson 1995) and the Mountain Goat (Mainguy et al. 2007). These populations are however predicted to remain susceptible to novel pathogen infections. The red squirrel population of the UK may provide a warning to such populations as it may be the first recorded example of a wildlife population with limited genetic diversity that expanded following a population bottle neck in the 18th century as a result of reforestation efforts (Shorten 1954) only to be decimated by an exotic viral infection in the 20th century.

It may be fortuitous, but no evidence of SQPV has been reported in continental European red squirrels despite the introduction on at least three occasions of eastern grey squirrels to Northern Italy between 1948 and the 1990s, followed by numerous translocations and undocumented releases (Martinoli et al. 2010; Bertolino et al.
The Italian red squirrel population is the most genetically diverse population analysed in this study with a large number of diverse MHC haplotypes associated with high levels of heterozygosity. The absence of SQPV along with a genetically diverse red squirrel population and low levels of diversity in Italian grey squirrels (Signorile et al. 2014) may have contributed to the relatively slow spread of grey squirrels in Northern Italy compared with those in the UK (Bertolino et al. 2014).

**MHC diversity and red squirrel conservation**

Surprisingly, the distribution of alleles in the red squirrel population on the Isle of Arran, located off the West coast of Scotland, suggests that they are more closely related to those from continental Europe than to other squirrels from the UK. The Scvu-DRB1a/Scvu-DRB2a haplotype dominant in mainland UK populations was not recorded and existing records indicate that red squirrels were introduced to the island between the 1930s and 1950s. This supports an earlier study which identified two mitochondrial haplotypes in the Arran population, one of which was also found in Belgium populations (Barratt et al. 1999). As the Arran population appears unique in the UK we suggest that animals from this population could be used to expand levels of diversity and contribute to long term population health in other Scottish red squirrel strongholds (and potentially other areas of mainland UK) with established red squirrel populations with limited MHC diversity. This approach may be preferable to introductions from continental Europe with the risk of introducing exotic pathogens. Currently red squirrel reintroduction strategies in the UK are focused on controlling the grey squirrel population and habitat restoration which favours red squirrels with little regards to population genetic diversity. We suggest that by incorporating a simple measure of MHC diversity in the reintroduction strategy overall population health would be improved in the longer term.
Figure Legends

Fig. 1
Map of Western Europe showing the population number and number of red squirrels sampled from each location in parenthesis.

Fig. 2
Multiple alignments of the predicted amino acid sequences derived from three red Squirrel DRB1 and DRB2 transcripts aligned with nineteen DRB1 and DRB2 allelic sequences derived from the genomic analysis of 100 red squirrels from the UK and continental Europe. Only unique allelic sequences are included. The full length DRB transcript derived from the tassel-eared squirrel (Sciurus aberti, Scab-DRB) is used as the reference sequence. Sequences are numbered from the first amino acid of the mature protein. The portion of the DRB-β1 domain encoded by the second exon is shaded and amino acid positions predicted by Reche and Reinherz (2003) to interact with peptides within the peptide binding domain are indicated with a *. Sequence identity is indicated by a . and missing sequence is indicated by a -.

Fig. 3
Maximum likelihood tree estimating the relationships between Squirrel DRB nucleotide sequences. The tree is generated using the HKY substitution model and rooted using the murine DRB orthologue, H2-EB1, (NM_01382). Only bootstrap values 60 or above are shown. Species designations are as follows; Scab, Sciurus aberti (tassel eared squirrel, M97616); Scvu, Sciurus vulgaris (Eurasian Red squirrel, LN832043 to LN832063), Scca, Sciurus carolinensis (eastern grey squirrel).

Fig. 4
Distribution of Scvu-DRB1 and DRB2 allelic diversity in each red squirrel population.
<table>
<thead>
<tr>
<th>Primer</th>
<th>Specificity</th>
<th>Template/Location</th>
<th>Sequence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Scvu351F</td>
<td>DRB1 and DRB2</td>
<td>gDNA, exon 2</td>
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| Table 1. PCR primers |

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| Table 2. Scvu-DRB1/DRB2 allelic nomenclature and associated accession numbers |

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Legend Table 3, N = Number of individuals genotyped; *, Belgian population; #, Italian population
References


Shuttleworth CM, Everest DJ, McInnes CJ, Greenwood A, Jackson NL, Rushton S, Kenward RE (2014) Inter-specific viral infections: Can the management of captive red squirrel collections help inform scientific research? Hystrix, the Italian J Mammalogy 25:18-24


Fig. 2