Preclinical deposition of pathological prion protein PrPSc in muscles of hamsters orally exposed to scrapie

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Recently, pathological prion protein PrPSc, the putative key constituent of infectious agents causing transmissible spongiform encephalopathies (TSEs), was found in muscles of rodents experimentally infected with scrapie and in patients with Creutzfeldt-Jakob disease (CJD). For the assessment of risk scenarios originating from these findings (e.g., alimentary transmission of pathogens associated with bovine spongiform encephalopathy [BSE] and chronic wasting disease [CWD] via tainted beef and game or iatrogenic dissemination of CJD agent through contaminated surgical instruments) more detailed information about the time course of PrPSc accumulation in muscles at preclinical and clinical stages of incubation is needed. Here we show that PrPSc in muscles of hamsters fed with scrapie can be detected prior to the onset of clinical symptoms, but that the bulk of PrPSc was deposited late in clinical disease. Additionally, regarding the question of how muscles become invaded, we report on the intramuscular location of PrPSc and substantial indications for centrifugal spread of infection from spinal motor neurons to myofibers. Our findings in a well-established animal model for TSEs contribute to a better assessment of the risks for public health emanating from “Prions in skeletal muscle” and provide new insights into the pathophysiological spread of TSE agents through the body.

Introduction
Transmissible spongiform encephalopathies (TSEs) such as scrapie in sheep, bovine spongiform encephalopathy (BSE) in cattle, chronic wasting disease (CWD) in elk and deer, or Creutzfeldt-Jakob disease (CJD) and its variant form (vCJD) in humans are a group of fatal neurodegenerative disorders of the CNS. Apart from neuropathological features such as gliosis, neuronal cell loss, and spongiform change, these diseases are consistently characterized by the deposition in the CNS of a misfolded form of prion protein (PrP), which is known as PrPSc (1). According to the prion hypothesis (2), the causative agents of TSEs (so-called “prions”) are composed essentially of PrPSc, and although the exact molecular structure of these pathogens remains elusive, there is substantial evidence that PrPSc can be used as a reliable biochemical marker for TSE agents (3–6). PrPSc deposition in TSEs is not necessarily confined to the CNS; it has also been observed in a variety of peripheral tissues from donors with experimental and nonexperimental TSEs (1). Irrespective of an incidental observation in experimental caprine scrapie (7), however, muscle tissue from TSE-affected animals or humans was essentially found to be negative for infectivity or PrPSc in a variety of studies (6, 8–10).

The detection of substantial amounts of PrPSc in muscle tissue of mice and hamsters experimentally infected with scrapie (11–13) and of patients with sporadic CJD (14), including those affected by inclusion body myositis (15), has been reported only recently in a series of articles by different research groups. These new findings have revived concerns about the potential risks for public health emanating from possibly unrecognized reservoirs for PrPSc and infectivity in muscles of animals and humans incubating TSEs. Particularly, the theoretical possibility of transmitting pathogens associated with BSE and CWD to humans via tainted beef and game, or that of iatrogenic dissemination of CJD or vCJD agent through contaminated surgical instruments used on muscle tissue, have gained considerable attention. While the assessment of such risk scenarios is certainly difficult, it could greatly benefit from a better knowledge of the time course of PrPSc deposition in muscles at preclinical and clinical stages of incubation.

So far, apart from studies on the tongue of hamsters infected by intralingual injection of TSE agent (13), findings have been reported only on PrPSc or TSE infectivity in muscles from donors already exhibiting clinical symptoms (7, 11, 12, 14). In this article we present data from a time-course study in hamsters fed with 263K scrapie, addressing the topical question of preclinical PrPSc deposition in skeletal muscle.

Following up on experimental pathogenesis studies in mice (16–19) and sheep (20), the animal model used in this study has provided key observations on the spread of infection through the body (4, 21–24) that have been confirmed in naturally occurring ovine scrapie (25), field cases of BSE (26), orally transmitted BSE in primates (27), and orally transmitted or naturally occurring CWD (28, 29). Additionally, more recent reports on the involvement of the sympathetic nervous system in vCJD (30) and the deposition of PrPSc in muscles of patients with sporadic CJD (14) have shown the relevance of our hamster model for peripheral PrPSc routing in TSE-affected humans.

Here we report that PrPSc in muscles of hamsters orally infected with scrapie was already detectable shortly before the onset of clinical symptoms, but that the bulk of PrPSc accumulated only...
late in clinical disease. Improved methods for paraffin-embedded tissue (PET) blotting and immunohistochemistry (IHC) provided detailed information about the intramuscular location of PrPSc and led to new conceptual insights into the pathophysiology of peripheral PrPSc routing.

**Results**

**Time course of PrPSc deposition in muscle tissue.** To assess whether, and to what extent, the biochemical marker for TSE infectivity, PrP Sc, accumulates at preclinical and clinical stages of incubation in skeletal, heart, and tongue muscles of hamsters orally challenged with 263K scrapie, the pathological prion protein was visualized by sensitive Western blotting after extraction in the form of PrP27-30 (protease-resistant core of the pathological prion protein PrPSc) from tissue samples by a high-yield purification method. PrP27-30 could not be detected in any of the examined muscle or sciatic nerve samples from six hamsters sacrificed at 100 days after infection (Figure 1A), that is, at about 60% of the mean incubation period until terminal disease (166 ± 8 days, expressed as the mean ± SD; n = 20, with animals also taken from other experimental series). The same holds true for four out of seven animals sacrificed before the onset of scrapie symptoms at 130 days after infection (Figure 1B). In two of the asymptomatic animals culled at 130 days after infection, however, at least one muscle (musculus [M.] tibialis cranialis, Figure 1C, or M. masseter, not shown) was found clearly positive for PrP27-30, and in

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**Figure 1**

Time course of PrPSc deposition in muscle tissue. Western blot detection of PrP27-30 extracted from different muscles and sciatic nerve of hamsters orally challenged with 263K scrapie and sacrificed at the following time points after infection. (A) At 100 days after infection, (B–D) 130 days after infection, (E) onset of clinical signs for scrapie (139–149 days after infection), and (F) at the terminal stage of disease (161–173 days after infection). Lanes with test samples: M1, M. biceps femoris (hindlimb); M2, M. tibialis cranialis (hindlimb); M3, M. triceps brachii (forelimb); M4, M. extensor carpi radialis (forelimb); M5, M. trapezius (shoulder); M6, M. masseter (head); M7, M. psoas major (back); T, tongue; H, heart; SN, sciatic nerve. Lanes with control samples: 1, proteinase K-digested brain homogenate from terminally ill scrapie hamsters containing 1 × 10⁻⁷ g (B and F) or 5 × 10⁻⁷ g (A, C–E) brain tissue; 2, skeletal muscle from an uninfected hamster spiked before extraction with 5 × 10⁻⁶ g (A, B, and F), 1 × 10⁻⁵ g (C and D), or 2 × 10⁻⁵ g (E) brain homogenate from terminally ill scrapie hamsters; 3, skeletal muscle from a mock-challenged hamster sacrificed at 173 days after infection (F). For each stage of incubation representative results are shown. (G) Time scale displaying the mean incubation period and the preclinical and clinical phases of incubation of hamsters orally infected with 263K scrapie. Small vertical arrows indicate time points at which animals were screened for PrPSc in muscles. dai, days after infection.
PrPSc as demonstrated by strong PrP27-30 immunostaining in the Western blot. Heart muscle was only very slightly positive in two out of six terminally ill animals (the sample displayed in Figure 1F, lane H, exhibits no signal for PrP27-30). Muscle samples from mock-challenged age-matched control hamsters consistently gave negative results (Figure 1F, lane 3, shows one representative sample: M. extensor carpi radialis). Detection thresholds for PrPSc/PrP27-30 in the Western blot analyses were in the range previously reported (12) as validated in this study by positive controls of muscle tissue from uninfected hamsters spiked with either 5 × 10^-6 g, 1 × 10^-5 g, or 2 × 10^-5 g of homogenized scrapie-infected hamster brain from terminally ill donors (Figure 1, A–F, lane 2).

As far as the blotting results shown in Figure 1 allow a comparison of the glycosylation and electrophoretic migration patterns of PrP27-30 from muscles, sciatic nerve, and brain, no significant differences can be discerned despite the divergent nature of these tissues. Our Western blot experiments, however, were primarily designed for the sensitive detection of pathological prion protein. A thorough molecular characterization of muscle-associated PrPSc, which needs to be performed in future studies, will require the Western blot colation of carefully adjusted amounts of PrP27-30 that produce optimized staining intensities and clearly visible unglycosylated bands.

**Intramuscular location of PrPSc.** The morphological PET blot analysis of the distribution of proteinase K-resistant PrP in muscle tissues revealed the deposition of PrPSc in muscle fibers (Figure 3, A, D, and E; arrowheads) and nerve fascicles (Figure 3, A and E; arrows). In nerve fascicles, which consistently appeared somewhat less-intensely stained than muscle fibers, the deposition was endoneural (inset in Figure 3E). In muscle fascicles, PrPSc was found in adjacent, as well as topologically distant, muscle fibers (prominent in Figure 3D; arrowheads), similar to a distribution that would be expected for motor units innervated by individual motor neurons in the spinal cord (31, 32). Within individual muscle fibers, PrPSc was found predominantly subsarcolemmal in the region of the fiber surface but also granularly scattered (Figure 3D). In some fibers PrPSc seems to be associated with structures reminiscent of neuromuscular junctions as exemplarily indicated by the strong staining at the right border of the myofiber marked by an asterisk (Figure 3A and inset).

To corroborate these PET blot findings, we tried to visualize PrPSc deposition in muscles by IHC. To reliably avoid misleading IHC immunostaining in muscle samples, a specifically designed protocol needed to be established that included pretreatment of specimens by hyaluronidase and proteinase K. The immunohistochemical examinations confirmed the subsarcolemmal location of PrPSc as well as the protein’s granularly scattered occurrence within muscle fibers (Figure 3F and inset). Because of the lower sensitivity of IHC as compared with PET blotting and other technical difficulties, however, immunohistochemical detection of PrPSc could be achieved only in a small proportion of the analyzed samples (Figure 3F). In the majority of specimens found to be positive by PET blotting, IHC produced negative results (Figure 3C). Muscle samples from mock-challenged control hamsters consistently gave negative PET blot and IHC results (not shown).

**Infectivity in muscle tissue.** To formally verify that PrPSc found in the muscles of our model animals correlates with TSE infectivity, we performed a supplementary series of bioassays in which hamsters were intracerebrally challenged with homogenized muscle tissue from terminally ill scrapie hamsters. Four different skeletal muscles and tongue specimens showed a consistent accumulation of PrPSc in muscles perorally infected with 263K scrapie (A and B). Animals for which all muscles and the sciatic nerve were found negative for PrPSc at 130 days after infection displayed less widespread and intense deposition of PrPSc in the brain stem (A) than observed in the corresponding brain region (B) of animals already showing PrPSc in their muscle tissue. Scale bars in A and B, 500 μm.

**Figure 2**

PET blotting of preclinical cerebral PrPSc deposition in the brain stem. The slices comprise the area of the dorsal motor nucleus of the vagus nerve that provides the initial site of PrPSc deposition in the brain of hamsters perorally infected with 263K scrapie (A and B). Animals for which all muscles and the sciatic nerve were found negative for PrPSc at 130 days after infection displayed less widespread and intense deposition of PrPSc in the brain stem (A) than observed in the corresponding brain region (B) of animals already showing PrPSc in their muscle tissue. Scale bars in A and B, 500 μm.
muscles (M. biceps femoris, M. triceps brachii, M. trapezius, M. psoas major) and the lingual muscle from two donor animals were titrated for infectivity. All reporter animals of which the incubation times were used to calculate infectivity titers succumbed to scrapie as verified by Western blot detection of PrPSc/PrP27-30 in their brains (Figure 4, lane 1).

The results of the infectivity bioassay are summarized in Table 1. Mortalities and incubation times for the recipients in the bioassay demonstrated a consistent presence of infectivity in the range of about $2 \times 10^4$ to $10 \times 10^5$ 50% intracerebral lethal doses (LD$_{50ic}$) per gram of tissue, except for the M. triceps brachii from the forelimb, which exhibited a surprisingly low infectivity titer of only 100–300 LD$_{50ic}$/g. Three hamsters inoculated with tissue from this muscle failed to exhibit clinical symptoms until 350 days after infection. Consistent with the low amount of infectivity revealed by bioassay for the inocula of M. triceps brachii, however, the inconspicuous reporter animals were subclinically infected as shown in Figure 4, lane 2.

The lingual muscle showed the highest infectivity titer of the examined specimens, a finding consistently mirrored by the Western blot results. The determined infectivity titers appear somewhat below the level expected on the basis of the observed PrPSc immunostaining intensities, however. The detected amount of PrP27-30 in the muscles from terminally ill hamsters is comparable to, or even higher than, that present in $5 \times 10^{-6}$ g of homogenized scrapie-infected hamster brain (Figure 1F, lanes M1–M7 and T versus lane 2). As previously discussed elsewhere (12), this would correspond to an infectivity titer of approximately $10^5$ LD$_{50ic}$/g if PrPSc in muscles correlated with infectivity at a ratio similar to that observed in the brain or spinal cord of 263K-infected hamsters. Thus, the bioassay findings possibly provide an underestimation of the infectivity in muscles since the titrated inocula, after digestion of samples in collagenase A, had been heated for technical reasons (see Methods) to 80°C before intracerebral administration. Theoretically, it appears conceivable that this treatment, which somewhat differs from conventional ways of tissue homogenization (11), may have partially inactivated the infectious scrapie agent originally present in situ.

Figure 3
Intramuscular location of PrPSc in terminally ill 263K scrapie hamsters visualized by PET blotting and IHC. (A) PET blot from M. triceps brachii. Individual muscle fibers display deposition of PrPSc predominantly subsarcolemmal in the region of the fiber surface, but PrPSc is also scattered within fibers (arrowheads). At some myofibers PrPSc immunostaining exhibits a structural pattern reminiscent of neuromuscular junctions (asterisk and inset). Only minor PrPSc deposits are visible within nerve fascicles (arrow). (B) Adjacent section to A, stained with H&E; arrow: nerve fascicle. (C) Adjacent section to A, immunohistochemically stained with mAb 3F4 without detectable PrPSc deposition; arrow: nerve fascicle. (D) PET blot from M. masseter; PrPSc in adjacent as well as topologically distant muscle fibers (arrowheads), distributed in a pattern reminiscent of that expected for motor units innervated by individual neurons in the spinal cord. Note: Tangentially cut muscle fiber (rhomb) with strong granularly scattered immunolabeling. Reproduction in higher magnification (inset: photomicrograph from a different section) illustrates prominent subsarcolemmal localization of PrPSc deposition in myofibers. (E) PET blot from lingual muscle; distinct endoneural PrPSc accumulation in nerve fascicles (arrows). One fascicle (asterisk) is reproduced in higher magnification in the inset. The intensity of PrPSc deposition in muscle fibers of the tongue (arrowheads) seems to be stronger than that observed in other muscles. (F) Sample from lingual muscle immunohistochemically stained with mAb 3F4; granular immunostaining illustrates subsarcolemmal as well as scattered PrPSc deposition in muscle fibers (arrowheads; inset: higher magnification of fiber marked by an asterisk). Scale bars in A–F, 100 μm.
Comprehensive inactivation studies, however, failed to reveal a decrease of infectivity in scrapie brain homogenate from 263K-infected hamsters heated under the conditions used in our titration experiments, that is, for 10 minutes to a temperature of 80°C (33). In light of the pertinent report by Rohwer (33), the sensitivity of the bioassay would appear unlikely to have been reduced — if at all — by more than 0.5 logs due to our method for the preparation of inocula. Therefore, the unexpectedly low infectivity titers found in muscle samples from our model animals may point to higher PrPSc/infectivity ratios in the examined specimens than previously established for CNS tissue (3–4). This explanation would also be consistent with the astonishingly low levels of infectivity measured in the M. triceps brachii despite the substantial amounts of PrP27-30 detected in that muscle.

**Discussion**

Our studies demonstrate that PrPSc can be detected in skeletal muscles of hamsters fed with scrapie before the animals show clinical signs of disease. Preclinical detection of pathological prion protein in muscle tissues was confined to relatively late stages of the asymptomatic phase of incubation, however. The bulk of muscle-associated PrPSc accumulated during clinical disease and all examined skeletal muscles and tongue specimens from terminally ill hamsters contained substantial amounts of PrPSc in accordance with previous observations (12). The accumulation of PrPSc found in the latter group of animals was accompanied by the consistent presence of at least 2 × 10⁴ to 10 × 10⁴ LD₅₀ of scrapie agent per gram of tissue in the M. biceps femoris, M. trapezius, M. psoas major, and lingual muscle. PET blot and IHC examinations corroborated the Western blot findings in our hamsters and provided new detailed information on the distribution and location of PrPSc in muscle tissue. Since the animal model used in this study in the past has been frequently found to exhibit key pathogenetic features of peripheral PrPSc routing (4, 12, 21–24) in several different alimentarily acquired and naturally occurring TSEs of animals and humans (14, 25–30), the findings reported here may well have several relevant implications:

Pathophysiology of peripheral PrPSc deposition in muscles. In experiments with transgenic mice exhibiting a four- to eightfold overexpression of the cellular isoform of the prion protein (PrPc) in myofibers under the control of myocyte-specific promoters, Bosque et al. (11) showed intrinsic replication of infectivity in muscles injected with scrapie agent. Here, with the more natural route of peroral infection, we have provided direct experimental evidence for the deposition of PrPSc in myofibers, particularly subsarcolemmally, and found that the deposition of this protein in muscles is also endoneurally associated with nerve fibers. Furthermore, the distribution patterns of PrPSc observed in muscles conspicuously resembled those known for the innervation of motor units (31, 32). Taken together, these findings are highly indicative of spread of infection from motor neurons in the spinal cord via axonal projections to neuromuscular junctions and on from there postsynaptically into muscle fibers. The results from the Western blot studies clearly show that PrPSc can be detected in muscles only shortly before the onset of clinical symptoms, that is, at a time point when the CNS is already heavily loaded with PrPSc and infectivity. This, again, would most plausibly be explained by projections of the peripheral nervous system mediating centrifugal infection of muscles from the spinal cord. Further time-course studies will be necessary, however, to validate the proposed neural spreading pathway and to elucidate, or rule out, the relevance of alternative mediators for muscle invasion such as lymph and blood or neural pathways linked with the centripetal ascension of TSE agent to the CNS.

**Table 1**

Detection of infectivity in muscles of terminally ill hamsters orally infected with 263K scrapie

<table>
<thead>
<tr>
<th>Donor</th>
<th>Sample (Muscle)</th>
<th>Mortality due to scrapie (until 350 dai)</th>
<th>Incubation time (days)</th>
<th>Infectivity titer (10⁺ LD₅₀/g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Hindlimb (M. biceps femoris)</td>
<td>4/4</td>
<td>131, 166, 215, 215</td>
<td>2.0</td>
</tr>
<tr>
<td>2</td>
<td>Hindlimb (M. biceps femoris)</td>
<td>5/5</td>
<td>134, 136, 180, 343, 349</td>
<td>2.0</td>
</tr>
<tr>
<td>1</td>
<td>Forelimb (M. triceps brachii)</td>
<td>5/5</td>
<td>187, 225, 254, 260, 292</td>
<td>0.3</td>
</tr>
<tr>
<td>1</td>
<td>Forelimb (M. triceps brachii)</td>
<td>2/5</td>
<td>267, 338</td>
<td>0.1</td>
</tr>
<tr>
<td>1</td>
<td>Shoulder (M. trapezius)</td>
<td>5/5</td>
<td>113, 134, 208, 232, 236</td>
<td>10.9</td>
</tr>
<tr>
<td>1</td>
<td>Shoulder (M. trapezius)</td>
<td>5/5</td>
<td>124, 127, 131, 166, 243</td>
<td>5.7</td>
</tr>
<tr>
<td>1</td>
<td>Back (M. psoas major)</td>
<td>5/5</td>
<td>120, 145, 208, 211, 232</td>
<td>7.6</td>
</tr>
<tr>
<td>1</td>
<td>Back (M. psoas major)</td>
<td>5/5</td>
<td>131, 131, 134, 141, 280</td>
<td>2.6</td>
</tr>
<tr>
<td>1</td>
<td>Lingual muscle</td>
<td>2/2</td>
<td>120, 131</td>
<td>8.1</td>
</tr>
<tr>
<td>2</td>
<td>Lingual muscle</td>
<td>3/3</td>
<td>121, 139, 164</td>
<td>11.7</td>
</tr>
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</table>

Aliquots (50 μl) of homogenized muscle samples from terminally ill hamsters orally challenged with 263K scrapie were inoculated intracerebrally into groups of two to five recipients. Mortality is shown until 349 days after infection, and the experiment was terminated at 350 days after infection.
Risks for public health and protection of consumers and patients.

What conclusions regarding the risks for public health emanating from "Prions in skeletal muscle" can be drawn from the findings described in this report?

For the reasons outlined above, the experimental paradigm used in our study is likely to provide a relevant model on the issue of muscle contamination. It probably reflects a worst-case rather than a best-case scenario, however, because the levels of infectivity and PrPSc produced in the CNS and other neural tissues of 263K scrapie-infected hamsters generally tend to be higher than in nonexperimental TSEs. Indeed, other than for caprine scrapie (7) or sporadic CJD (14), no infectivity or PrPSc has been detected so far in skeletal muscles of cattle with BSE (10).

If the findings in the hamster model were confirmed, to a varying degree, in ovine scrapie, BSE, or CWD, significant deposition of PrPSc in muscles of affected sheep, cattle, and elk or deer would be expected to occur only at relatively late stages of preclinical infection, with the bulk of muscle-associated PrPSc accumulating during clinical disease. At those stages of incubation, however, the TSE routine tests currently in use should allow both an identification of affected animals (34) with high reliability and their safe disposal. Thus, our findings strongly support the schemes for active TSE testing of animals that have already been implemented for the nosocomial transmission of human TSE agents. Epidemiological evidence substantiating this hypothetical hazard has not been reported, however, and in any case the risk can be considered as effectively addressed by comprehensive precautionary measures already implemented for infection control in nosocomial settings. Particularly, recommended procedures for the routine maintenance (cleaning, disinfection, and sterilization) of surgical instruments (35, 36) aimed at the reliable removal and inactivation of unrecognized contaminations with infectious agents are expected to substantially reduce iatrogenic risks possibly originating from TSE agents in muscles — if they are firmly observed.

Since 2002, a considerable body of new data on PrPSc and TSE infectivity in muscles of experimentally infected rodents and patients with sporadic CJD has been compiled (11–14). It remains to be examined in future studies whether, and to what extent, these findings can be transferred to nonexperimental scrapie, BSE, and CWD, or to vCJD in humans. Special attention should be paid to food products containing ruminant or cervid tongue, given the observations reported here and elsewhere (13). In any case, the data available so far leave no doubt that, 40 years after Pattison and Millson (7) reported finding the scrapie agent in muscle tissue of an experimentally infected goat, a topical field of TSE research has been (re)opened.

Methods

Animal inoculations and preparation of tissue samples

All animal experiments were performed in accordance with European and German legal and ethical regulations and approved by the responsible review boards and authorities. Outbred Syrian hamsters were perorally infected or mock-challenged with a single dose of homogenized 263K scrapie or normal hamster brain (corresponding to 10 mg of tissue), respectively, as previously described (4). Scrapie-infected animals were humanely sacrificed for Western blot examinations at 100 days after infection (n = 6), at 130 days after infection (n = 7), at the onset of at least two clinical signs for scrapie, such as tremor of head or whole body, incoordination of gait, or difficulty in rising up from a supine position (n = 5: 139, 145, 145, 146, and 149 days after infection), and at the terminal stage of disease (n = 6: 161, 163, 165, 170, 171, and 173 days after infection). For PET blotting of muscle samples and infectivity bioassays, five additional hamsters in the terminal stage of scrapie were sacrificed at 171, 173 and 174 and at 163 and 168 days after infection, respectively. Control animals (mock-challenged age-matched hamsters) were sacrificed at corresponding time points. Heart muscle (apex), lingual muscle (approximately 5 mm from the tip of the tongue), sciatic nerves, and seven different skeletal muscles from the hindlimb (M. biceps femoris; M. tibialis cranialis), the forelimb (M. triceps brachii [caput laterale]; M. extensor carpi radialis), the shoulder (M. trapezius), the head (M. masseter), and the back (M. psoas major) were dissected for Western blot examinations, and visible nerve fibers were removed from the muscle specimens. All samples for Western blot analyses were cut into small pieces and stored at −80°C until examination. The mass of the muscle samples ranged from approximately 40 to approximately 100 mg, and that of the sciatic nerves from approximately 5 to approximately 10 mg. Sampling from hamsters sacrificed for PET blot analysis and infectivity bioassays was performed as described below. To avoid cross contamination, instruments used for the preparation of samples were carefully cleaned after removal and processing of each specimen.

Western blot examinations

Muscle and nerve specimens to be examined by Western blotting were processed as previously described (12) for the extraction of PrPSc in the form of PrP27-30 (3, 37); the final pellets were resuspended in 15 μl of 2x sample loading buffer and heated to 100°C for 5 minutes before Western blotting using the anti-PrP mAb 3F4 (38).

Proteinase K-digested homogenate from scrapie hamster brains, used as a PrP27-30 reference in the Western blotting analyses, was prepared as outlined previously (12, 37).

SDS-PAGE and Western blot analyses were performed as described elsewhere (3, 37) with recently published modifications (12). PrP signals were visualized on a X-OMAT AR (Kodak; Sigma-Aldrich, Steinheim, Germany) film. Film was exposed for 30 minutes.

PET blot examinations

PET blotting of brain samples. Brains were immediately removed from sacrificed hamsters and immersed for 48 hours in 4% phosphate-buffered formalin as described by Schulz-Schaeffer et al. (39). After treatment with formic acid for 1 hour at room temperature, brains were placed in 4% phosphate-buffered formalin for a further 24 hours and subsequently trimmed coronally into 2- to 3-mm-thick slices. Samples were embedded in paraffin wax, and PET blotting was carried out as described (39), with slight modifications. Six-micrometer microtome slices were mounted onto nitrocellulose membranes and dried flat at 50°C for 16 hours. After washing, sections were digested with 250 μg/ml proteinase...
K (Roche Molecular Biochemicals, Mannheim, Germany) in PK digestion buffer containing 10 mM Tris-HCl (pH 7.8), 100 mM NaCl, and 0.1% Brij 35 for 8 hours at 55°C. Sections were then denatured in 3 M guanidine thiocyanate and blocked with 0.2% casein before labeling with mAb 3F4.

**PET blotting of muscle samples.** Morphological analysis of the distribution and location of PrP Sc in muscle tissue by PET blotting was performed on lingual muscle (approximately 5 mm from the tip of the tongue) and the seven different skeletal muscles specified above. The specimens were taken from three hamsters that developed terminal symptoms of scrapie after peroral infection at 171, 173, and 174 days after infection and from corresponding mock-infected controls. Animals were transcardially perfused with periodate-lysine-paraformaldehyde (22), and removed tissue samples were immersed additionally for 5 hours in periodate-lysine-paraformaldehyde. After incubation for 8 hours at 4°C in phosphate-buffered saline, muscle samples were transferred to 70% ethanol, processed in a tissue processor, and subsequently embedded in paraffin wax. For PET blot examinations of muscle tissue (2- to 3-μm-thick sections), modifications of the original protocol (39) were necessary to remove connective tissue. After prewetting blots with TBST (10 mM Tris-HCI, pH 7.8, 100 mM NaCl, 0.05 % Tween 20) sections were digested with 1.5 mg/ml collagenase A (Roche Molecular Biochemicals) in a buffer containing 10 mM Tris-HCI, pH 7.8, 100 mM NaCl, 100 mM CaCl2, and 0.1% Brij 35 for 30 minutes at 60°C followed by digestion using 250 μg/ml proteinase K (Roche Molecular Biochemicals) in PK digestion buffer (see above) for 8 hours at 55°C.

**Immunohistochemical examinations**

Examination of the distribution and location of PrP Sc in muscle tissue by IHC was performed on 2- to 3-μm-thick sections of paraffin-embedded muscle specimens prepared as described above (see PET blot examinations). Sections were pretreated with 2 mg/ml hyaluronidase (Sigma-Aldrich) in sodium acetate buffer (pH 5.2; 21 mM acetic acid, 79 mM sodium acetate) for 30 minutes at 37°C and then exposed at the same temperature and for the same time to 50 μg/ml proteinase K in TBS (10 mM Tris HCl, 133 mM NaCl, pH 7.4) containing 2 mM CaCl2. After incubation with the mAb 3F4 (1:500) and secondary alkaline phosphatase-coupled goat anti-mouse Ab (1:500; Dako, Hamburg, Germany) for 90 minutes each, immunostaining was performed using new fuchsin as chromogen. Subsequently, slides were counterstained with hemalun.

**Infectivity bioassay**

Infectivity titers in selected muscle specimens were assessed by an incubation time-interval assay (40, 41). For this purpose, four different skeletal muscles (M. biceps femoris, M. triceps brachii, M. trapezius, and M. psoas major) and the tip of the tongue were removed from two terminally ill scrapie hamsters sacrificed at 163 and 168 days after infection. Tissue specimens were homogenized in a way similar to that for the Western blot examinations. After removal of visible nerve fibers, samples were washed three times in TBS and incubated in a rocking device at 37°C for 3.5 hours in TBS containing 2 mM CaCl2 and 0.25% collagenase A (Roche Molecular Biochemicals). Collagenase A digestion was performed in a total volume of 500 μl per sample. Following ultrasonification, samples were heated to 80°C for 10 minutes in order to accomplish inactivation of the enzyme and of accidental microbial contaminations prior to intracerebral inoculation. Aliquots (50 μl) of the muscle homogenates were intracerebrally inoculated into groups of two to five recipient hamsters each, and the amount of infectivity in the samples was assayed as previously described (3–4) using the incubation times shown in Table 1 (mean error of assay ± 0.4 logs). Proteinase K-digested homogenates were prepared from the brains of all reporter animals (including those that failed to show scrapie symptoms), as described elsewhere (12, 37), and subjected to Western blot examination for PrP Sc/PrP27-30 as outlined above. The bioassay experiment was terminated at 350 days after infection.

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