Research Paper

Prion sequence polymorphisms and chronic wasting disease resistance in Illinois white-tailed deer (Odocoileus virginianus)

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Abbreviations: BSE, bovine spongiform encephalopathy; CWD, chronic wasting disease; GPS, global positioning system; HWE, hardy-weinberg equilibrium; IDNR, Illinois department of natural resources; PRNP, prion gene; PrP, encoded prion protein; PrPsc, abnormal disease causing conformation of the prion protein; SNP, single nucleotide polymorphism; CJD, Creutzfeldt-Jakob disease

Key words: CWD, PRNP, synonymous polymorphism, cumulative polymorphisms, haplotype

Nucleic acid sequences of the prion gene (PRNP) were examined and genotypes compiled for 76 white-tailed deer from northern Illinois, which previously tested positive for chronic wasting disease (CWD), and 120 negative animals selected to control for geographic location and age. Nine nucleotide polymorphisms, seven silent and two coding, were found in the sampled population. All observed polymorphisms except two of very low frequency were observed in both negative and positive animals, although five polymorphic loci had significantly different distributions of alleles between infected and non-infected individuals. Nucleotide base changes 60C/T, 285A/C, 286G/A and 555C/T were observed with higher than expected frequencies in CWD negative animals suggesting disease resistance, while 153C/T was observed more than expected in positive animals, suggesting susceptibility. The two coding polymorphisms, 285A/C (Q95H) and 286G/A (G96S), have been described in white-tailed deer populations sampled in Colorado and Wisconsin. Frequency distributions of coding polymorphisms in Wisconsin and Illinois deer populations were different, an unexpected result considering the sampled areas are less than 150 km apart. The total number of polymorphisms per animal, silent or coding, was negatively correlated to disease status. The potential importance of silent polymorphisms (60C/T, 153C/T, 555C/T), either individually or cumulatively, in CWD disease status has not been previously reported.

Introduction

Chronic wasting disease is the only prion disease established in the western United States, and is found in white-tailed deer, mule deer, elk and moose.¹,² Factors related to CWD transmission are of particular interest because of its spread in the wild and ability to become established under natural population densities. Environmental transmission of CWD has been demonstrated in captive herd holding pens contaminated with carcass residue or fecal material,³,⁴ and may also occur by infectious prions present in blood and saliva of deer with CWD.¹ The first reported occurrences in wild deer and elk were found in Colorado and Wyoming in 1985. CWD has since spread slowly, often displaying large geographical gaps.⁶

The first positive cases of CWD east of the Mississippi river were detected during 2002 in Wisconsin,⁷ and another outbreak was discovered in Illinois later that same year.⁸ It is unclear whether the two outbreaks are related or of independent origin despite their geographic proximity of less than 150 km. Substantial differences exist between the sites, ranging from a moderately rugged agriculture and deciduous forest mix with distributed deer habitat in the Wisconsin outbreak to flat, large-field agriculture and highly urban landscapes with narrowly connected small habitat fragments in the Illinois outbreak.

Prion protein sequence is important in many aspects of prion disease including etiology, pathology and transmission. A number of PrP polymorphisms that alter resistance to prion disease have been documented in many species,⁹ and several may alter CWD susceptibility in deer. For example Q95H and G96S polymorphisms have been reported in white-tailed deer from Colorado,¹⁰ and south central Wisconsin,¹¹,¹² although small sample sizes precluded a clear conclusion about genetic susceptibility. In mule deer the S225F polymorphism was weakly related to CWD susceptibility.¹³ A S138N polymorphism reported in deer,¹¹,¹⁴ has subsequently been shown to reside within a processed pseudogene.¹⁵ White-tailed deer from the western United States may also have a G65E or A116G polymorphism,¹⁶ although these have not been related to CWD.

This research examined the relationship between prion polymorphisms and CWD disease status in free-ranging white-tailed deer from northern Illinois as indicated by allelic frequency distributions of positive and negative animals. Additionally, comparisons between PRNP polymorphisms in Illinois deer and those observed in other CWD outbreaks will provide a better assessment of the
underlying mechanisms that link PRNP with CWD susceptibility and resistance.

Results

PRNP sequences were determined for 196 deer, 76 CWD positives and 120 negative controls. In the 196 samples, nine single nucleotide polymorphisms (SNP) were detected (Fig. 1), eight of which had been previously described (nucleotide 286G/A,14 nucleotides 60C/T, 153C/T, 438C/T, 555C/T,16 nucleotide 285A/C,11 nucleotides 243T/A, 438C/T,10 nucleotide 676C/A EMBL AY425673).

Blast and literature searching indicated the 378G/A SNP has not been previously reported, consistent with our observation in a single individual. Seven of these polymorphic loci were silent and translated to a synonymous amino acid sequence while two of the SNP translated to a change in amino acid sequence of the protein; nucleotide 285A/C to amino acid Q95H and 286G/A to G96S. All of the observed allele frequencies of polymorphic alleles were less than 50% which indicated that the database derived consensus sequence was also the wild type genotype in Illinois white-tailed deer.

Polymorphisms and CWD status. Five of the nine polymorphic alleles were significantly related to CWD status according to Chi-square tests (Table 1). Loci 60, 285, 286 and 555 had higher than expected frequencies of polymorphisms in CWD negative animals while the opposite was true at locus 153 where observed frequencies of polymorphic alleles were higher than expected in CWD positive animals. Logistic regression also indicated that polymorphisms at loci 60, 285 and 286 were protective against CWD (p < 0.05) as each had odds ratios less than one (Table 1). All loci were in Hardy-Weinburg equilibrium (HWE) except for locus 286, which was found to significantly deviate from the expected frequency (HWE p < 0.01).

The relationship of CWD with polymorphic genotype frequency (Table 2) was similar to that for polymorphic allele frequency. Individuals heterozygous for polymorphisms at 60, 285 and 286 had significantly reduced odds ratios. However, the low number of individuals homozygous for these polymorphisms prevented calculation of meaningful risk reduction for these genotypes. Polymorphism at locus 555 was more common (42.4% overall) and individuals with a homozygous 555T/T genotype (Table 2) had a reduced odds ratio of 0.29 for CWD. However, the odds ratios for the polymorphic 555T allele (Table 1), as well as for the heterozygous genotype 555C/T (Table 2), overlapped 1 and so were not considered significant.

The polymorphism at locus 153 was a risk factor for CWD (Table 1), with 153T alleles more than doubling the risk of being CWD positive (odds ratio of 2.66). Similarly, heterozygous genotypes resulted in a 2.6-fold increase in CWD risk (Table 2). Although only four deer were found to be polymorphic homozygous at 153 (153T/T), this genotype produced a near 6-fold increase in the odds of CWD. Confidence limits for 153T/T, however, included 1 and so were not considered significant.

Statistical power to detect differences between positive and negative animals was good for most alleles although the low frequencies observed for some polymorphic alleles limited power. For example, the polymorphic coding alleles had power of 0.70 and 0.99 with an α of 0.05 and a two-tailed test. In contrast, the frequency of individual polymorphic alleles at 243, 378, 438 and 676 resulted in a power of less than 0.50 limiting the potential to detect an actual difference between positive and negative animals in this study.

Linkage disequilibrium between loci was observed in both positive and negative populations with the strongest association occurring...
between loci 60 and 285 ($\chi^2 = 70.5$, p < 0.01). Fifteen deer were polymorphic at either locus 60 or locus 285 and nineteen deer had both polymorphisms. Including both correlated loci in a multiple logistic regression model resulted in adjusted odds ratios of 0.18 and 0.21 for heterozygous genotypes 60C/T or 285A/C respectively.

Haplotype frequencies were estimated from unphased genotypes and the nine most common haplotypes (frequencies greater than 1%) are reported in Table 3. These haplotypes were present in 97% of deer sampled. Thirty-three different haplotypes were reconstructed in the sampled population with three haplotypes being distinct to the CWD positive population and nine of the haplotypes restricted to negative animals. The permutation test performed on the haplotypes determined that CWD positive and negative animals had significantly different distributions of haplotype frequencies (p < 0.01), suggesting that genetic variation between CWD positive and negative populations exceeds genetic variation within the two groups. Unfortunately, low frequencies in the majority of haplotypes prevented us from using the phased data in further exploratory analyses.

In the population sampled, 89% of the CWD positive animals had the consensus genotypes at both coding polymorphisms while 57% of the negative animals had the consensus genotypes at these loci (285A = Q95 and 286G = G96). No deer with two coding SNP, either homozygous for either polymorphic coding allele, or heterozygous for both coding alleles, were CWD positive although negative deer with these genotypes were observed (p < 0.01).

Because it appeared an additional copy of coding SNP reduced the percentage of CWD positive animals, we examined the relationship of multiple polymorphisms and disease status (Fig. 2). If deer were grouped by the cumulative number of polymorphic alleles per animal, a significant Wald Chi-Square was observed for the model (p < 0.01). Reverse coding or omission of locus 153 from models did not alter outcome, as both models were significant and produced similar odds ratio estimates (p < 0.01).

To examine relative contributions of expressed and silent changes to CWD resistance, coding and silent polymorphisms were modeled separately and combined for multivariate analysis. Single variable models produced odds ratios of 0.19 for each coding polymorphic allele, and odds ratios of 0.52 for each silent polymorphic allele (Fig. 2). Multivariate modeling of silent and coding polymorphisms resulted in significant Wald Chi-Square (p < 0.05) tests for both

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**Table 2: Ratio of observed and expected genotype frequencies for nucleotide polymorphisms**

<table>
<thead>
<tr>
<th>Locus</th>
<th>Genotype</th>
<th>n</th>
<th>Frequency in negatives</th>
<th>Frequency in positives</th>
<th>Odds ratio</th>
<th>Wald 95% confidence interval</th>
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<tbody>
<tr>
<td><em>60</em>*</td>
<td>CC</td>
<td>166</td>
<td>78.4</td>
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<td>0.21</td>
<td>0.07–0.63</td>
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<td></td>
<td>CT</td>
<td>29</td>
<td>20.8</td>
<td>5.3</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>TT</td>
<td>1</td>
<td>0.8</td>
<td>0</td>
<td>nc</td>
<td></td>
</tr>
<tr>
<td>*153</td>
<td>CC</td>
<td>155</td>
<td>85.8</td>
<td>68.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>CT</td>
<td>37</td>
<td>13.3</td>
<td>27.6</td>
<td>2.60</td>
<td>1.25–5.40</td>
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<td></td>
<td>TT</td>
<td>4</td>
<td>0.8</td>
<td>4.0</td>
<td>5.94</td>
<td>0.60–58.52</td>
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<td>243</td>
<td>TT</td>
<td>191</td>
<td>97.5</td>
<td>97.4</td>
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</tr>
<tr>
<td></td>
<td>TA</td>
<td>4</td>
<td>1.7</td>
<td>2.6</td>
<td>ns</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AA</td>
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<td>0.8</td>
<td>0</td>
<td>ns</td>
<td></td>
</tr>
<tr>
<td><em>285</em>*</td>
<td>AA</td>
<td>173</td>
<td>83.3</td>
<td>96.1</td>
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<tr>
<td></td>
<td>AC</td>
<td>22</td>
<td>15.8</td>
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<td>0.06–0.76</td>
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<tr>
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<td>0</td>
<td>nc</td>
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<tr>
<td>*286</td>
<td>GG</td>
<td>153</td>
<td>68.3</td>
<td>93.4</td>
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<tr>
<td></td>
<td>GA</td>
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<td>22.5</td>
<td>6.6</td>
<td>0.21</td>
<td>0.08–0.59</td>
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<td>9.2</td>
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<td>99.2</td>
<td>100</td>
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<tr>
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<td>0</td>
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<td></td>
</tr>
<tr>
<td></td>
<td>AA</td>
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<td>0</td>
<td>0</td>
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<tr>
<td>438</td>
<td>CC</td>
<td>179</td>
<td>90</td>
<td>93.4</td>
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</tr>
<tr>
<td></td>
<td>CT</td>
<td>17</td>
<td>10</td>
<td>6.6</td>
<td>ns</td>
<td></td>
</tr>
<tr>
<td></td>
<td>TT</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>ns</td>
<td></td>
</tr>
<tr>
<td>*555</td>
<td>CC</td>
<td>65</td>
<td>31.6</td>
<td>35.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>CT</td>
<td>96</td>
<td>44.2</td>
<td>35.6</td>
<td>1.14</td>
<td>0.60–2.16</td>
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<tr>
<td></td>
<td>TT</td>
<td>35</td>
<td>24.2</td>
<td>16.0</td>
<td>0.29</td>
<td>0.11–0.80</td>
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<tr>
<td>676</td>
<td>CC</td>
<td>192</td>
<td>96.7</td>
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</tr>
<tr>
<td></td>
<td>CA</td>
<td>3</td>
<td>2.5</td>
<td>0</td>
<td>ns</td>
<td></td>
</tr>
<tr>
<td></td>
<td>AA</td>
<td>1</td>
<td>0.8</td>
<td>0</td>
<td>ns</td>
<td></td>
</tr>
</tbody>
</table>

Bold text indicates loci where nucleotide sequence resulted in a change of expressed amino acid; other polymorphisms were synonymous. *Indicates a significant association (p < 0.05) between each genotype and CWD status using chi-square expected frequency obtained from PROC FREQ. Odds ratio value obtained with PROC LOGISTIC using each SNP as a class variable regressed against CWD as an outcome. Only odds ratios with significant (p < 0.05) parameter estimates are reported. Polymorphisms with non-significant odds ratios and 95% confidence limits overlapping 1 are denoted by ns. **Indicates loci with significant linkage disequilibrium (p < 0.01). "nc" indicates no cases so odds ratios were not calculable.
variables and significant Log Likelihood ratio tests (p < 0.01). Including independent variables for both coding and silent polymorphisms also decreased AIC values compared to single variable models, indicating that multivariate analysis improved the overall fit of the regression model.

We found no relationships between CWD status and age or gender although the experimental design was not intended to test this question. Polymorphisms were not related to age or gender according to chi-square tests. Protease resistant PrP was detected by immunohistochemistry in retropharyngeal lymph nodes of all CWD positive deer and in obex from just over 75% of positive deer. No relationship was apparent between genotypes and presence or absence of protease resistant PrP in the obex.

The non-coding pseudogene was present in approximately 15% of the animals sampled. Sequencing of the pseudogene PCR products revealed that the observed polymorphisms were indeed present in the functional PRNP gene nucleotide sequences, and were not detected in the pseudogene. All pseudogene PCR products displayed a 138S/N substitution, as had been previously reported. Presence of the pseudogene was not significantly correlated to CWD test status, age or gender of the animal according to chi-square tests.

Geographic variation in polymorphisms. Polymorphisms with significant association to disease status were distributed across the study area and have been found in other geographical locations. Low frequency polymorphisms had a non-uniform distribution across the study area, as would be expected for a gene introduced by a recent migrant or mutation, but low numbers limited spatial analysis. The same coding polymorphisms observed in Illinois have been reported in Wisconsin deer 150 km to the northwest. Based on our observed genotype frequencies and published data,12 the proportion of deer with consensus genotypes were similar. Approximately 57% of the Illinois negative deer had a homozygous consensus sequence compared to 51% of Wisconsin negative deer, while 95% of the Illinois positive deer had a consensus sequence compared to 82% of the Wisconsin CWD positive deer. However, genotype frequencies of the two coding polymorphisms (Fig. 3) differed according to chi-square test (p < 0.05). Illinois had a higher frequency of the Q95H polymorphism than Wisconsin with 17% percent of negative deer heterozygous or homozygous for Q95H in Illinois while only 4% of Wisconsin negative deer had this polymorphism. Conversely, G96S was more common in the Wisconsin herd with 44% of negative and 17% of CWD positive deer heterozygous or homozygous for this polymorphism compared to 32% of negative and 7% of CWD positive deer from Illinois. Deer with either SNP had a lower susceptibility to CWD than those with the consensus in both states.

**Discussion**

Polymorphisms that alter resistance to prion disease are present in many species. For example, in sheep PrP containing the V136 R154 Q171 polymorphic sequence confers scrapie susceptibility while a combination of A136 R154 R171 alleles confers scrapie resistance. Selection for the resistant alleles has been used in the European Union to reduce the prevalence of scrapie. Wildlife managers are similarly interested in the potential of genetic resistance in managing CWD, both for understanding or predicting spread of disease, and for the possibility of use in repopulation following herd reductions. Furthermore, information about polymorphisms and susceptibility in additional species could contribute to the general understanding of prion disease mechanisms.

This study identified allelic variants at nucleotide positions 60, 285, 286 and 555 (codons 20, 95, 96 and 185) of the white-tailed deer prion gene associated with resistance and at nucleotide 153 (codon 51) associated with susceptibility to CWD. The association with resistance to CWD of polymorphism 285A/C which codes for Q95H and 286G/A which codes for G96S, confirms previous reports of a genetic trend or association in enclosed herds, and in a free-ranging white-tailed deer population. Recently it was reported that S96 (286G/A) transgenic mice inoculated with CWD displayed high levels of resistance to multiple strains of the disease when compared to their G96 transgenic counterparts, which is consistent with a genetic trend in wild herds.

### Table 3 Reconstructed frequencies of common haplotypes for CWD positive animals and controls

<table>
<thead>
<tr>
<th>Haplotype definition</th>
<th>Frequency in (−) animals</th>
<th>Frequency in (+) animals</th>
</tr>
</thead>
<tbody>
<tr>
<td>CCTAGGCCC</td>
<td>33.0</td>
<td>39.8</td>
</tr>
<tr>
<td>CCTAGGGCTC</td>
<td>28.1</td>
<td>31.6</td>
</tr>
<tr>
<td>CTTAAGCTC</td>
<td>12.2</td>
<td>2.6</td>
</tr>
<tr>
<td>CTTAGGGCCC</td>
<td>10.3</td>
<td>16.3</td>
</tr>
<tr>
<td>TCTCGGCCC</td>
<td>4.6</td>
<td>0.0</td>
</tr>
<tr>
<td>CTTAGGTCC</td>
<td>3.9</td>
<td>2.8</td>
</tr>
<tr>
<td>TCTAGGCC</td>
<td>1.4</td>
<td>1.8</td>
</tr>
<tr>
<td>TCTAGGCCA</td>
<td>1.3</td>
<td>0.0</td>
</tr>
<tr>
<td>CTTCGGCC</td>
<td>1.1</td>
<td>1.5</td>
</tr>
</tbody>
</table>

Haplotypes were generated from unphased genotypes with the Bayesian (ELB) algorithm in Phase ver 2.1. The distribution of haplotype frequencies was significantly different between CWD positive and CWD negative animals (p < 0.01). Only haplotypes with frequencies greater than 1% are reported.

![Figure 2. Probability of CWD as predicted by the number of polymorphic alleles.](image)
by analogy with herd immunity, resistant populations might be important in CWD spread within the proportion of even fairly rare individual genotypes with a degree of genetic resistance might be important in CWD spread within the proportion of even fairly rare individual genotypes with a degree of genetic resistance. O'Rourke, et al. reported three polymorphisms to disease status is the close proximity of Q95H and G96S to proteolytic cleavage sites at amino acid residues 82 and 93. In human Creutzfeldt-Jakob disease (CJD), polymorphisms in the amino acid sequence have been associated with alterations in location of these cleavage sites. In addition, polymorphisms quite distal to cleavage sites may alter conformation sufficiently to change proteolytic susceptibility. Prion protein from human homozgyous for M129 has a secondary cleavage site shifted towards the C-terminus of the protein. At higher copper concentrations, each Cu²⁺ ion is coordinated by a single histidine imidazole and an amide nitrogen from glycine. At lower copper concentrations, all six histidines are involved in binding two Cu²⁺ ions. The Q95H polymorphism may represent a gain in potential copper binding in vivo. If this is the case, copper binding would appear protective since Q95H animals were more likely to be CWD negative.

An alternative explanation for the relationship of the coding polymorphisms to disease status is the presence of a secondary cleavage site shifted towards the C-terminus. Synonymous or silent polymorphisms are non-coding and do not alter the amino acid sequence of translated protein. Furthermore, none of our observed synonymous polymorphisms have been previously related to CWD susceptibility. So it was initially surprising that one of the lower odds ratios for CWD corresponded to the 60C/T polymorphism which is not only silent but also codes a region altered phenotype. First, a synonymous sequence may be linked to an expressed sequence elsewhere in the same protein or in a different protein. Such a case occurs in atypical, asymptomatic scrapie, where two synonymous polymorphisms in PRNP have an apparent linkage with the ARH allele, which is under-represented in atypical scrapie. This linkage results in the synonymous polymorphisms also being under-represented in atypical scrapie despite not coding for a protein difference. Linkage that may occur between closely related proteins is illustrated by a synonymous polymorphism in the prion related doppel gene that is associated with BSE in Fleckvieh cattle. Similarly, a synonymous polymorphism within acetyl-coenzyme A dehydrogenase is illustrated by a synonymous polymorphism in the PRNP gene that is associated with BSE in Fleckvieh cattle.
carboxylase, the rate limiting enzyme in lipogenesis, that is positively associated with both milk fat and lactose content,\(^5\) indicates potential linkage across multiple proteins.

Alternatively to linkage effects, synonymous polymorphisms may cause phenotypic variation by less well understood mechanisms such as by altering RNA stability, tRNA binding or translational protein folding. There is considerable evidence for conformational variants of PrP\(^{Sc}\) that transmit distinct pathologies or prion disease strains into animals with identical protein sequence,\(^3\) so it is conceivable a translation folding event could alter disease susceptibility. There is also some evidence that RNA mediates enhanced PrP folding during PrP\(^{Sc}\) conversion,\(^5\) and so polymorphic mRNA might also alter translational folding and subsequently, potential for PrP\(^{Sc}\) formation.

This scenario has some support in a recent observation that multiple synonymous polymorphisms have an additive effect on protein function, similar to our observation in Figure 2. The MDR1 gene encodes a membrane transport protein important in multiple drug resistance which contains a single synonymous polymorphism, 3435C/T, that sometimes has an effect on drug transport,\(^6\) as a result of increased mRNA turnover. However, with the addition of two or three further synonymous polymorphisms in MDR1, drug transport is consistently reduced at equivalent mRNA and protein levels in a variety of cell lines.\(^5\) Conformation dependent antibody inhibition indicated reduced function was a result of altered protein folding which would have become progressively different as the number of synonymous polymorphisms increased. The inverse relationship between number of SNP, whether silent or coding, and CWD status could be the result of a similar process.

Although we examined only polymorphisms within \(PRNP\) recent reports of multiple insertion-deletion polymorphisms located in the promoter region of bovine \(PRNP\) that are significantly associated with BSE,\(^5\) demonstrate an additional mechanism for synonymous polymorphisms to impact disease. Changes in the promoter region of \(PRNP\) apparently alter binding of transcription factors which in turn affect PrP expression.\(^5\)\(^,\)\(^6\)

In addition to genetics, both environmental and physiological factors such as deer densities, habitat composition, and age must be considered when interpreting disease occurrence and pathology. In Wisconsin for example, it appears that older deer have a higher prevalence of CWD than younger animals.\(^7\) Differential binding capacities of various soil particles to PrP\(^{Sc}\)\(^6\) suggests that environmental factors, such as soil composition, could influence CWD infection and transmission. These epigenetic factors may have contributed to the inability of our data to confirm a relationship between genotypes and histochemical locations of resistant PrP in lymph node or oesophagus, which can be indicative of disease progression.\(^10\)

In general, it appears likely that most white-tailed deer residing in the northern Illinois study area have some degree of genetic susceptibility to CWD despite differences in genotypic frequencies between positive and negative animals. Except for SNP with low frequency (2.5% or less), CWD was observed in deer heterozygous for all observed polymorphisms. The association between the level of allelic variation within the study area and CWD status has interesting implications for CWD management and understanding of CWD transmission in free ranging animals. If further research shows that genetic diversity within the \(PRNP\) gene does reduce disease incidence or transmission, it would emphasize the importance for CWD management teams to understand and manage ecological barriers to gene flow in deer while conducting disease surveillance efforts.

**Methods**

**Deer sampling.** A CWD surveillance program for free-ranging wild white-tailed deer, consisting of public hunting in the traditional late fall season and sharp shooting by field biologists in the early spring, was conducted by the Illinois Department of Natural Resources (IDNR) between Winter 2002 and Spring 2006. Hunter harvested deer were located to the nearest section, based on hunter description of locations using detailed topographical maps at deer check stations. A section is a 1.6-by-1.6 square km parcel of known location. Deer population control efforts by field biologists provided detailed spatial information about individual deer including Global Positioning System (GPS) coordinates and time of harvest. Demographic data, including sex and age estimated by dentition were collected from all deer sampled.

Samples of oesophagus and retropharyngeal lymph nodes were tested using USDA approved immunohistochemical procedures to detect prion protein resistant P\(^{3-4}\) at the Illinois Department of Agriculture Diagnostic Laboratories in Galesburg or Centralia and most positive samples were confirmed at the National Veterinary Services Laboratories. When possible, frozen tissue samples were archived for both CWD positive and negative deer.

Samples from all CWD positive deer with frozen tissue available were used for DNA extraction, PCR and \(PRNP\) sequencing. Given the nature of sample collection during surveillance efforts, we could not do a perfect paired-case control design though for each CWD positive case, negative controls were selected on the basis of age, sex and geographic location. We controlled for age estimated by dentition because of reported age dependency for CWD prevalence in Wisconsin.\(^7\) The prevalence of CWD has also been reported to be gender dependent,\(^1\) so whenever possible, both female and male negative controls of the same age were selected for each CWD positive deer. Furthermore, with the geographic spread of new CWD cases over time and the fragmented habitat in the study area, we felt it was important to also control for geographic variation in the samples. Deer harvest locations were known to the nearest section (1.6 x 1.6 km) and controls were selected on this basis. To check for geographic control we used ArcGIS and calculated that 66% (50 of 76) of positives were from the same section as their matched negative control, 18% (14 of 76) from an adjacent section, and 16% (12 of 76) were 3.2–8.2 km from the controls. These longer distances are not problematic when compared to deer movements in habitat similar to ours (i.e., intensive row-crop agriculture). For example, Brinkman et al.\(^1\) found the mean distance between summer and winter home range was 10.1 km for female deer in a landscape dominated by row-crops and over 85% of all deer moved. Furthermore, winter home range sizes average 5.2 km\(^2\) (2–6 sections). In habitats similar to our study area, Nixon et al.\(^1\) evaluated white-tailed deer dispersal in Illinois agricultural environments, their findings indicate average distance moved between 30 and 44 km for males and 37 to 41 km for females in East central, West central and Northern Illinois. Thus, 89% of our controls likely fell within the same home range as the positive and
the other 15% were within seasonal movement ranges. Whenever possible control animals were selected from the same harvest year as the CWD case.

The study area included four counties in northern Illinois with less than 1/3 of the area containing suitable deer habitat as described by Roseberry and Woolf. The mean age of deer in the study was estimated to be 1.8 years based on dentition and consisted of 72 males (30 CWD positive) and 124 females (46 CWD positive). All of the deer tested were free-ranging but all came from counties with a human population exceeding two/hectare, which is three times greater than the state average population density excluding the city of Chicago. Both CWD positive and control animals were sampled from deer habitat within metropolitan and rural areas.

**PCR.** Genomic DNA was isolated from skeletal muscle using the Wizard Genomic DNA Purification Kit (Promega, Madison, WI) in accordance with manufacturer’s instructions. Forward and reverse PCR primers previously published,11 were used to amplify the coding region for the mature prion protein. The forward primer, CWD-13 (5'-TTTTGTCAGATAATGTCATCAGTGTGAAA-3') overlaps the junction of Intron 2 and Exon 3 of the prion gene. The reverse primer CWD-LA (5'-AGAAGATAATGAAAACAGGAAG-3').10 If the pseudogene was contained within the functional gene, an additional primer set forward primer 369 and Comparative Genomics using a BigDye Terminator Sequencing Kit (Applied Biosystems) were used to align each sequence with the consensus to identify Single Nucleotide Polymorphisms (SNP).

**Workbench, SDSC) were used to align each sequence with the consensus to identify Single Nucleotide Polymorphisms (SNP).**

**Analysis.** The CLUSTALW and FASTA alignment programs (Biology Workbench, SDSC) were used to align each sequence with the consensus to identify Single Nucleotide Polymorphisms (SNP). The Keck Center yielded the highest quality reactions with 99.8% sequence homology between duplicates. The Wizard Genomic DNA Purification Kit (Promega, Madison, WI) in accordance with manufacturer’s instructions. Forward and reverse PCR primers previously published,11 were used to amplify the coding region for the mature prion protein. The forward primer, CWD-13 (5'-TTTTGTCAGATAATGTCATCAGTGTGAAA-3') overlaps the junction of Intron 2 and Exon 3 of the prion gene. The reverse primer CWD-LA (5'-AGAAGATAATGAAAACAGGAAG-3').10 If the pseudogene was detected in an animal, both the pseudogene and functional gene PCR reactions were sequenced to determine the true origin of polymorphisms.

A 40 μl reaction was assembled for amplification of the functional gene and pseudogene consisting of 24.2 μl of dH2O, 0.8 μl of 10 mM dNTP, 8 μl of TaqMaster, 4 μl of 1× TaqBuffer, 0.4 μl of Eppendorf Taq (5 Units/μl), 0.8 μl of 10 mM forward primer, 0.8 μl of 10 μM reverse primer and 1.1 μl (=20 μg/μl) of genomic DNA. Thermal cycler conditions for pseudogene and functional gene PCR are as previously described.10

**PRNP sequence evaluation.** PCR products were purified using the Wizard SV Gel and PCR Clean-Up System (Promega) and then sequenced at the University of Illinois Keck Center for Functional and Comparative Genomics using a BigDye Terminator Sequencing kit and an ABI 3730XL DNA Sequencer (Applied Biosystems). To check sequence accuracy and repeatability, replicate samples of DNA were submitted and also sent to two off-site sequencing centers. The Keck Center yielded the highest quality reactions with 99.8% sequence homology between duplicates.

CodonCode Aligner software (CodonCode Corp.,) was used to evaluate chromatograms produced from the ABI sequencing reactions. Only nucleotide sequences possessing Phred quality scores of 20 or higher were imported into Biology Workbench for further analysis. The CLUSTALW and FASTA alignment programs (Biology Workbench, SDSC) were used to align each sequence with the consensus to identify Single Nucleotide Polymorphisms (SNP).

Observed polymorphisms were compared to a database derived consensus sequence. All previously published white-tailed deer prion nucleotide sequences from the GenBank and EMBL-Bank databases were used to compile a consensus PRNP sequence using MultAlin software,21 with a protein translation generated from the nucleotide consensus sequence using Six Frame software (Biology WorkBench, SDSC). A Blast search indicated the consensus nucleotide sequence was not equivalent to any individual published sequence. The similar AF156185.1 (EMBLCDs:AAF80284) had T at the 438C/T SNP while AY286008.1(EMBLCDs:AAP37447) had T at the 555C/T SNP. The protein translated sequence of the consensus sequence was equivalent to both of these entries.

**Statistical analysis.** Logistic regression and chi-squared tests were performed with SAS for Windows, version 9.1 (SAS Institute Inc.). Chi-square tests were performed for each polymorphic allele and genotype to check for differences between observed and expected frequencies in both CWD negative and positive animals using an α level of 0.05. When needed, the Mantel-Haenzel Chi-square Test was employed to correct for low frequency genotypes. Deviations from Hardy-Weinburg equilibrium were tested using the program GenAlEx version 6.22

Additionally, logistic regression was used to determine the relationship between polymorphisms and disease status. Each allele was classified as 1 (polymorphic) or 0 (consensus) and regressed against the probability of being CWD positive. Genotypes were also compared to consensus genotypes in an additive model as discussed by North et al.23 In this method genotypes were classified as 0 = homozygous for the consensus genotype, 1 = heterozygous, 2 = homozygous for the polymorphism. For all logistic regression models, odds ratios less than 1 were equated with probability of disease below 50% and were therefore considered protective against CWD. Odds ratios greater than 1 were equated with probability of disease greater than 50% and were therefore considered risk factors for CWD.24

To determine the cumulative effects of polymorphisms, the total number of polymorphic alleles per animal was summed across all loci and regressed against CWD status. Observed allele frequencies were higher than expected for locus 153, so coding for this variable was reversed for cumulative regression models (2 = homozygous for the consensus genotype, 1 = heterozygous, 0 = homozygous for the polymorphism). The total number of polymorphic alleles (0, 1, 2, 3, 4, 5 or 6) was examined in one model, the number of coding polymorphic alleles (0, 1 or 2) was examined in a second model, silent polymorphic alleles (0, 1, 2, 3, 4, 5 or 6) were evaluated in a third model, and coding and silent polymorphic alleles were tested simultaneously in a fourth model. To be sure that the susceptibility conferred by locus 153 did not have excessive influence, models were also run with this polymorphism omitted.

To determine our ability to detect significant differences in polymorphisms between positive and negative animals, power analyses were performed for each polymorphic allele and cumulative logistic regression predictor variables using SamplePower for Windows, version 2.0 (SPSS, Inc.).

Haplotypes were generated from unphased genotypes using Phase, version 2.1.25.26 This program was also used to test for linkage disequilibrium between genotypes, and random distribution of haplotype frequencies between cases and controls. We also used logistic regression and chi-square tests to determine the relationship between disease status and sex, age and gender. Finally, allele frequencies of the Q95H and G96S coding polymorphisms were compared...
between the Illinois and Wisconsin outbreaks. Wisconsin allele frequencies were calculated from published data. 12

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