Disparate Modes of Evolution Shaped Modern Prion (PRNP) and Prion-Related Doppel (PRND) Variation in Domestic Cattle

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Abstract

Previous investigations aimed at determining whether the mammalian prion protein actually facilitates tangible molecular aspects of either a discrete or pleiotropic functional niche have been debated, especially given the apparent absence of overt behavioral or physiological phenotypes associated with several mammalian prion gene (PRNP) knockout experiments. Moreover, a previous evaluation of PRNP knockout cattle concluded that they were normal, suggesting that the bovine prion protein is physiologically dispensable. Herein, we examined the frequency and distribution of nucleotide sequence variation within the coding regions of bovine PRNP and the adjacent Doppel (PRND) gene, a proximal parologue to PRNP on BTA13. Evaluation of PRND variation demonstrated that the gene does not depart from a strictly neutral model of molecular evolution, and would therefore not be expected to influence tests of selection within PRNP. Collectively, our analyses confirm that intense purifying selection is indeed occurring directly on bovine PRNP, which is indicative of a protein with an important role. These results suggest that the lack of observed fitness effects may not manifest in the controlled environmental conditions used to care for and raise PRNP knockout animals.

Introduction

Transmissible spongiform encephalopathies (TSE) are a class of fatal neurodegenerative diseases that affect humans as well as livestock and wildlife in farmed and natural environments [1]. Human TSEs, such as Creutzfeldt-Jakob disease (CJD), variant CJD, Gerstmann Sträusler-Scheinker disease, and kuru are typically identified via observed clinical signs and post-mortem analyses [2]. Animal TSEs have largely been classified in the same manner and include transmissible mink encephalopathy, scrapie of sheep and goats, chronic wasting disease in free-ranging and captive species of Cervidae, feline spongiform encephalopathy, and bovine...
spongiform encephalopathy [2]. Importantly, the most profound unifying feature of these diseases is the accumulation of an infectious protease-resistant isoform (PrPSc) of the host-encoded cellular prion protein (PrPC) within tissues of the central nervous system [1].

Notably, the prion protein gene (PRNP) is present in all vertebrate species [3], and the striking degree of amino acid conservation observed across a wide variety of highly divergent taxa suggests an important functional role for PrPC [4]. Nevertheless, this evolutionary observation has been paradoxically shrouded by several knockout studies that failed to elucidate one or more overt physiological roles for PrPC [5–7]. The implication that PrPC deficient cattle may be considered safe mainstays for enhancing future agricultural products [8] is also in direct conflict with a previous investigation that provided unequivocal population and phylogenetic evidence for intense purifying selection constraining the long-term evolution of bovine PRNP [9]. Strong levels of purifying selection similar to that observed for bovine PRNP are normally only detected for genes encoding functionally important endogenous host proteins such as histones or ubiquitin [10, 11]. In the absence of function, what might explain such intense purifying selection?

The primary goal of this study was to determine whether strong purifying selection is acting directly on bovine PRNP [9], or if the selective signal actually radiates from a proximal bovine protein coding gene. Notably, the two proximal genes flanking bovine PRNP on BTA13 are ZMYND11 (~202 kb upstream) and PRND (~31 kb downstream). ZMYND11 encodes a transcriptional regulator protein that binds to adenovirus E1A proteins [12] while PRND encodes the doppel protein (Dpl), an evolutionarily related paralogue of PRNP believed to be involved in sperm maturation and capacitation [13]. Given the physical proximity between PRNP and PRND [14], as well as phylogenetic evidence that the two genes have been evolutionarily co-selected [15], we tested the hypothesis that strong selection within bovine PRND may be explanatory for selective signals previously detected in bovine PRNP [9]. Collectively, our analyses provide statistical support for intense purifying selection operating on bovine PRNP, with bovine PRND variation exhibiting no evidence for deviation from a strictly neutral model.

**Materials and Methods**

**DNA panels**

To comprehensively evaluate nucleotide sequence variation within bovine PRNP and PRND, we compiled and utilized data derived from 228 DNA samples previously employed to investigate bovine PRNP [16–18], including representatives of Bos taurus taurus, Bos taurus indicus, and their hybrids (composites). For PRND analysis, we used these same bovine samples, which included DNA extracted from 39 Holstein steers [17] and 189 commercially available spermatozoa samples of unrelated sires from the following 41 cattle breeds: Angus (4), Beefmaster (4), Belgian Blue (4), Blonde D’Aquitaine (5), Braford (4), Brahman (28), Brahmosin (2), Brangus (12), Braunvieh (5), Brown Swiss (4), Charolais (5), Chianina-Chianguis (5), Corriente (1), Gelbvieh (4), Gir (12), Guzerat (1), Hereford (3), Holstein (4), Limousin (3), Maine Anjou (4), Murray Gray (2), Nelore (8), Normande (1), Piedmontese (2), Red Angus (4), Red Brangus (1), Red Poll (1), Romagnola (2), Salers (3), Santa Gertrudis (9), Scottish Highland (1), Senepol (2), Shorthorn (19), Simbrah (3), Simmental (8), Tabapua (1), Tarentaise (1), Texas Longhorn (4), Three-way-cross (2), and White Park (1) [16, 18].

**PRND sequencing and multiple sequence alignments**

Polymerase chain reaction primers amplifying a 991 bp product encompassing the entire PRND coding region were designed from the Genbank reference sequence DQ205538 (5’-AGATCACTATCCTGAAATGGTG-3’, 5’-TTTAGGTAAGCCTGGAG-3’). Each 25-μL PCR
reaction contained 50 ng of genomic DNA, 1x PCR buffer with 1.5 mM MgCl₂, 1 mM each dNTP, 0.8 μM of each primer, and 1.5 units of Taq polymerase. Amplification conditions were as follows: 94°C for 1 min; 35 cycles of 94°C for 30s, 56°C for 30s, 72°C for 30s; 72°C for 2 min; 4°C hold. PCR products were visualized and verified on 2% NuSieve gels (Cambrex, Rockland, ME) and subsequently treated with ExoSAP-IT (GE Healthcare, Piscataway, NJ) for purification. Purified PRND PCR products were directly sequenced using the amplification primers and a pair of internal primers (5’-TGCCAAGTACCTCCAG-3’, 5’-TTTCCTTGATGACATTGG-3’) in conjunction with standard dye terminator cycle sequencing technology. Individual PRND contig sequences were assembled for each sample using Lasergene 6 (DNASTAR, Inc., Madison, WI). Thereafter, PRND sequences were aligned using ClustalX [19] and submitted to GenBank (accession numbers JF808218-JF808446). Likewise, bovine PRNP exon 3 sequences [9, 16, 17] were also aligned using ClustalX [19] as previously described [9].

Haplotype inference and network analysis
For haplotypes that were not phase-resolved through a second round of PCR, cloning, and bidirectional sequencing [9], we assembled unphased diploid genotypes for nucleotide sequence variation observed within the coding regions of bovine PRNP and PRND, including both single nucleotide polymorphisms (SNPs) and insertion-deletion mutations (indels). Bovine PRNP and PRND haplotype reconstructions were performed with PHASE 2.1 [20, 21] using all intragenic polymorphisms, all cattle (n = 228), and the—X10 option as previously described [22]. Haplotype phases previously established for 112 of the PRNP samples [9] were designated as phase-known for haplotype reconstruction.

Median joining haplotype networks for bovine PRNP and PRND were constructed using the program Network 4.5.1.0 (Fluxus Technology Ltd) in conjunction with the suggested character weights of 10 for SNPs and 20 for indels. Network branch angles were adjusted to ensure clarity without modifying branch lengths.

Sequence analysis
Phased PRNP and PRND sequences were used in conjunction with the software program DnaSP 5.1 [23] to estimate the number of potentially synonymous and non-synonymous nucleotide sites, the number of synonymous and non-synonymous polymorphisms, and the number of synonymous (dₐ) and non-synonymous (dₛ) substitutions per site with Jukes-Cantor correction [24, 25]. Of the 456 phased resolved PRNP sequences evaluated, 19 had alignment gaps in the octapeptide repeat region and were excluded from the synonymous and non-synonymous analysis.

Tests of selection
To evaluate potential deviations from a strictly neutral model of molecular evolution, we used the Z-test implemented in MEGA 4 [26] to evaluate the null hypothesis that dₙ = dₛ (strict neutrality; two-tailed test) and the research hypothesis dₙ < dₛ (one-tailed test) using pair-wise deletion of alignment gaps with Jukes-Cantor correction. For pairwise tests of selection, we estimated the variance of (dₙ - dₛ) via bootstrap analysis with 1000 replicates. Frequency distribution tests, including Tajima’s D [27] and Fu and Li’s Tests (D* and F*) [28], were performed in DnaSP v5.1 [23] using all PRNP and PRND coding region polymorphisms (excluding gaps). Significance was assessed for each test by estimating confidence intervals through coalescent simulations using the observed number of segregating sites with 5,000 replicates [23]. All tests were conducted both with and without the 19 PRNP haplotypes possessing alignment gaps. The program GARD (Genetic Algorithm for Recombination Detection) was
used to detect the presence of recombination in the \textit{PRND} and \textit{PRNP} sequences [29]. No overt evidence of recombination was detected. The potential for episodic and pervasive selective pressures was estimated for the unique haplotypes using the tree-based analysis programs MEME (Mixed-Effects Model of Evolution [30]), FUBAR (Fast, Unconstrained Bayesian AppRoximation [31]), and BS-REL (Branch-site Random-effects Likelihood [30]), which are part of the HyPhy software suite [32], as executed in the Datamonkey webserver [33]. Chi-square with Yates correction was used to assess the overall magnitude of differences in the distributions of polymorphisms at synonymous and non-synonymous sites with respect to \textit{PRNP} and \textit{PRND} (http://www.quantpsy.org/chisq/chisq.htm).

\section*{Results}

\subsection*{General nucleotide data}

To facilitate a detailed comparative analysis between bovine \textit{PRNP} and \textit{PRND} variation, we computed the number of potentially synonymous and non-synonymous nucleotide sites [24, 25], the total number of synonymous and non-synonymous SNPs, and the number of synonymous and non-synonymous substitutions per site (Table 1). The coding sequence of bovine Doppel is 537 bp in length, and 6 SNPs were identified among the 228 samples at positions 141 (A/G), 149(A/G), 172(A/G), 285(C/T), 395(A/G), and 528(A/T). The three SNPs detected at sites 141, 285, and 528 were predicted to encode synonymous substitutions, while SNPs at positions 149, 172, and 395 were predicted to encode amino acid replacements (R50H, A58T, Q132R, respectively; IUB/IUPAC Amino Acid Codes). All 6 \textit{PRND} polymorphisms were detected among samples representing \textit{B. t. taurus} and the composite cattle, but only 3 were predicted in samples representing \textit{B. t. indicus} cattle (141, 172, 395). Collectively, 30 synonymous SNPs and one non-synonymous SNP (S154N; \textit{B. t. indicus} and composite cattle) were predicted in the coding sequences of bovine \textit{PRNP}, as previously described [9].

\subsection*{Haplotype data}

The 31 polymorphic nucleotide sites in \textit{PRNP} yielded 31 distinct haplotypes (S1 Table), with corresponding frequency distributions among \textit{B. t. taurus}, \textit{B. t. indicus}, and composite cattle.

\begin{table}[h]
\centering
\caption{Nucleotide data for \textit{PRND} and \textit{PRNP} genes.}
\label{table1}
\begin{tabular}{llllllll}
\hline
 & #alleles & #syn sites a & # syn mut b & # non-syn sites c & # non-syn mut d & $d_s$ e & $d_n$ f & $d_n/d_s$ \\
\hline
\textbf{PRND} & & & & & & & & \\
\textit{B. t. taurus} & 278 & 119.06 & 3 & 414.94 & 3 & 0.0036 & 0.0013 & 0.3705 \\
\textit{B. t. indicus} & 100 & 119.36 & 1 & 414.64 & 2 & 0.0028 & 0.0024 & 0.8514 \\
Composite & 78 & 119.12 & 3 & 414.88 & 3 & 0.0043 & 0.0017 & 0.3869 \\
Total & 456 & 119.14 & 3 & 414.86 & 3 & 0.0037 & 0.0018 & 0.4796 \\
\hline
\textbf{PRNP} & & & & & & & & \\
\textit{B. t. taurus} & 264 & 186.33 & 23 & 605.67 & 0 & 0.0064 & 0.0000 & 0.0000 \\
\textit{B. t. indicus} & 98 & 186.33 & 10 & 605.67 & 1 & 0.0102 & 0.0003 & 0.0255 \\
Composite & 75 & 186.33 & 10 & 605.67 & 1 & 0.0070 & 0.0001 & 0.0128 \\
Total & 437 & 186.33 & 30 & 605.67 & 1 & 0.0076 & 0.0001 & 0.0105 \\
\hline
\end{tabular}
\begin{flushleft}
\textsuperscript{a} number of potentially synonymous sites \\
\textsuperscript{b} number of synonymous mutations observed \\
\textsuperscript{c} number of potentially non-synonymous sites \\
\textsuperscript{d} number of non-synonymous mutations observed \\
\textsuperscript{e} synonymous substitutions per site \\
\textsuperscript{f} non-synonymous substitutions per site
\end{flushleft}
\end{table}
that were similar to a previous PRNP analysis [16, 18]. The 6 polymorphic nucleotide sites in PRND produced 9 individual haplotypes (Table 2). PRND haplotypes #1 and #2 accounted for the majority of all haplotypes predicted in B. t. taurus (85%), B. t. indicus (63%), and composite (79%) cattle. Notably, four PRND haplotypes were exclusive to B. t. taurus in our samples, which resulted from 3 putative SNPs that were not detected among B. t. indicus cattle. Interestingly, haplotypes #5, #6, and #9 were identified primarily in B. t. indicus (37%) compared to B. t. taurus (1%; See Table 1). Median joining haplotype networks constructed as putative representations of bovine PRNP and PRND evolution (Fig 1) provide evidence for only a few major haplotypes as well as haplotype sharing across the three investigated bovine lineages (B. t. taurus, B. t. indicus, and composites). Moreover, specialized beef and dairy breeds could not be differentiated based on PRNP or PRND haplotypes, which is concordant with a recent study on bovine Toll-like receptor evolution [34].

Tests of selection

The observed ratio of synonymous to non-synonymous polymorphisms predicted for bovine PRNP was highly skewed (30:1, respectively), whereas PRND exhibited no skewness (3:3, respectively). To further assess and compare the potential for functional and/or selective constraint(s) acting on bovine PRNP and PRND, we computed Tajima’s D, Fu and Li’s D’, and Fu and Li’s F’ (Table 3). All frequency distribution tests indicated that variation within bovine PRNP does not adhere to a strictly neutral model, which is concordant with a previous study demonstrating that bovine PRNP is subject to strong purifying selection [16]. In contrast, frequency distribution tests carried out for PRND revealed no evidence for departure from a strictly neutral model (Table 3).

The Z-test was conducted to determine if the rate of change between synonymous sites and non-synonymous sites was strictly neutral (d_N = d_S) or directional (d_N < d_S) for PRND and PRNP (Table 3). Variation within bovine PRND did not deviate to a strictly neutral model, which is concordant with a previous study demonstrating that bovine PRNP is subject to strong purifying selection [16]. In contrast, frequency distribution tests carried out for PRND revealed no evidence for departure from a strictly neutral model (Table 3).

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Fig 1. Median joining haplotype networks for PRNP and PRND. Median joining haplotype networks were constructed for bovine PRNP and PRND using character weights of 10 for SNPs and 20 for indels. Network branch angles were adjusted to ensure clarity without modifying branch lengths.

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PRNP and PRND revealed a significant difference between the two genes \((P = 0.005)\), whereas the rate of nonsynonymous change was not found to differ \((P = 0.413)\). The difference in synonymous changes, but not nonsynonymous changes, further supports the supposition that bovine PRNP is under purifying selection, with selective signals that cannot be attributed to PRND.

To further clarify inferences drawn from our initial tests of selection, we also sought to investigate the potential for episodic selection within bovine PRND and PRNP. Analyses performed using the programs MEME and BS-REL [30] failed to detect evidence for episodic selection within PRND and PRNP (S1 File). Moreover, using the program FUBAR [31], we observed site-specific evidence for pervasive purifying selection within both PRND \((n = 3 \text{ sites/codons})\) and PRNP \((n = 9 \text{ sites/codons})\); whereas evidence for pervasive diversifying selection was only detected in PRND \((n = 2 \text{ sites/codons}; \text{S1 File})\). Collectively, these analyses further support the conclusion that nearly all nucleotide sites within the coding sequence of bovine PRND adhere to a strictly neutral model of molecular evolution, whereas the coding sequence of bovine PRNP is subject to intense purifying selection, thereby suggesting a potentially important role for bovine PrP.

**Discussion**

Herein, we have demonstrated that strong purifying selection on bovine PRNP cannot be attributed to selective pressures that are acting on a neighboring coding region (i.e., PRND), as bovine PRND variation does not depart from a strictly neutral model of molecular evolution. We also show that B. t. taurus and B. t. indicus share the major haplotypes for both genes, thereby suggesting that the majority of the polymorphisms, and the differences in selective constraints between PRNP and PRND, likely occurred before taurine and indicine divergence.

Notably, we cannot dispute the fact that prion knockouts analyzed to date lack gross evidence of deleterious effects [36]. However, we do dispute the conclusion that a lack of deleterious effects suggests that PRNP is simply dispensable [5], as biological dispensability, the manifestation of disease, and evolutionary rate have a significant relationship [37–39]. Interestingly, prior analyses of essential genes, Mendelian disease genes, and complex disease genes, as compared to non-essential and non-disease genes, demonstrate evidence of strong purifying selection [38, 39]. Relevant to this study, a protein under intense purifying selection (i.e., bovine PRNP), with even a small but measureable fitness effect, may be essential for the functional viability of certain cells and their corresponding tissues [37, 40–42]. For this reason, it

Table 3. Tests of selection for PRND and PRNP genes.

|         | Fu and Li D* | Fu and Li F* | Tajima’s D | Z-test \(d_n = d_s\) | Z-test \(d_n < d_s\) |
|---------|--------------|--------------|------------|-----------------------|-----------------------|
| **PRND**|              |              |            |                       |                       |
| B. t. taurus | 1.04         | 0.84         | 0.07       | -0.871                | 0.917                |
| B. t. indicus | 0.83         | 1.48         | 2.21       | -0.117                | 0.116                |
| Composite | 1.14         | 0.84         | -0.16      | -0.899                | 0.938                |
| All Cattle | 1.00         | 1.01         | 0.55       | -0.800                | 0.824                |
| **PRNP** |              |              |            |                       |                       |
| B. t. taurus | -4.89 a      | -4.53 a      | -2.00 a    | -2.374 a              | 2.435 a              |
| B. t. indicus | 0.67         | 0.32         | -0.47      | -2.656 a              | 2.756 a              |
| Composite | 0.04         | -0.54        | -1.39      | -2.341 a              | 2.416 a              |
| All Cattle | -3.64 a      | -3.55 a      | -1.95 a    | -2.753 a              | 2.797 a              |

\(a\) Statistically significant \((P < 0.05)\)

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PRNP and PRND revealed a significant difference between the two genes \((P = 0.005)\), whereas the rate of nonsynonymous change was not found to differ \((P = 0.413)\). The difference in synonymous changes, but not nonsynonymous changes, further supports the supposition that bovine PRNP is under purifying selection, with selective signals that cannot be attributed to PRND.

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should be noted that fitness effects may simply not have manifested in the controlled environmental conditions utilized for prion knockout cattle, or perhaps they were not recognizable through the limited assessment of these animals [5]. Examples exist in the literature of gene knock out effects observed to act only in a gender specific manner or in an age-dependent manner [43–46]. PRNP knockout mice have in fact been shown to have numerous subtle phenotypic differences [36]. Other than resistance to TSE, the first documented phenotypic change in PrP knockout mice was disturbances in sleep and altered circadian rhythms, as compared to wild-type mice, when housed in constant darkness [47, 48]. The knock out cattle used as evidence to suggest that PRNP is a dispensable gene were all castrated males studied as mature adults with no physiological or immunological stress placed on the animal(s). In general, given many accounts of gender-specific and age-specific effects observed for gene knockouts, as well as specific examples for PRNP reviewed by Steele and colleagues [36], it is possible that no gross changes were observed for castrated males under the carefully controlled environments (i.e., feeding and housing) used for the knock out cattle studies. These studies lack the exhaustive assessment of all developmental and physiological conditions necessary to assert any claims as to the biological dispensability of PRNP.

Fitness effect(s) or dispensability is not the only contribution to evolutionary rate. The number of protein-protein interactions has a strong correlation to both evolutionary rate and fitness [37]. Evidence that 30 different proteins likely interact with PrP [49], could at least, in part, explain the high degree of conservation commonly observed for the mammalian PRNP gene. Putative functional roles for PrP are likewise multifaceted with numerous cellular pathways influenced by PrP [50–52]. PrP has been shown to have a role in epithelial to mesenchymal transition [53], and several studies report protective roles for PrP, including protection against oxidative stress [54, 55]. PrP may also play a direct role in Alzheimer’s disease, another neurodegenerative disorder [54, 56]. High levels of PrP expression are found in placenta, indicating a potential role in reproduction [57]; this role ties closely with evidence that Dpl and a third member of the prion gene family, Shadoo (Sho), can both interact with PrP, as well as function in place of PrP in aspects of reproduction [58]. Consistent with this is the observation that in the absence of PrP, aberrant expression of Dpl in the CNS results in mice that develop ataxia due to apoptosis of cerebellar cells [59], suggesting that Dpl is interacting with ligands that would normally interact with PrP. Expression of PrP in these tissues is sufficient to negate this effect [60]. Sho exhibits PrP-like neuroprotective properties with regard to Dpl-induced neurotoxicity in the CNS [49], indicating that Sho is also likely to be capable of interacting with the same ligands. A clear overlap of interactions exists between the various members of the prion gene family, which may be one plausible explanation for the lack of a discernable phenotype in PRNP knockout animals.

Supporting Information

S1 File. BSREL and FUBAR analysis output.
(DOCX)

S1 Table. PRNP haplotype information.
(XLSX)

Acknowledgments

Mention of trade names or commercial products in this article is solely for the purpose of providing specific information and does not imply recommendations or endorsement by the US Department of Agriculture. USDA is an equal opportunity provider and employer.
Author Contributions
Conceived and designed the experiments: EMN BWB CMS. Performed the experiments: AMO. Analyzed the data: BWB CMS. Contributed reagents/materials/analysis tools: BWB CMS EMN. Wrote the paper: EMN BWB CMS AMO.

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