In most forms of prion disease, infectivity is present primarily in the central nervous system or immune system organs such as spleen and lymph node. However, a transgenic mouse model of prion disease has demonstrated that prion infectivity can also be present as amyloid deposits in heart tissue. Deposition of infectious prions as amyloid in human heart tissue would be a significant public health concern. Although abnormal disease-associated prion protein (PrPSc) has not been detected in heart tissue from several amyloid heart disease patients, it has been observed in the heart tissue of a patient with sporadic Creutzfeldt-Jakob Disease (sCJD), the most common form of human prion disease. In order to determine whether prion infectivity can be found in heart tissue, we have inoculated formaldehyde fixed brain and heart tissue from two sCJD patients, as well as prion protein positive fixed heart tissue from two amyloid heart disease patients, into transgenic mice overexpressing the human prion protein. Although the sCJD brain samples led to clinical or subclinical prion infection and deposition of PrPSc in the brain, none of the inoculated heart samples resulted in disease or the accumulation of PrPSc. Thus, our results suggest that prion infectivity is not likely present in cardiac tissue from sCJD or amyloid heart disease patients.

Transmissible spongiform encephalopathies (TSEs) or prion diseases are a group of rare, fatal, and infectious neurodegenerative diseases that include scrapie in sheep, bovine spongiform encephalopathy in cattle (BSE), and sporadic Creutzfeldt-Jakob disease (sCJD) in humans. The conversion of the soluble and normally protease-sensitive host prion protein (PrPC) into a highly aggregated and protease-resistant form, PrPSc, is a central event in prion pathogenesis and PrPSc is the primary component of the infectious prion. In humans, prions may be acquired by ingestion or inoculation of prion-contaminated materials, by exposure to prion-contaminated surgical instruments or via transplantation of tissues that can harbor prion infectivity. In the latter instance, cornea, dura mater, and cadaver-derived human growth hormone have transmitted sCJD (1), while variant CJD (vCJD), which is linked to exposure to BSE (2), has been transmitted via blood (3, 4) and blood products (5). Thus, from a public health perspective, it is critical to understand which tissues harbor infectivity for each type of prion disease in order to prevent accidental transmission of prions via inoculation, transplantation, or prion-contaminated medical instruments.

PrPSc is expressed in almost every tissue in the body and can be found at high levels in several organs such as the brain, heart, and spleen (6, 7). Although theoretically PrPSc could accumulate in any tissue that expresses PrPC, its distribution can vary. For example, PrPSc in human vCJD is found not only in the brain and spinal cord but also in the spleen, tonsils, and peripheral lymph nodes (8, 9). In the most common form of human prion disease, sCJD, PrPSc is detected in central nervous system (CNS) tissues (9–11) and occasionally in other tissues such as skeletal muscle (12–14). Since the distribution of PrPSc varies depending upon the type of prion disease, the risk of transmission of prion infectivity from peripheral (i.e., non-CNS) tissues has to be determined experimentally for each prion disease type.

Humans expressing PrP molecules without the glycosylphatidylinositol (GPI) cell membrane anchor accumulate abnormal PrP amyloid in the brain and develop a form of prion disease (15). After infection with mouse scrapie, transgenic mice expressing mouse prion protein without the GPI membrane anchor not only develop a prion disease due to accumulation of PrP amyloid in the brain (16) and other tissues (17) but also develop an infectious cardiac amyloidosis with high levels of PrPSc and infectivity in the heart (18). These data raised the possibility not only that PrPSc might be deposited in the heart of individuals with prion disease but also that abnormal forms of PrP could be an unrecognized component of amyloid heart disease in humans (19). Consistent with this possibility, PrPSc has been reported in the heart of one sCJD patient (20), in muscle tissue (including heart) of BSE-infected nonhuman primates (21), in a BSE-infected macaque with a prion-amyloid cardiomyopathy (22), and in several chronic wasting disease models (23, 24). In contrast, several studies have failed to detect PrPSc in either sCJD heart tissue (9, 12, 19) or human amyloid hearts (19), and heart tissue from several sCJD patients did not transmit disease to nonhuman primates (25).

However, prion infectivity can be present in a tissue even in the absence of detectable PrPSc. For example, in both wild-type mouse...
models of BSE (26) and transgenic mouse models of prion infection (27–29), significant infectivity can be detected in brain material that is negative for PrPSc by both immunoblotting and immunohistochemistry. Similarly, even though PrPSc is restricted primarily to tissues of the CNS in sCJD (9–11), transmission studies in nonhuman primates occasionally detected infectivity in PrPSc-negative tissues such as lung, liver, and kidney (25). The occasional detection of prion infectivity in non-CNS tissue in sCJD could be the consequence of a low titer of infectivity in the tissue and/or a barrier to prion infection between human and nonhuman primates, either of which could decrease the efficiency of prion transmission across species. Species barriers in prion diseases are defined by extremely long incubation times at first passage compared to incubation times at later passages (30, 31). They are strongly influenced by the sequence of host PrP C and can be overcome using transgenic mice expressing PrPSc with the same amino acid sequence as PrPSc (32).

In order to determine whether prion infectivity is present in human heart tissue, we have inoculated archival formaldehyde-fixed and paraffin-embedded brain and heart tissue from two sCJD patients, as well as PrP-positive heart samples from two amyloid heart disease patients, into transgenic mice overexpressing human prion protein. These mice have no significant species barrier to infection with the most common form of sCJD. Our results show that although both brain samples triggered a prion infection, neither clinical nor subclinical prion infection occurred in mice inoculated with any of the heart tissue samples. Although the samples we tested are limited, our data suggest that prion infectivity is not likely present in either sCJD heart or PrP-positive amyloid heart tissue.

MATERIALS AND METHODS

Antibodies. The anti-hamster PrP monoclonal antibody 3F4 has been described previously (33) and recognizes an epitope encompassing amino acid residues 109 to 112 present in both hamster and human PrP (34). The 3F4 antibody was made in-house at the Rocky Mountain Laboratories from the tissue culture supernatant of hybridoma cells. PV30 denotes an affinity-improved variant of the recombinant anti-PrP antibody P that recognizes a peptide epitope composed of PrP residues 96 to 105 (35). Regions of the parental antibody P sequence were randomized using an overlap PCR methodology employing an NNS doping strategy (36). Two independent mouse antibody-phyage display libraries were constructed. In the first, a total of six residues within the first and third light-chain-complementary-determining regions (CDR1 and CDR3, respectively) were targeted for mutagenesis. In the second, four residues within the heavy-chain CDR3 were randomized. The two resulting libraries contained an average of 5 × 10^6 variants. Phage library selection was carried out in solution against synthetic biotinylated bovine and human PrP (resids 90 to 145). Using this strategy, over 30 different variants of the parental antibody P with improved kinetic constants were recovered. Of this number, Fab clone PV30 demonstrated an 88-fold improvement in binding kinetics for human PrP 23–231 when measured against the parental antibody P using surface plasmon resonance (K_D = 0.18 ± 0.07 nM for PV30, versus 16.61 ± 0.1 nM for antibody P).

Tissue samples. For all tissues and homogenates studied, the superscript “FFPE” is used to denote material derived from formalin-fixed and paraffin-embedded tissue while the superscript “Froz” is used to denote material derived from frozen tissue.

Formalin-fixed, paraffin-embedded brain and heart tissue from patients diagnosed with CJD in 1979 (CJD1FFPE) and 1981 (CJD2FFPE), as well as heart tissue from two amyloid heart disease patients (AHD1FFPE and AHD2FFPE), were obtained from the Armed Forces Institute of Pathology. Additional FFPE amyloid diseased hearts were obtained from individual cases from repositories at the University of California, San Francisco, CA (Stephen DeArmond), the VA Medical Center, La Jolla, CA (Paul Wolf), the Amyloid Research Center at the Royal Free and University College Medical School, London, United Kingdom (Mark Pepsy), and the Department of Repository and Research Services, Armed Forces Institute of Pathology, Washington, DC (Sherman A. McCall). Normal heart (NHT1FFPE and NHT2FFPE, left ventricle) and normal brain (NB1FFPE, cerebral cortex) tissue were kindly provided by Gerald Bordin, Department of Pathology, Scripps Green Hospital, La Jolla, CA.

Prnp genotype. DNA was extracted from FFPE tissue using the Puregene DNA purification kit (Qiagen) according to the manufacturer’s instructions. Extracted DNA was amplified by PCR, and the product was digested with the restriction endonuclease NspI (New England Biolabs) as described earlier (37). The size of the NspI digestion product differs depending upon whether there is a codon for methionine or valine at amino acid position 129. Using DNA extracted from FFPE tissue sections, we were able to determine the Prnp genotype of AHD2FFPE as homozygous for methionine at codon 129 but have as yet been unable to clearly determine the genotypes of the samples CJD1FFPE, CJD2FFPE, and AHD1FFPE.

Mouse lines and prion isolates. The transgenic mouse lines overexpressing human PrPSc (Tg66) and hamster PrPSc (Tg7) have been described previously (38, 39). The control sCJD stock brain homogenate was derived from a frozen brain sample of sCJD MM type 1 from a patient homozygous for methionine at codon 129 in PrP (CJDfroz) and was kindly provided by Robert Rohwer (University of Maryland, Baltimore, MD). The CJDfroz stock had a titer in Tg66 mice after intracranial (i.c.) inoculation of 2 × 10^5.5 ID50/g of brain (ID50 is the dose at which 50% of the animals become infected). The stock frozen brain homogenate of the hamster-adapted scrapie strain 263K (263KFroz) had an i.c. titer of 2 × 10^9.3 ID50/g of brain.

Inoculation of FFPE scrapie-infected transgenic mouse brain. All animal experimental protocols were reviewed and approved by the Rocky Mountain Laboratories Animal Care and Use Committee (Protocol 2007-10). The Rocky Mountain Laboratories are fully accredited by the American Association for Laboratory Animal Care, and this study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health.

A frozen brain from Tg7 mice that had been i.c. inoculated with hamster 263Kfroz scrapie was fixed for 4 days in formalin and then paraffin embedded. The FFPE brain was stabbed with a 271/2 gauge needle four times for 10 s per stab in order to coat the needle with infected brain material. The needle was then used to inoculate Tg7 mice i.c., and the mice monitored for scrapie. After the needle stabs, the same block was heated to remove the tissue from the paraffin and the tissue inoculated in Pro-par clearant (Anatech, Ltd.) overnight at room temperature, followed by additional 2- and 1-h incubations in Pro-par the following day. The brain was rehydrated by rinsing in decreasing amounts of ethanol (100, 95, 80, and 50%) for 30 min for each rinse. The brain material was homogenized in phosphate-buffered saline (PBS) to give a final 20% (wt/vol) solution. The homogenate was vortexed, sonicated for 5 min, diluted 1:1 in 0.64 M sucrose, and centrifuged for 1 min at 1,000 rpm. The supernatant was removed and diluted 1:10 in PBS–2% fetal bovine serum (FBS), and then 50 μL was inoculated i.c. into Tg66 mice, which were subsequently monitored for disease.

Preparation of human FFPE brain tissue was similar to that for the Tg7 brain with the following minor modifications. The rehydrated tissue, which ranged in weight from 20 to 201 mg, was homogenized in PBS–0.16 M sucrose to give a final 10% (wt/vol) solution. This solution was vortexed, sonicated for 5 min and centrifuged for 1 min at 1,000 rpm. The supernatant was removed and diluted 1:10 in PBS–2% FBS and then 50 μL was inoculated i.c. into Tg66 mice. Formalin-fixed and paraffin-embedded human heart tissue was processed in the same way except that, due to the small amount of tissue available, the solution was not centrifuged prior to inoculation.
Human tissue immunohistochemistry. Normal and sCJD human brain and heart tissue sections were deparaffinized in Pro-Par clearing, followed by rehydration in decreasing amounts of ethanol. The slides were rinsed in distilled water and treated with formic acid (99%) for 10 min, followed by a 5-min rinse in double-distilled water. Slides were immersed in 1 M citrate buffer (pH 6.0), placed in a Biocare decloaking chamber, and treated at 120°C and at 20 lb/in² for 20 min. After cooling, slides were stained using an automated Ventana Nexus Stainer (Ventana Medical Systems). The primary antibody was the mouse monoclonal anti-PrP antibody 3F4, which was added to the slides undiluted for 30 min. The secondary antibody was biotinylated horse anti-mouse IgG (Vector Laboratories) diluted 1:250 in 1% normal goat serum (30 min), followed by a 1:3 dilution of peroxidase-conjugated streptavidin (Biogenex) for 30 min. The chromogen used for development was Ventana AEC (Ventana Medical Systems). The primary antibody was the mouse monoclonal anti-PrP antibody derived from hybridoma tissue culture supernatant. The secondary antibody was either a 1:3,000 dilution of horseradish peroxidase-conjugated sheep anti-mouse IgG (GE Healthcare) for ECL or Advanced ECL (GE Healthcare) according to the manufacturer’s instructions. Blots were developed using a 1:3,000 dilution of the 3F4 monoclonal antibody derived from hybridoma tissue culture supernatant. For preabsorbed antibody, PV30 (1 mg/ml) was mixed with 5 mg of a biotinylated PrP peptide derived from amino acids 89 to 112 that contains the PV30 epitope. The solution was incubated with mixing for 1 h at 4°C, followed by 3 h at 37°C with occasional stirring. After a 4,000 rpm spin to remove any precipitate, the supernatant was passed over an avidin column, and the pass-through was collected. This was concentrated to the original starting volume, and 50 µl was used per slide.

Slides were stained with Thioflavin S (ThioS) using a 1% solution of ThioS in distilled water. Slides were deparaffinized, washed in water, and then incubated in 88% formic acid for 30 min, followed by another wash water. Sections were treated with 25 µg of PK/ml for 10 min at 26°C, rinsed in water, and hydrolytically autoclaved in 10 mM Tris-HCl (pH 7.5) for 15 min. The primary antibody was either the anti-PrP antibody PV30 at a dilution of 1:400 in 5% dried milk for 1 h at room temperature (50 µl/slide) or preabsorbed PV30. For preabsorbed antibody, PV30 (1 mg/ml) was mixed with 5 mg of a biotinylated PrP peptide derived from amino acids 89 to 112 that contains the PV30 epitope. The solution was incubated with mixing for 1 h at 4°C, followed by 3 h at 37°C with occasional stirring. After a 4,000 rpm spin to remove any precipitate, the supernatant was passed over an avidin column, and the pass-through was collected. This was concentrated to the original starting volume, and 50 µl was used per slide.

Slides were stained with Thioflavin S (ThioS) using a 1% solution of ThioS in distilled water. Slides were deparaffinized, washed in water, and then incubated in 88% formic acid for 30 min, followed by another wash water. Sections were treated with 25 µg of PK/ml for 10 min at 26°C, rinsed in water, and hydrolytically autoclaved in 10 mM Tris-HCl (pH 7.5) for 15 min. The primary antibody was either the anti-PrP antibody PV30 at a dilution of 1:400 in 5% dried milk for 1 h at room temperature (50 µl/slide) or preabsorbed PV30. For preabsorbed antibody, PV30 (1 mg/ml) was mixed with 5 mg of a biotinylated PrP peptide derived from amino acids 89 to 112 that contains the PV30 epitope. The solution was incubated with mixing for 1 h at 4°C, followed by 3 h at 37°C with occasional stirring. After a 4,000 rpm spin to remove any precipitate, the supernatant was passed over an avidin column, and the pass-through was collected. This was concentrated to the original starting volume, and 50 µl was used per slide.

Western blotting. Brains from Tg66 mice were homogenized in PBS to give a final 20% (wt/vol) solution. Tris-HCl (pH 8.3), sodium deoxycholate, and Triton X-100 were added to the homogenate to give final concentrations of 0.1 M, 1%, and 1%, respectively. PK was then added to a final concentration of 63.3 µg/ml, and the reaction was incubated for 30 min at 37°C. Protease activity was inhibited by the addition of 0.1 M phenylmethylsulfonyl fluoride to a final concentration of 0.01 M, a volume of 2× sample loading buffer (1 M Tris-HCl [pH 6.8], 10% glycerol, 6 mM EDTA, 20% sodium dodecyl sulfate [SDS], 0.04% bromophenol blue) equal to the sample volume was added, and the sample was boiled for 3 min. A total of 0.6- to 0.7-mg brain equivalents were loaded onto a Novex 16% Tris-glycine gel (Invitrogen). After electrophoresis samples were transferred to polyvinylidene difluoride membrane (Millipore). Membranes were developed using either enhanced chemiluminescence (ECL) or Advanced ECL (GE Healthcare) according to the manufacturer’s instructions. Blots were developed using a 1:3,000 dilution of the 3F4 monoclonal antibody derived from hybridoma tissue culture supernatant. The secondary antibody was either a 1:3,000 dilution of horseradish peroxidase-conjugated sheep anti-mouse IgG (GE Healthcare) for ECL or a dilution of 1:250,000 for Advanced ECL.

RESULTS

Lack of a significant species barrier to human prion infection in mice overexpressing human prion protein. To determine whether or not there was a species barrier to infection between Tg66 mice and human sCJD, mice were inoculated with a stock homogenate made from a frozen brain sample of a common form of CJD, sCJD MM type 1 (CJD85/91). Consistent with previous reports (38), Tg66 mice developed disease after ~169 days (Table 1). Clinically ill mice demonstrated spongiform change and both synaptic and perineuronal deposition of PrPSc in multiple areas, including the pons (Fig. 1A), thalamus, and hypothalamus (data not shown). There was no significant change in the incubation time of human CJD (CJD85/91) that was passed three times through Tg66 mice (Table 1). The fact that the disease incubation time of CJD (CJD85/91) did not significantly change over multiple passages in Tg66 mice indicated that, as expected, there was no significant species barrier between Tg66 mice and infection with the most common form of sCJD.

Detection of hamster scrapie infectivity in FFPE hamster brain tissue. The amount of FFPE human tissue available for inoculation was limited, and two techniques were tested to determine a relatively efficient way to recover enough tissue for inoculation into mice: (i) needle stab and (ii) removal and rehydration of tissue from the paraffin block. In order to determine the viability of the two techniques, the brain from a clinically positive Tg7 mouse infected with the hamster scrapie strain 263K (263KFPPE) was formalin fixed for 4 days and paraffin embedded. The FFPE brain was stabbed with a needle that was then used to inoculate Tg7 mice i.c. The FFPE brain was then removed from the block, rehydrated, and homogenized, and the resulting homogenate (263KFPPE) was inoculated i.c. into Tg7 mice. As shown in Table 2, slightly over half of the Tg7 mice inoculated with the 263KFPPE needle stab developed scrapie after ~100

## Table 1. Lack of a significant species barrier to sCJD MM type 1 prion infection in Tg66 mice

<table>
<thead>
<tr>
<th>Passage</th>
<th>Clinical/total</th>
<th>Mean dpi ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>17/17</td>
<td>168.7 ± 12.8</td>
</tr>
<tr>
<td>2</td>
<td>19/19</td>
<td>182.6 ± 21.6</td>
</tr>
<tr>
<td>3</td>
<td>7/7</td>
<td>171 ± 6.4</td>
</tr>
</tbody>
</table>

* Each passage represents Tg66 mice inoculated with 50 µl of a 10⁻³ dilution of brain homogenate. The inoculum at passage 1 was sCJD MM type 1 (CJD85/91).

† That is, the number of Tg66 mice with clinical disease over the total number of inoculated mice.

‡ That is, the mean disease incubation time in days postinfection (dpi) ± standard deviation (SD) in days.
days, almost double the 52-day incubation time of mice inoculated with a brain homogenate stock of 263K hamster scrapie (Table 2, 263KFroz). In contrast, all Tg7 mice inoculated with 263KFFPE homogenate developed scrapie after 70 days (Table 2).

A standard hamster 263K scrapie incubation time curve in Tg7 mice was used to roughly estimate the amount of infectivity recovered by needle stab at 1/10 dilution of the starting material was inoculated intracranially.

**TABLE 2** Infection of transgenic mice overexpressing hamster PrP<sup>Sc</sup> with frozen or FFPE 263K scrapie-infected hamster brain

<table>
<thead>
<tr>
<th>Sample</th>
<th>Brain sample&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Clinical/total&lt;sup&gt;b&lt;/sup&gt;</th>
<th>Mean dpi ± SD&lt;sup&gt;c&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td>263K&lt;sub&gt;Froz&lt;/sub&gt;</td>
<td>Frozen homogenate</td>
<td>7/7</td>
<td>52 ± 5</td>
</tr>
<tr>
<td>263K&lt;sub&gt;FFPE&lt;/sub&gt;</td>
<td>FFPE-stab</td>
<td>6/10</td>
<td>99 ± 10</td>
</tr>
<tr>
<td>263K&lt;sub&gt;FFPE&lt;/sub&gt;</td>
<td>FFPE-homogenate</td>
<td>9/9</td>
<td>70 ± 7</td>
</tr>
</tbody>
</table>

<sup>a</sup> Samples were inoculated into Tg7 mice overexpressing hamster PrP<sup>Sc</sup>. For both homogenate samples, 50 µl of a 10<sup>-2</sup> dilution of the starting material was inoculated intracranially.

<sup>b</sup> That is, the number of mice with clinical scrapie over the total number of inoculated mice.

<sup>c</sup> That is, the mean disease incubation time in days postinfection (dpi) ± standard deviation (SD) in days.

**FIG 1** Intracranial inoculation of FFPE sCJD brain tissue causes clinical or subclinical prion infection in Tg66 mice. (A) H&E (left panel)- and 3F4 (right panel)-stained sagittal sections from a clinically positive Tg66 mouse 167 days after inoculation with control CJD<sup>Froz</sup> brain homogenate. Spongiosis and PrP<sup>Sc</sup> deposition (brown stain) are present in the pons. Scale bar, 100 µm. (B) Section of cortex stained with the anti-PrP monoclonal antibody 3F4 (red stain). The arrow in the right panel indicates a unicentric PrP<sup>Sc</sup> plaque. Scale bar, 100 µm. (C) Western blot analysis of brain homogenates from Tg66 mice inoculated with homogenates made from FFPE sCJD brain samples CJD<sub>1FFPE</sub> and CJD<sub>2FFPE</sub>. Each lane represents 0.7-mg brain equivalents of an individual mouse brain, except for the lane labeled CJD<sub>Froz</sub>, which represents brain homogenate from the sCJD MM type 1-positive control sample. The CJD<sub>1FFPE</sub> sample lanes 1 to 3 represent three mice that were euthanized for non-prion-disease-related causes at 113, 127, and 131 days postinfection. The blot was developed with the anti-PrP mouse monoclonal antibody 3F4 and the ECL development system. Molecular mass markers are indicated on the left. (D) H&E-, 3F4-, or Thioflavin S (ThioS)-stained sagittal brain sections from a Tg66 mouse at 234 (CJD1<sub>FFPE</sub>) or 648 (CJD2<sub>FFPE</sub>) days postinoculation with brain homogenate derived from FFPE sCJD brain samples. Spongiosis and PrP<sup>Sc</sup> deposition (brown stain) are present in the midbrain in CJD1<sub>FFPE</sub>-inoculated mice and in the pons in CJD2<sub>FFPE</sub>-inoculated mice. The arrows in the top two right panels indicate a plaque visible in both the H&E- and 3F4-stained sections. ThioS-positive plaques were present only in animals inoculated with the CJD2<sub>FFPE</sub> sample. Scale bar, 100 µm.

**Detection of PrP<sup>Sc</sup> in archival formalin-fixed human brain tissue samples.** Formalin-fixed and paraffin-embedded brain and heart tissue from two patients diagnosed with sCJD in 1979 and 1981 were obtained from the Armed Forces Institute of Pathology. Since these individuals were diagnosed prior to the discovery of prion protein, it was unclear whether or not the fixed brain samples were positive for PrP<sup>Sc</sup>. In order to confirm that PrP<sup>Sc</sup> was present, sections were taken from each sample and stained with...
the mouse monoclonal antibody 3F4, which recognizes an epitope within human PrP. As shown in Fig. 1B, brain tissue from the first sample (CJD1FFPE) showed significant PrPSc deposition and spongiform pathology. PrPSc was found in dense perivascular deposits, as well as in both coarse and synaptic-like deposits. Brain tissue from the second sample (CJD2FFPE) was also positive for PrPSc but, unlike CJD1FFPE, demonstrated occasional unicentric plaque-like deposits of PrPSc (Fig. 1B, arrow) in addition to occasional perineuronal and synaptic-like deposition. In the limited sections analyzed, spongiform change did not appear to be as extensive in CJD2FFPE as in CJD1FFPE. In contrast to the brain samples, neither heart tissue sample showed any PrP positivity using the anti-PrP antibody 3F4 (Fig. 2A). PrPSc was therefore easily detectable in brain, but not heart, samples from both sCJD patients. The presence of PrPSc in the brains of these patients definitively confirmed the original diagnosis of sCJD.

Detection of sCJD infectivity in archival FFPE human brain tissue. In order to determine whether sCJD infectivity was present in the FFPE brain tissue samples, both CJD1FFPE and CJD2FFPE samples, as well as a non-CJD FFPE brain (NBr1FFPE) were removed from the block, rehydrated, and homogenized in PBS to make a final 10% (wt/vol) brain homogenate. This material was then inoculated i.c. into Tg66 mice, and inoculated animals were monitored for disease. No clinical disease was observed in mice inoculated with NBr1FFPE (Table 3). In contrast, Tg66 mice inoculated with the CJD1FFPE sample developed clinical disease with an incubation time of ~214 days, longer than control mice inoculated with homogenate derived from the fixed human heart sample CJD1FFPE (Fig. 2A). PrPSc was therefore detectable in brain, but not heart, samples from both sCJD patients. The presence of PrPSc in the brains of these patients definitively confirmed the original diagnosis of sCJD.

FIG 2 Lack of PrPSc and spongiform pathology after i.c. inoculation of FFPE sCJD heart tissue into Tg66 mice. (A) Section of heart stained with the anti-PrP mouse monoclonal antibody 3F4 from a patient diagnosed with sCJD in 1979 (CJD1FFPE) and 1981 (CJD2FFPE). Both sections are negative for PrP. Scale bar, 100 μm. (B) H&E- and 3F4-stained sagittal brain section from Tg66 mice 755 days postinoculation with heart homogenate derived from samples CJD1FFPE and CJD2FFPE. Unlike mice inoculated with the corresponding brain material (Fig. 1D), there is no spongiform change or PrPSc deposition in the midbrain in CJD1FFPE-inoculated mice or in the pons in CJD2FFPE-inoculated mice. Scale bar, 100 μm. (C) Western blot analysis of brain homogenates from Tg66 mice inoculated with FFPE heart homogenates derived from samples CJD1FFPE and CJD2FFPE. Each lane represents 0.7-mg brain equivalents of an individual mouse brain. MoCJD1FFPE, brain homogenate from a Tg66 mouse inoculated with CJD1FFPE; NHt1FFPE, brain homogenate from a Tg66 mouse inoculated with FFPE normal human heart sample. The blot was developed with the anti-PrP mouse monoclonal antibody 3F4 and the ECL advanced development system. Molecular mass markers are indicated on the left.

mice that were lost due to intercurrent disease between 113 and 131 days (Table 3) had little or no detectable PrPSc (Fig. 1C, CJD1FFPE lanes 1 to 3). Perineuronal and synaptic-like deposition of PrPSc, as well as spongiform change, was seen primarily in the midbrain (Fig. 1D) and pons in clinically ill mice. PrPSc was not deposited as amyloid, as indicated by the lack of thioflavin S (ThioS) staining (Fig. 1D). A second passage of CJD1FFPE into Tg66 also caused clinical disease after ~180 days (data not shown), confirming the presence of sCJD prion infectivity in the original sample. Thus, sufficient prion infectivity was present in the CJD1FFPE sample to initiate prion infection and PrPSc formation in all surviving Tg66 mice.

In contrast to both the CJD1FFPE sample and the positive control sample CJDProx, a homogenate derived from the fixed human brain sample CJD2FFPE did not cause clinical disease in Tg66 mice after i.c. inoculation (Table 1). However, all mice inoculated with CJD2FFPE were positive for PrPSc by Western blot analysis (Fig. 1C), suggesting that they were subclinically infected. These results were confirmed via immunohistochemical analysis using the 3F4 antibody (Fig. 1D). Unlike CJD1FFPE, mice inoculated with the CJD2FFPE sample had synaptic-like PrPSc staining as well as large, PrPSc-positive plaques in the pons (Fig. 1D), hypothalamus, cortex, and corpus callosum, as well as subpially (data not shown). These plaques were ThioS positive (Fig. 1D), indicating that PrPSc was deposited as amyloid. In addition, both spongiform change and large plaques were visible by hematoxylin and eosin (H&E) staining (Fig. 1D). A second passage of CJD2FFPE into Tg66 is in progress and has resulted in clinical disease roughly 400 days postinoculation (data not shown), confirming that the first passage into Tg66 mice induced a subclinical prion infection. Thus, using Tg66 mice, we were able to detect both clinical and subclinical
heart tissue from samples CJD1FFPE and CJD2FFPE contained brain or a non-CNS tissue (41). Therefore, if the formalin-fixed the same whether the infectious homogenate is derived from the pathological profile associated with a given prion isolate remains determined, none of the inoculated mice had shown any sign of clinical monitored for disease. After 755 days, when the experiment was termi-

PBS to make a final 10% (wt/vol) heart homogenate. This material were removed from the block, rehydrated, and homogenized in formalin-fixed and paraffin-embedded CJD1FFPE and CJD2FFPE heart samples, both of which were determined to be negative for PrPSc by immunohistochemistry using the 3F4 antibody (Fig. 2A), were removed from the block, rehydrated, and homogenized in PBS to make a final 10% (wt/vol) heart homogenate. This material was then inoculated i.c. into Tg66 mice, and the mice were mon-
disease (Table 3). There was no PrPSc deposition in any area of the brain (Fig. 2B), including the regions of the brain that had been made positive for PrP, an example of which was then inoculated i.c. into Tg66 mice, and the mice were monitored for disease. After 755 days, when the experiment was termi-
nated, none of the inoculated mice had shown any sign of clinical disease (Table 3). There was no PrPSc deposition in any area of the brain (Fig. 2B), including the regions of the brain that had been strongly PrPSc positive following inoculation of the corresponding FFPE brain samples. The lack of PrPSc in the brain was confirmed by Western blot analysis (Fig. 2C). Thus, we were unable to detect any sign of prion infectivity in heart tissue from two sCJD-positive individuals.

Detection of prion protein in amyloid heart disease tissue. We have previously shown that transgenic mice expressing PrPc without the GPI membrane anchor can accumulate high levels of PrPSc amyloid and infectivity in the heart after infection with mouse scrapie (18). These results suggested the possibility that abnormal prion protein deposited as amyloid in human heart tissue might also be infectious. We tested heart tissue samples from 42 cases of human amyloid heart disease for the presence of protease-resistant prion protein. Of the 42 samples tested, 6 were positive for PrP, an example of which (AHD1FFPE) is shown in Fig. 3. ThioS staining confirmed the presence of cardiac amyloid in AHD1FFPE (Fig. 3A) but not in normal heart tissue (Fig. 3B). Cells in the AHD1FFPE sample were positive using the anti-PrP antibody PV30 (Fig. 3C), which has a very high affinity for human PrPc (see Materials and Methods). The positive signal remained following PK diges-
tion (Fig. 3E), suggesting the presence of protease-resistant PrP. The specificity of the stain for PrP was confirmed by the loss of positive signal using PV30 antibody that had been pre-


dation time ± standard deviation (SD) in days or the range in days of non-TSE-related intercurrent deaths. dpi, days postinfection.

d The number excludes three intercurrent deaths at 113, 127, and 131 days postinfection (dpi). Brain tissue from these mice had little or no PrPSc (see Fig. 1).

d This number excludes one intercurrent death at 88 dpi.

levels of sCJD infectivity in FFPE brain tissue which, at the time of inoculation, was >26 years old.

Lack of prion infectivity in sCJD heart tissue. The neuro-
pathological profile associated with a given prion isolate remains the same whether the infectious homogenate is derived from the brain or a non-CNS tissue (41). Therefore, if the formalin-fixed heart tissue from samples CJD1FFPE and CJD2FFPE contained prion infectivity, they should trigger the same pathology as the corresponding fixed brain tissue when inoculated into Tg66 mice. Formalin-fixed and paraffin-embedded CJD1FFPE and CJD2FFPE heart samples, both of which were determined to be negative for PrPSc by immunohistochemistry using the 3F4 antibody (Fig. 2A), were removed from the block, rehydrated, and homogenized in PBS to make a final 10% (wt/vol) heart homogenate. This material was then inoculated i.c. into Tg66 mice, and the mice were monitored for disease. After 755 days, when the experiment was terminated, none of the inoculated mice had shown any sign of clinical disease (Table 3). There was no PrPSc deposition in any area of the brain (Fig. 2B), including the regions of the brain that had been strongly PrPSc positive following inoculation of the corresponding FFPE brain samples. The lack of PrPSc in the brain was confirmed by Western blot analysis (Fig. 2C). Thus, we were unable to detect any sign of prion infectivity in heart tissue from two sCJD-positive individuals.

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DISCUSSION

We were unable to induce clinical or subclinical prion infection following inoculation of fixed heart tissue from two sCJD patients into transgenic mice overexpressing the human prion protein. One caveat to our experiments is that we used FFPE tissue in our inoculations. It has been known since the 1930s that scrapie can survive formalin fixation (42) but, as shown here (Table 2) and in previous work (40), fixation in formalin can reduce infectivity by several logs. This could have an impact in tissues with lower levels of prion infectivity. For example, a recent study has shown that, while homogenates derived from vCJD appendix could transmit disease to 50% of transgenic mice expressing human PrP\(^{\text{FIP}}\), an FFPE vCJD appendix did not cause an infection (43). We therefore cannot exclude the possibility that low levels of infectivity which might have been present in sCJD heart tissue were inactivated following formalin fixation and paraffin embedding. However, our results are in agreement with previous studies that used sensitive Western blotting techniques (9, 12), immunohistochemistry (19), or inoculation of nonhuman primates (25) to assay for sCJD infectivity in heart tissue. Thus, our results are consistent with the conclusion that prion infectivity is not likely to be present in the hearts of patients with sCJD.

A common amino acid polymorphism at codon 129 in human PrP, encoding either a methionine or valine, can influence the efficient transmission of human prions into transgenic mice expressing human PrP (44–46). With the exception of AHD2\(^{\text{FFPE}}\),
which is homozygous for methionine at codon 129 (see Materials and Methods), we have as yet been unable to clearly determine the amino acid residue at codon 129 of our FFPE tissue samples. It is therefore possible that a mismatch at codon 129 could influence the transmission of infectivity into Tg66 mice, which overexpress human PrP with a methionine at codon 129. It is important to note, however, that inoculation of sCJD from different Prnp genotypes can still lead to efficient clinical and/or subclinical infection in transgenic mice overexpressing human PrP with a methionine at codon 129 (45). Thus, if sufficient levels of infectivity were present, we would still expect that, regardless of genotype, some percentage of Tg66 mice inoculated with PrP-positive FFPE heart tissue would be positive for PrPSc by immunohistochemistry or Western blotting.

The identification of an infectious cardiac amyloidosis with high levels of PrPSc amyloid and infectivity in the heart of a transgenic mouse model of scrapie (18) raised the possibility that abnormal forms of PrP might also be an unrecognized component of amyloid heart disease in humans. Interestingly, Jansen et al. (15) found mutations in human Prnp that resulted in the absence of the GPI anchor in PrP, which predisposed to amyloid plaque formation in the CNS. Given the multiorgan distribution of PrPSc amyloid in mice expressing PrP\textsuperscript{C} without the GPI anchor (17) and the recent identification of a prion amyloid myopathy in a BSE-infected macaque (22), PrPSc amyloid might also be present outside of the CNS in humans with anchorless PrP\textsuperscript{C}. However, the presence of PrP\textsuperscript{Sc} and prion infectivity in the heart and other organs has not yet been assessed in the human cases where PrP does not contain the GPI anchor (15). Consistent with the possibility that amyloid heart tissue might contain prion infectivity, we were able to detect PrP positivity in 6/42 (14%) human amyloid heart disease samples. However, inoculation of two of these positive samples into Tg66 mice did not lead to disease (Table 3), indicating that the PrP detected was not infectious PrP\textsuperscript{Sc}.

Although the signal in the PV30-positive amyloid heart samples does not appear to be infectious PrP\textsuperscript{Sc}, the fact that it could be blocked by a PrP peptide specific to PV30 (Fig. 3F) suggests that it could a form of PrP that is abnormal but noninfectious. It is well known that normal PrP\textsuperscript{C} expression can increase under stressful conditions such as hypoxia (47) or ischemia (48) and that the overexpression of normal PrP\textsuperscript{C} in muscle tissue can lead to increased protease resistance and myopathy (49, 50). It is therefore possible that the PrP signal detected by PV30, which was cell-associated (Fig. 3C and E), was a result of some form of cellular stress which led to an increase in PrP\textsuperscript{C} expression and the production of abnormal forms of PrP that were more protease resistant. This would explain why PrP-specific signal was detected by the PV30 antibody even after PK digestion. Consistent with this possibility, cell-associated PrP\textsuperscript{C} staining was not detected in nonamyloid heart tissue (Fig. 3D) or in heart tissue from the two sCJD cases exam-
ined (Fig. 2A and data not shown). That PrP was detected only in a minority of the amyloid heart disease cases examined may explain why PrP-positive staining was not seen in a previous report that examined a smaller number of human amyloid heart samples (19).

One advantage of the present study over previous infectivity studies of sCJD heart is in the assay system used, i.e., the Tg66 mice. Unlike the nonhuman primate studies, and as evidenced by the fact that there was no change in incubation time when sCJD MM type I was passaged in Tg66 mice three consecutive times, there is no significant species barrier to transmission of the most common form of sCJD. Furthermore, the mice appear to be sensitive enough to detect relatively low levels of infectivity. Inoculation of the CJD1FFPE sample led to clinical disease in all recipient mice, indicating that the tissue analyzed contained more than a single 50% lethal dose (LD50), i.e., the dose of inoculum that would lead to fatal disease in 50% of the animals. However, while inoculation of the CJD2FFPE brain sample also led to PrPSc accumulation in all of the mice, there was insufficient infectivity to induce clinical disease. This indicated that less than one LD50 was present in the CJD2FFPE sample. Thus, these data suggest that the Tg66 mice are a sensitive enough assay system to detect less than one LD50/50 mL of 1% brain homogenate. By comparison, it has been conservatively estimated that a highly sensitive Western blotting technique can detect brain PrPSc levels equivalent to ≥1,000 LD50/mL (i.e., ≥20 LD50/50 mL) (12).

Earlier studies had demonstrated that CJD infectivity could be recovered from FFPE tissue that had been stored for 7 months (51) and 21 months (52). Our results demonstrate that FFPE sCJD brain tissue stored for as long as 28 years still contains sufficient infectivity to infect 100% of inoculated animals. Thus, FFPE sCJD tissue can harbor >1 LD50 of sCJD infectivity. The fact that prions can survive for decades in archival tissues reinforces the fact that extra precautions should always be used when manipulating or disposing of archival sCJD tissues.

Our results also suggest that the characteristics of different sCJD prion isolates can be maintained following formalin fixation and storage over decades. Deposition of PrPSc in the brain samples analyzed differed between the two isolates. In particular, unicentric PrPSc plaques were frequently found in the CJD2FFPE sample but not in the CJD1FFPE sample (Fig. 1B). The pattern of PrPSc deposition in the brains of Tg66 mice inoculated with CJD1FFPE or CJD2FFPE brain material was consistent with this difference. Mice inoculated with CJD2FFPE had abundant PrPSc-positive plaques, whereas CJD1FFPE inoculated mice did not (Fig. 1D). To the best of our knowledge, these are the first data to suggest that formalin fixation, followed by paraffin embedding and long-term storage, does not eliminate the differences in neuropathological phenotypes induced by different prion isolates.

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