PERMANENT GENETIC RESOURCES

Eight polymorphic microsatellite loci developed and characterized from Townsend’s big-eared bat, Corynorhinus townsendii

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Abstract

Two of the five subspecies of the western big-eared bat, Corynorhinus townsendii, are listed as federally endangered with the remaining three being of conservation concern. Knowing the degree of connectivity among populations would aid in the establishment of sound conservation and management plans for this taxon. For this purpose, we have developed and characterized eight polymorphic microsatellite markers.

Keywords: Corynorhinus townsendii, microsatellite, Townsend’s big-eared bat

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Townsend’s big-eared bat, Corynorhinus townsendii, is a North American bat of the family Vespertilionidae. There are five subspecies in the USA and Mexico (Piaggio & Perkins 2005) and two, Corynorhinus townsendii townsendii and Corynorhinus townsendii pallescens, are listed as species of Special Concern or sensitive species by state and federal agencies (Pierson et al. 1999), while Corynorhinus townsendii ingens and Corynorhinus townsendii virginianus are federally listed as Endangered. Although many local, state and federal agencies have developed management plans for C. townsendii that include monitoring and protection, little is known about their population structure or connectivity.

Population-level genetic data would significantly increase our understanding of C. townsendii. Only five of 15 microsatellite primers designed for other microchiropteran species (Burland et al. 1998; Vonhof et al. 2002) amplified and were variable in C. townsendii (Piaggio et al. in press). To increase the number of markers for this species, we developed and characterized eight new microsatellite loci.

Tissue samples were obtained from 25 individuals from Colorado (C. t. pallescens) and from 29 individuals from Idaho (C. t. townsendii). One individual from Colorado was used in the enrichment and development process. We targeted these localities because most current population-level research of C. townsendii is focused on these subspecies. We developed a microsatellite library following methods adapted from Glenn & Schable (2005). Genomic DNA was digested with the enzyme RsAl and fragments were ligated using double-stranded SNX-24 linkers. This library was hybridized to 12 biotinylated microsatellite oligonucleotide probes with dinucleotide and trinucleotide repeats (e.g., GT, CA, TG, CAC and CAG). Hybridized fragments were captured on streptavidin-coated Dynabeads (Dynal Biotech). Microsatellite-enriched fragments were amplified and cloned with the TOPO TA cloning kit (Invitrogen). Insert sequences from 96 colonies were obtained with M13 forward and reverse primers and visualized on an ABI 3730xl genetic analyser (Applied Biosystems). Forty-one clones had recognizable microsatellite sequences, of which 88% (36) had adequate flanking regions to design primers, which was accomplished with Staden package (Staden et al. 1998), trol(O (Castelo et al. 2002; Martins et al. 2006), and web-based Primer 3 (Rozen & Skaltsky 2000) software packages.

Polymerase chain reactions (PCR) were carried out using 0.5 μL each of 10 μm 5’ fluorescently end-labelled primers (Table 1), 3.0 μL nanopure water, 5.0 μL ReddyMix (AGene), and 1.0 μL of DNA (6–25 ng DNA/μL). The thermal profile

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for all loci was an initial denaturation at 94 °C for 2 min (B02, G07 and C02H08), 3 min (G12B11, H10E11 and E09B10) or 4 min (F09F10 and G02H10) followed by 35 cycles of 94 °C for 30 s, annealing at 51 °C (G12B11 and H10E11), 52 °C (E09B10, F09F10 and G02H10) or 55 °C (B02, G07 and C02H08), 3 min (G12B11, H10E11 and E09B10) or 4 min (F09F10 and G02H10) followed by 35 cycles of 94 °C for 30 s, annealing at 51 °C (G12B11 and H10E11), 52 °C (E09B10, F09F10 and G02H10) or 55 °C (B02, G07 and C02H08) for 45 s. Cycling was followed with a 7-min extension at 72 °C (B02, G07 and C02H08) or a 30-min extension at 60 °C. Of the 36 primer pairs that were designed and tested, eight pairs amplified and were variable in both populations.

PCR products were genotyped on an ABI 3130 genetic analyser and analysed with strand software (Hughes 1998; Locke et al. 2000). Genotypic disequilibrium between pairs of loci was tested using fstat 2.9.3 (Goudet 2001). Hardy–Weinberg Equilibrium (HWE), number of alleles and expected and observed heterozygosities were estimated in Arlequin (Excoffier et al. 2005) and each locus was tested for null alleles using Micro-Checker (van Oosterhout et al. 2004). We found no evidence of linkage disequilibrium between loci. The number of alleles ranged from six to 20 per locus (Table 1). Two loci in each population demonstrated significant deviations from HWE (Table 1) after sequential Bonferroni correction (Rice 1989); however, only locus Coto_H10F_E11R violated HWE in each population. Moderate (0.05–0.20), Chapuis & Estoup (2007) null allele frequencies were found at some loci (Table 1) which could be the result of a Wahlund effect or the presence of true null alleles, although we have no evidence of the latter since all individuals yielded amplification products (i.e. we found no null homozygotes). These eight new markers, plus previously characterized markers developed from other Vespertilionids (Piaggio et al. in press), now make it possible to undertake detailed population genetic studies of C. townsendii.

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References


