Scrapie in Swine: a Diagnostic Challenge

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Abstract
A naturally occurring prion disease has not been recognized in swine, but the agent of bovine spongiform encephalopathy does transmit to swine by experimental routes. Swine are thought to have a robust species barrier when exposed to the naturally occurring prion diseases of other species, but the susceptibility of swine to the agent of sheep scrapie has not been thoroughly tested. We conducted this experiment to test the susceptibility of swine to U.S. scrapie isolates by intracranial and oral inoculation. Scrapie inoculum was a pooled 10% (w/v) homogenate derived from the brains of clinically ill sheep from the 4th passage of a serial passage study of the U.S scrapie agent (No. 13–7) through susceptible sheep (homozygous ARQ at prion protein residues 136, 154, and 171, respectively). Pigs were inoculated intracranially (n=19) with a single 0.75 mL dose or orally (n=24) with 15 mL repeated on 4 consecutive days. Necropsies were done on a subset of animals at approximately six months post inoculation (PI): the time the pigs were expected to reach market weight. Remaining pigs were maintained and monitored for clinical signs of transmissible spongiform encephalopathies (TSE) until study termination at 80 months PI or when removed due to intercurrent disease (primarily lameness). Brain samples were examined by immunohistochemistry (IHC), western blot (WB), enzyme immunoassay (EIA), and for a subset of pigs in each inoculation group, bioassay in mice expressing porcine prion protein. At six-months PI, no evidence of scrapie infection was noted by any diagnostic method. However, at 51 months of incubation or greater, 5 animals were positive by one or more methods: IHC (n=4), WB (n=3), or EIA (n=4). Furthermore, positive bioassay results were obtained from all inoculated groups (oral and intracranial; market weight and end of study) suggesting that swine are potential hosts for the agent of scrapie.

Keywords
prion, scrapie, swine, transmissible spongiform encephalopathy (6)

Disciplines
Large or Food Animal and Equine Medicine | Veterinary Infectious Diseases | Veterinary Pathology and Pathobiology

Comments

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Scrapie in Swine: a Diagnostic Challenge

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A naturally occurring prion disease has not been recognized in swine, but the agent of bovine spongiform encephalopathy does transmit to swine by experimental routes. Swine are thought to have a robust species barrier when exposed to the naturally occurring prion diseases of other species, but the susceptibility of swine to the agent of sheep scrapie has not been thoroughly tested. We conducted this experiment to test the susceptibility of swine to U.S. scrapie isolates by intracranial and oral inoculation. Scrapie inoculum was a pooled 10% (w/v) homogenate derived from the brains of clinically ill sheep from the 4th passage of a serial passage study of the U.S scrapie agent (No. 13–7) through susceptible sheep (homozygous ARQ at prion protein residues 136, 154, and 171, respectively). Pigs were inoculated intracranially (n=19) with a single 0.75 mL dose or orally (n=24) with 15 mL repeated on 4 consecutive days. Necropsies were done on a subset of animals at approximately six months post inoculation (PI): the time the pigs were expected to reach market weight. Remaining pigs were maintained and monitored for clinical signs of transmissible spongiform encephalopathies (TSE) until study termination at 80 months PI or when removed due to intercurrent disease (primarily lameness). Brain samples were examined by immunohistochemistry (IHC), western blot (WB), enzyme immunoassay (EIA), and for a subset of pigs in each inoculation group, bioassay in mice expressing porcine prion protein. At six-months PI, no evidence of scrapie infection was noted by any diagnostic method. However, at 51 months of incubation or greater, 5 animals were positive by one or more methods: IHC (n=4), WB (n=3), or EIA (n=4). Furthermore, positive bioassay results were obtained from all inoculated groups (oral and intracranial; market weight and end of study) suggesting that swine are potential hosts for the agent of scrapie.

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initial passage is usually not efficient between species, but subsequent passages can lead to stabilization of disease in a new species\(^7\). The major influence on species barrier is amino acid sequence differences between the donor and recipient hosts\(^6,8\) and the effect on prion protein structure and folding\(^9\).

The susceptibility and incubation periods of prion disease in an individual are highly dependent on the amino acid sequence of the prion protein in mice\(^10\), sheep\(^11–13\), cervids\(^14,15\), and humans\(^16\). However, compared to sheep, cervids, and humans, the porcine prion protein (PrP) is highly homogenous\(^17,18\) with only four known single nucleotide polymorphisms (G11A, G615C, G684A, T726G) one of which (G615C) that results in a serine to asparagine substitution at codon 4 that is not part of the mature prion protein\(^19\). As compared to cattle, pigs have 5 unique amino acids that appear in the prion protein sequence of the globular domain and 4 of these appear in helix 3\(^20\). While epidemiologic evidence would suggest pigs are not susceptible to prion disease, in vitro assays demonstrate that recombinant porcine PrP converts to amyloid fibrils faster than human or bovine sequences\(^21\).

Several experimental studies have been conducted to test the susceptibility of pigs to prion diseases. The agent of BSE transmitted to pigs after multiple-route (concurrent intracranial, intravenous, and intraperitoneal) parenteral inoculation with an incubation period of 17–38 months and resulted in severe neuropil vacuolation and abnormal prion protein (PrP\(_{Sc}\)) accumulation in the brain\(^3\). Further, infectivity was demonstrated in brain, spinal cord, stomach, small intestine, and pancreas by bioassay in C57BL/6J mice\(^22\). However, attempts to transmit BSE to pigs after oral dosing with up to 1.2 kg of brain material from infected cattle was unsuccessful\(^23\). Studies utilizing transgenic mice expressing the porcine prion protein suggest that initial low dose exposure to BSE prions may result in a subclinical infection that results in a high attack rate and shortened incubation periods on second passage\(^20\).

Piglets exposed orally to brain material from scrapie-infected sheep from the UK lacked any evidence of disease and bioassay of tissues from these pigs failed to reveal infectivity in tissues\(^23\). However, mice expressing porcine PrP are susceptible to atypical (Nor-98) scrapie with a low attack rate that adapt to 100% attack rate with a rapid incubation on subsequent passages\(^24\), similar to the adaptation that was observed in experiments that were conducted with BSE\(^20,24\). Interestingly, sheep passed BSE transmits more efficiently to pigs than BSE from cattle and was associated with PrP\(_{Sc}\) in variety of peripheral tissues\(^25\).

We conducted a study to determine if swine were susceptible to an isolate of the US scrapie agent by either the intracranial or oral route. Initially, we used transgenic mice expressing porcine prion protein (PoPrP-Tg002) to test their susceptibility to the agent of sheep scrapie. No mice in this study developed clinical signs before the end of the experiment (700 days), but 5/30 tested positive by enzyme immunoassay (EIA; BSE-Scrapie Antigen Test Kit, IDEXX, Westbrook, ME). For the swine study, weaned piglets were inoculated with a 10% (w/v) homogenate made from the brains of clinically ill sheep from the 4th passage of a serial passage study of the U.S scrapie agent (No. 13–7) through sheep homozygous ARQ at prion protein residues 136, 154, and 171, respectively\(^26\). Intracranial inoculations\(^27\) (n=19) were with a single 0.75 mL dose of homogenate. Oral inoculations (n=24) were 15 mL doses repeated on 4 consecutive days. Necropsies were done on a subset of animals at approximately six months post inoculation (PI): the time the pigs were expected to reach market weight. Remaining pigs were maintained and monitored for clinical signs of transmissible spongiform encephalopathies (TSE) until study termination at 80 months PI or when removed due to intercurrent disease. Brain samples were examined by immunohistochemistry (IHC)\(^28\), western blot (WB)\(^12\), and enzyme immunoassay (EIA)\(^29\), and for a subset of pigs in each inoculation group, bioassay in mice expressing porcine PrP.

All pigs examined at market weight tested negative by traditional diagnostic methods. However, 5/10 aged pigs that were necropsied at 51 months of age or later tested positive using immunohistochemistry (IHC; n=4), western blot (WB; n=3), and/or EIA (n=4). In all hematoxylin and eosin stained slides examined by light microscopy, there was no evidence of spongiform encephalopathy beyond that present in negative control pigs. Examination of slides stained by an immunohistochemical method with monoclonal antibody L42 that targets to amino acids 145–163 of the ovine prion protein sequence\(^30\) demonstrated immunoreactivity that was largely confined within neurons (Fig. 1) and was present in hippocampus, cerebellum, thalamus, brainstem, and spinal cord. Findings from HE and IHC were similar to scrapie in cattle where spongiform change was not noted and IHC positives had prominent intraneuronal labeling\(^31\). In pigs that were WB positive, the molecular profile was distinct from the original scrapie inoculum in that the nonglycosylated band migrated lower than sheep scrapie and that the profile had a most prominent monoglycosylated rather than diglycosylated band (Fig. 2).

One pig from each inoculation group was bioassayed in PoPrP-Tg002 mice. Bioassay results were determined by EIA. Positive results were obtained from the market weight pigs that were orally (5/29) or intracranially (2/27) inoculated. The sample that was selected from the orally inoculated aged pig (incubation period=50 months) also had a low number of positive results (2/24) in mouse bioassay. Interestingly, the aged pigs from the intracranially inoculated pigs that were IHC, WB, and/or EIA positive that were selected for bioassay yielded 0/29 and 0/30 positive samples, which
was unexpected. This interesting result contrasted those of BSE samples where second passage results in a 100% attack rate with a relatively short incubation period\textsuperscript{20,24}. Of great interest are the mice that tested ELISA positive after bioassay of brain material from an orally inoculated pig that was examined at market weight. These mice were collected at the end of the mouse study that was scheduled to last for 700 days post-inoculation. The positive mice had low EIA optical density scores. No vacuolar change was noted on sections available for lesion profiling. A second passage from a positive mouse with a positive EIA was attempted in mice expressing porcine prion protein (PoPRP-Tg002) or mice expressing sheep prion protein (Tg338). The second passage in PoPRP-Tg002 mice is ongoing, but passage to Tg338 mice resulted in 16/20 positive mice. This work demonstrates that pigs are susceptible to the US scrapie agent by oral and intracranial inoculation. Aged animals inoculated intracranially were detectable by traditional diagnostic methods, however, orally

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**Fig. 1.** Immunoreactivity (red) in central nervous system tissues of pigs with scrapie is predominantly intraneuronal. High magnification of the brainstem at the level of the obex with intraneuronal immunoreactivity after staining with an automated protocol using monoclonal antibody L42.

**Fig. 2.** Western blot analysis demonstrating unique PrP\textsuperscript{Sc} profile in pigs with scrapie. The positive sample from a pig inoculated with the agent of scrapie has a prominent monoglycosylated band and lower migration relative to the sheep scrapie inoculum. Blot developed with monoclonal antibody L42.
exposed aged or market weight animals, while positive by bioassay were not detected by any standard diagnostic technique. Further, no animals demonstrated clinical signs of disease or had spongiform change in their brains.

Pigs were exposed to the agent of BSE in contaminated meat and bone meal in many European countries, but no naturally occurring cases have been described and surveys of meat and bone meal fed pigs failed to demonstrate any evidence of TSE in pigs. Pigs challenged orally with BSE failed to develop evidence of infection despite observation for up to seven years. These previously published studies along with results of the current study suggest that swine are capable of harboring a prion disease, although epidemiologic evidence is not in support of this occurring under production conditions.

We have demonstrated that swine are susceptible to the agent of sheep scrapie. While no pigs examined at market weight were positive by traditional diagnostic methods, there were 5 aged pigs with positive IHC, WB, and/or ELISA from brain tissue. These pigs did not have definitive clinical illness or spongiform change during the course of this experiment. Positive bioassay results from the brains of market weight pigs suggest that even orally inoculated pigs do harbor a low level of infectivity. If scrapie were to occur in pigs, it would represent a significant diagnostic challenge because of long incubation periods, failure of pigs to develop clinical signs, and the inability of traditional diagnostic methods to detect infected animals early in the course of disease.

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