Polymerisms of the prion protein gene (PRNP) in Alaskan moose (Alces alces gigas)

H. J. Huson and G. M. Happ

Institute of Arctic Biology, University of Alaska Fairbanks, Fairbanks, AK 99775–7000, USA
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Source/description: The prion protein (PRNP) gene of mammals encodes a prion protein (PrP), which is expressed in many tissues including the brain. Misfolded PrP conformers are responsible for neurodegenerative diseases known as spongiform encephalopathies. Transmissible spongiform encephalopathies (TSEs) include bovine spongiform encephalopathy, ovine scrapie, human Creutzfeldt–Jakob disease and chronic wasting disease (CWD).1,2 In mule deer (Odocoileus hemionus), white-tailed deer (Odocoileus virginianus) and Rocky Mountain elk (Cervus elaphus), First found in Colorado, CWD has now been identified in the eastern USA, as far south as New Mexico and as far north as west-central Canada.1,3 Polymorphisms of PRNP appear to be linked to susceptibility to TSE in numerous species including free-ranging white-tailed deer4 and mule deer.5 In mule deer, the SS genotype at residue 225 is associated with a higher incidence of CWD.3 Differences in PrP amino acid sequence are believed to be species barriers to disease transmission.6 However, Wyoming moose sequences that were previously deposited in GenBank (AY225484 and AY225485) are similar to the sequence of Odocoileus. CWD has not been observed in Rocky Mountain moose (Alces alces shirasi) or in caribou at higher latitudes (Rangifer tarandus), yet both species overlap the geographical range of Odocoileus species. We report here the PRNP sequences for 44 Alaskan moose (Alces alces gigas).

Polymerase chain reaction conditions and sequence analysis: Genomic DNA was purified from blood samples of 44 moose (Alces alces gigas) that were sampled from eight locations across Alaska (Fig. S1). DNA purification protocols, primers, amplification conditions and sequence analysis methods are provided in Appendix S1.

Polymorphisms: Two unique sequences (i.e. alleles) were found in the sequences of 44 individual moose (DQ154297 and DQ154298); these differed only at codon 209. The allele encoding methionine was present with a frequency of 0.45, and the allele encoding isoleucine was present with a frequency of 0.55. The diploid genotypes did not depart significantly from Hardy–Weinberg predictions ($\chi^2 = 0.4$, $P < 0.01$).

Comments: The conservation of amino acid sequences in the PrP of moose, caribou and deer is striking (Table 1) and consistent with the fact that all three genera are in the subfamily Capreolinae. In comparison with caribou, Alaskan moose samples show six synonymous substitutions (bases 195, 231, 324, 360, 384 and 674), presumably reflecting purifying selection for the unique conformation of the globular N-terminal domain of cervid prions.7 CWD has been transmitted to moose by an oral route in an experimental laboratory setting.8 Genetic similarities, susceptibility in the laboratory setting and overlapping geographical ranges suggest the lack of a barrier to the transmission of prion disease from mule and white-tailed deer to moose. The absence of reports of CWD transmission to moose in natural settings may reflect ecological or epidemiological factors. Moose tend to be more solitary than deer of the genus Odocoileus, and dense social aggregations might be prerequisites of CWD epizootic outbreaks in cervids.

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References
8 Williams E. S. (2005) Vet Pathol 42, 530–49.

Correspondence: G. M. Happ (ffgmh@uaf.edu)

Supplementary Material
The following supplementary material is available as part of the online article from http://www.blackwell-synergy.com:
Appendix S1 Table of differences among prion alleles, Materials and Methods.
Figure S1 Locations of sites where moose were sampled in Alaska.

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**Table 1** A comparison of differences among the prion protein alleles of moose (DQ154297 and DQ154298), caribou (DQ154293) and mule deer (AY228473).

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<td>C</td>
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<td>T</td>
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<tr>
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<td>T</td>
<td>A</td>
<td>A</td>
<td>C</td>
<td>G</td>
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<td>V</td>
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Unlisted bases and amino acids were identical.
Supplementary Materials
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Supplementary Figure 1: Locations of sites where moose were sampled in Alaska.

Materials and Methods
Biological Materials - Whole blood samples were obtained from Alaska moose (Alces alces gigas). Samples were from eight different interior and southwest Alaska sites. Genomic DNA was extracted using the QIAGEN QIAamp DNA Blood Mini Kit.
QIAGEN protease was utilized for cell lysis. The genomic DNA was eluted with deionized water. Forty-four Alaska moose were successfully sequenced.

**Polymerase Chain Reaction** - PCR reactions were performed with 50-100ng DNA in a 50ul PCR reaction containing 1x PCR buffer as supplied by Qiagen (with no extra magnesium), 0.2mM each dNTP, 0.5pmol/ul of both forward and reverse primer, and 0.025U/ul Qiagen HotstarTaq Polymerase. A modified forward primer, Ce19v2-CTTTATTTTGCAGATAAGTC and reverse primer of Ce778-AGAAGAGATAATGAAAAACAGGAAG were utilized (O’Rourke et al. (1999) *J Gen Virol. 80, 2765-9.*). An MJ Research PTC-225 Peltier Thermal Cycler was used with the following settings: 95°C for 15min, then 40 cycles of 94°C for 30sec, 51.5°C for 30sec, and 72°C for 1min, with a final extension of 72°C for 10min.

**PCR Purification, Cycle Sequencing, Sequence Purification** - PCR purification was performed on all samples using the QIAGEN QIAquick PCR Purification Kit. All samples were eluted with 50ul deionized water. Cycle sequencing was performed using Big Dye Terminator Version 3 from Applied Biosystems. Sequence purification was performed with sephadex G-50 DNA grade and the samples sequenced on an Applied Biosystems 3100 Genetic Analyzer.

**Sequence Analysis** - Raw sequence data was analyzed using the Sequencher (GeneCodes Inc.) and BioEdit ([www.mbio.ncsu.edu/BioEdit/bioedit.html](http://www.mbio.ncsu.edu/BioEdit/bioedit.html)) programs to yield a 771 base pair product coding for the prion protein.