Progress and problems in the biology, diagnostics, and therapeutics of prion diseases

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Abstract

The term "prion" was introduced by Stanley Prusiner in 1982 to describe the atypical infectious agent that causes transmissible spongiform encephalopathies, a group of infectious neurodegenerative diseases that include scrapie in sheep, Creutzfeldt-Jakob disease in humans, chronic wasting disease in cervids, and bovine spongiform encephalopathy in cattle. Over the past twenty years, the word "prion" has been taken to signify various subtly different concepts. In this article, we refer to the prion as the transmissible principle underlying prion diseases, without necessarily implying any specific biochemical or structural identity. When Prusiner started his seminal work, the study of transmissible spongiform encephalopathies was undertaken by only a handful of scientists. Since that time, the "mad cow" crisis has put prion diseases on the agenda of both politicians and the media. Significant progress has been made in prion disease research, and many aspects of prion pathogenesis are now understood. And yet the diagnostic procedures available for prion diseases are not nearly as sensitive as they ought to be, and no therapeutic intervention has been shown to reliably affect the course of the diseases. This article reviews recent progress in the areas of pathogenesis of, diagnostics of, and therapy for prion diseases and highlights some conspicuous problems that remain to be addressed in each of these fields.
The term “prion” was introduced by Stanley Prusiner in 1982 to describe the atypical infectious agent that causes transmissible spongiform encephalopathies, a group of infectious neurodegenerative diseases that include scrapie in sheep, Creutzfeldt-Jakob disease in humans, chronic wasting disease in cervids, and bovine spongiform encephalopathy in cattle. Over the past twenty years, the word “prion” has been taken to signify various subtly different concepts. In this article, we refer to the prion as the transmissible principle underlying prion diseases, without necessarily implying any specific biochemical or structural identity. When Prusiner started his seminal work, the study of transmissible spongiform encephalopathies was undertaken by only a handful of scientists. Since that time, the “mad cow” crisis has put prion diseases on the agenda of both politicians and the media. Significant progress has been made in prion disease research, and many aspects of prion pathogenesis are now understood. And yet the diagnostic procedures available for prion diseases are not nearly as sensitive as they ought to be, and no therapeutic intervention has been shown to reliably affect the course of the diseases. This article reviews recent progress in the areas of pathogenesis of, diagnostics of, and therapy for prion diseases and highlights some conspicuous problems that remain to be addressed in each of these fields.

Prion pathogenesis, diagnostics, and therapy: where do we stand?

Prion diseases, also known as transmissible spongiform encephalopathies (TSEs), are invariably fatal neurodegenerative disorders affecting a broad spectrum of host species and arise via genetic, infectious, or sporadic mechanisms (Table 1). In humans, prion diseases result from infectious modes of transmission (variant Creutzfeldt-Jakob disease [vCJD], iatrogenic CJD, Kuru); inherited modes of transmission in which there is nonconservative germ line mutation of the PRNP gene open reading frame (familial CJD, Gerstmann-Sträussler-Scheinker Syndrome, Fatal Familial Insomnia) (1, 2); and modes of transmission that have as yet been neither determined nor understood (sporadic CJD [sCJD]). The clinical symptoms associated with each of the human prion disease forms vary dramatically (2).

Nomenclature applied to prion biology continues to be complex and confusing to nonspecialists. Here we utilize the term “prion” to denote the causative agent of prion diseases, without implying associated structural properties. We refer to the disease-associated prion protein (PrPSc), a disease-specific isoform of the host-encoded cellular prion protein (PrP*), which accumulates in individuals affected with most forms of TSE (Figure 1) (3). While PrPSc is classically defined as partially protease-resistant, aggregated PrP, it has recently been shown that PrPSc may undergo disease-associated structural modifications that do not impart properties of inherent protease resistance (4). In light of this, it is advisable that PrPSc be defined on the basis of disease-associated structural modifications rather than properties of protease resistance.

Prion diseases are conceptually recent; the first cases of Creutzfeldt-Jakob disease were described eight decades ago (5, 6), yet the protein-only theory of prion infection was originally formulated in 1967 (7) and later refined and the term “prion” coined in 1982 (8). The precise physical nature of the prion agent is still the subject of intense scientific controversy. PrPSc may or may not be congruent with the infectious agent. It remains to be formally proven whether the infectious unit consists primarily or exclusively of: (a) a subspecies of PrPSc; (b) an intermediate form of PrP (PrP*) (9); (c) other host-derived proteins (10); or (d) nonprotein compounds (which may include glycaminoglycans and maybe even nucleic acids) (11). We still do not know, therefore, whether the prion hypothesis is correct in its entirety.

As with any other disease, a thorough mechanistic understanding of pathogenesis is the best foundation for devising sensitive predictive diagnostics and efficacious therapeutic regimens. The purpose of the present article is to discuss some aspects of the state of the art in prion science and their impact on prion diagnostics, primarily with respect to peripherally acquired prion disease. As of now, no causal therapies can be offered to prion disease victims. Yet we are witnessing the emergence of an impressive wealth of therapeutic approaches, some of which certainly deserve to be tested for their validity.

Progress in understanding prion pathogenesis

Prion pathogenesis is a dynamic process that can be broken down into spatially and temporally distinct phases: (a) infection and...
Peripheral replication; (b) transmigration from the periphery to the CNS (also termed “neuroinvasion”); and (c) neurodegeneration. But what are the mechanisms underlying neuroinvasion, and which cellular compartments are involved in replication and neuroinvasion of prions?

Peripheral replication

Cell tropism of prions varies dramatically among animal species and is also in part dependent on the particular strain of prion agent. For example, prions are lymphotropic in sheep scrapie and vCJD (12) but less so in sCJD (13) and bovine spongiform encephalopathy (BSE). Different prion strains can lead to different routes of peripheral replication in experimental models of scrapie (14, 15), and, therefore, strain-encoded properties might also determine the route of peripheral replication. With respect to peripheral pathogenesis of prion diseases, it is well established that replication of the prion agent occurs in high titers in lymphoid tissues such as spleen and lymph nodes well before neuroinvasion and subsequent detection in the CNS (16).

Upon oral challenge, an early rise in prion infectivity is observed in the distal ileum of infected organisms. This applies to several species but has been most extensively investigated in sheep scrapie and BSE. Different prion strains can lead to different routes of peripheral replication in experimental models of scrapie (14, 15), and, therefore, strain-encoded properties might also determine the route of peripheral replication. With respect to peripheral pathogenesis of prion diseases, it is well established that replication of the prion agent occurs in high titers in lymphoid tissues such as spleen and lymph nodes well before neuroinvasion and subsequent detection in the CNS (16).

Neuroinvasion

The resistance to prions of mice that lack expression of PrP(C, encoded by Prnp (a single-copy gene located on chromosome 2 in mice and 20 in humans), is well documented (19, 20). While the
precise physiological function of PrP"C is unclear, expression of it is absolutely required for transportation of the infectious agent both from the peripheral sites to the CNS (21) and within the CNS (22). However, reconstitution of Prnp knockout (Prnp"/"o) mice with WT bone marrow is insufficient to restore neuroinvasion in Prnp"/" mice (21). Hence one could argue that the elemental compartment required for prion neuroinvasion is stromal and must express PrP". Nevertheless, in adoptive transfer experiments on Prnp"/"o mice with WT bone marrow, the capability of the spleen to accumulate prions of the RML strain is restored (21, 23). This suggests that the interface between cells of the immune system and the CNS. (H) Shortened prion disease incubation period in CXCR5"/" mice inoculated intraperitoneally, relative to WT controls.

But just which cell types are involved in neuroinvasion? The innervation pattern of lymphoid organs is primarily sympathetic (32). Sympathectomy delays the onset of scrapie, while sympathetic hyperinnervation enhances splenic prion replication and neuroinvasion, which suggests that innervation of secondary lymphoid organs is the rate-limiting step to neuroinvasion (33). However, there is no physical synapse between FDCs and sympathetic nerve endings (34). So how can prions transmigrate from FDCs to sympathetic nerve fibers? A series of recent experiments (discussed below) may go some way toward providing answers.

FDC positioning is a primary determinant of velocity of neuroinvasion

We investigated how the distance between FDCs and splenic nerves affects the velocity of neuroinvasion, utilizing mice deficient in the CXC chemokine receptor 5 (CXCR5), which directs lymphocytes toward specific microcompartments (35). While density, distribution, and branching patterns of sympathetic nerve processes in CXCR5"/" spleens are normal, the distance between FDCs and nerve endings is greatly reduced (36).

After peripheral administration of high doses of prions, velocity of pathogenesis was similar in CXCR5"/" and WT mice; however, delivery of smaller inocula resulted in a dose-dependent increase in incubation periods in WT mice that was not evident in CXCR5"/" mice. Peripheral prion pathogenesis in CXCR5"/" mice is therefore more efficient upon incremental reduction of the inoculum.

What is the basis of this reduced incubation period? Kinetics measurements of prion infectivity titers in the thoracic spinal cord provided the answer: following peripheral administration, only traces of infectivity were found in WT spinal cords at 80 days post-inoculation (dpi), whereas infectivity rose to measurable levels in the spinal cords of CXCR5"/" mice already at 60 dpi. This suggests that increased velocity of prion entry into the CNS in CXCR5"/" mice is due to the repositioning of FDCs near highly innervated, splenic arterioles (Figure 2). This was validated by the finding that incubation periods were prolonged in CXCR5"/" mice treated with soluble LTβR-Ig to deplete mature FDCs.

Hence topographical relationships within lymphoid organs contribute to prion neuroinvasion. However, it remains to be determined whether this results from passive diffusion of prions or whether mobile cells (e.g., germinal center B cells) are involved in an active transport process.

This study also raises the possibility that spread of infection to peripheral nerves occurs more rapidly when FDCs are in close proximity to nerves in lymphoid tissue other than spleen, such as Peyers’s patches. Indeed, FDCs are crucial to disease progression but only during a short window of time following oral scrapie challenge (17). This implicates the efficiency of neuroimmune transfer of prions as a primary determinant of neuroinvasion. The detection of PrP"Sc in spleens of sCJD patients (12) suggests that the interface between cells of the immune system and...
Peripheral nerves (the neuroimmune connection) might also be of relevance in sporadic prion disease.

The neurodegeneration issue

There has certainly been progress in understanding the events underlying peripheral prion pathogenesis and neuroinvasion (37). However, prions exert their destructive effects exclusively within the CNS. The precise cause of neurodegeneration remains poorly understood and is a point of contention among prionologists. It seems unlikely that PrPSc is directly toxic, since tissue devoid of PrPSc that subsequently accumulates PrPSc remains healthy and free of pathology (20, 38). During the conversion process, PrPSc levels may be depleted, yet this is also an unlikely cause of pathology, since ablation of PrPSc does not result in scrapie-like symptoms (39), even when ablated postnatally (40).

Lindquist and colleagues have suggested a mechanism that may account for prion-associated toxicity: (a) expression of a PrP variant resident in the cytosol was strongly neurotoxic in cultured cells and transgenic mice, which suggests a common framework for diverse PrP neurodegenerative disorders (41); and (b) PrP, retrogradely transported out of the endoplasmic reticulum, produced amorphous aggregates of PrP possessing partial protease K resistance in the cytosol. Once conversion occurred, it was self-sustaining (42). It will be interesting to determine whether the disease generated in these mice is, in some way, transmissible. However, while the results obtained here are certainly intriguing, it should be noted that reports elsewhere, although not refuting these observations, argue against the contribution of such potential neurotoxic PrP species (43, 44). Similarly, it has been reported that PrPSc in some forms of prion disease assumes a transmembrane topology, C-terminal transmembrane PrP (CmPrP), and that the extent of neurotoxicity is a result of concentration of CmPrP, thereby arguing that CmPrP may represent a major toxic moiety (45, 46). However, while we still do not understand the biochemical events involved in cytosolic or CmPrP-induced neurotoxicity, elucidation of this may aid in the much-needed identification of therapeutic targets. Additionally, in-depth characterization of transgenic mice expressing amino-terminally truncated PrPSc (47), in which cerebellar neurodegeneration occurs, may not only aid in the elucidation of the molecular events responsible for potentially common neurodegenerative processes but perhaps also provide clues to the physiological function of PrPSc itself.

Prion diagnostics

The ability to secure early diagnosis is vital for therapeutic interventions to be of real value. With respect to animals destined for the human food chain, there is the additional demand to determine presence of the prion agent in tissues in asymptomatic organisms, well before the appearance of any clinical symptoms. This applies equally to the detection of prions in humans, who may participate in tissue donation programs.

Prions were transmitted via blood transfusion in sheep using blood obtained from infected animals prior to the onset of clinical symptoms (48, 49). If the same route applies to humans, this could represent a nightmare scenario for the blood transfusion services (50). A transfusion recipient received blood from an individual harboring the vCJD agent 3.5 years prior to the development of...
any clinical signs of prion disease in the donor. The unfortunate recipient developed disease 6.5 years after the transfusion.

**Detection of PrPsc**

To be truly useful, prion diagnostics should identify “suspect” cases as early during pathogenesis as possible. However, the currently available methods are quite insensitive when compared with those available for other infectious diseases. PrPsc represents the only disease-specific macromolecule identified to date, and all approved commercial testing procedures are based on the immunological detection of PrPsc. While around 50 companies are reported to be developing prion diagnostic assays, all commercial test kits validated for use by the European Union rely on proteolytic removal of endogenous PrP prior to detection of PrPsc (Table 2). In addition, the conformation-dependent immunoassay (4) utilizes the differential binding of antibodies to native or denatured PrPsc.

Circumvention of the protease digestion step might theoretically yield increased sensitivity of PrPsc-based detection methods and make these methods more amenable to high-throughput technologies. However, it has proved difficult to discriminate between PrP and PrPsc with antibodies, despite some early reports (51). Interestingly, tyrosine-tyrosine-arginine (YYP) motifs (52) were reasoned to be more solvent-accessible in the pathological isoform of PrP, and a monoclonal antibody directed against these motifs was reported to be capable of selectively detecting PrPsc across a variety of platforms. However, YYR motifs are certainly not unique to pathological prion proteins, and it remains to be determined whether this reagent can really improve the sensitivity of detection of prion infections.

Deposition of PrPsc in lymphoid tissues of human prion disease patients has long been believed to be restricted to vCJD. However, recent results (12) imply that PrPsc is present in spleens and muscle tissue from as much as one third of patients with sCJD. It is presently unclear whether the patients with extraneural PrP represent a specific subset of CJD patients or whether the extraneural-negative patients may simply deposit PrPsc in muscle and spleen at levels that are below the detectability threshold of the assay. If the latter scenario proves true, and if the assay sensitivity can be raised, minimally invasive muscle biopsies may replace brain biopsy in clinical CJD diagnostics.

**Surrogate markers and prion infectivity**

While presence of PrPsc secures diagnostic association with the presence of prion disease, PrPsc is not always easily detectable in several forms of prion disease (53–55). In order to enhance the safety of the blood supply and of products of bovine origin, absolute specificity in securing diagnosis of asymptomatic prion disease may not be required. Instead, it may be prudent to accommodate less than 100% specificity with a panel of surrogate markers capable of identifying donated blood units from “suspect” individuals rather than requiring definitive diagnosis. It could be envisaged that wide-scale primary screens accommodate a certain degree of loss of specificity to identify samples to be re-tested in a secondary screen utilizing more specific (and likely labor-intensive) criteria.

Several research efforts have been directed at identifying transcripts and proteins differentially expressed in tissues of prion-infected animals relative to disease-free controls (56–58). However, these have mostly focused primarily either on prion-infected neural cell lines or on CNS tissue, frequently with emphasis on late-stage disease. Ideally, these surrogate markers should be detectable (and differentially expressed) in easily accessible body fluids, such as blood or urine. At present, only one extraneural gene was reported to be differentially expressed during prion infection (59): erythroid differentiation–related factor (EDRF; also known as erythroid-associated factor) levels were progressively reduced in spleens of prion-infected mice throughout pathogenesis and also in blood of experimentally infected mice, cattle with BSE, and sheep with clinically manifest scrapie.

Assessment of the levels of surrogate markers in healthy individuals is crucial in order to define the normal range of expression (according to age, sex, etc.) in order to determine what represents abnormal levels. In this respect, it is worth noting that determination of normal expression range must utilize appropriate controls. For example, EDRF transcript levels have recently been reported to show a broad range of variation in healthy human subjects (60). However, since EDRF is an erythroid-specific transcript, it would be imperative to utilize other erythroid transcripts as internal controls to normalize for variations in numbers of circulating cells in which the transcript under study is expressed relative to total cells. More searches for surrogate markers would certainly be useful, and it is likely that surrogate markers of prion disease, particularly if they are detectable in body fluids, will expand the panel of tools available for screening for prion infections.

It is also worth noting here that recent advances in neuroimaging techniques, particularly with respect to MRI, may lead to clinically useful methods of assessment of prion disease in humans, perhaps even the ability to distinguish between sCJD and vCJD (61). For example, in vCJD the pulvinar sign (a high T2 MRI signal in the posterior thalamus) has been suggested to be relatively specific for vCJD, being present in approximately 75% of vCJD patients tested (62). In sCJD, fluid-attenuated inversion recovery and diffusion-weighted MRI sequences appear to be associated with high sensitivity and specificity. MRI imaging techniques such as these may therefore represent a relatively noninvasive method to corroborate suspicion of clinical presentation of human prion disease.

While surrogate markers such as S-100, neuron-specific enolase, and 14-3-3 protein have been suggested as potential biomarkers of prion disease using body fluids such as cerebrospinal fluid (CSF) (63, 64), it is worth remembering that these are clearly surrogate markers of general neurodegenerative disease and are not therefore predictive for human prion disease. For example, one study reported false positives of 14-3-3 detection in CSF samples of patients with herpes simplex encephalitis, hypoxic brain damage, atypical encephalitis, intracerebral metastases of a bronchial carcinoma, and metabolic encephalopathy (65).

It should not be forgotten that there is no ultimate consensus on the nature of the prion: PrPsc itself might represent a surrogate marker of prion disease (66). The real gold standard of prion diagnostics is the detection of prion infectivity (whether or not PrPsc is present). Until recently, the only method available to assay for prion infectivity was the use of the mouse bioassay, in which serial dilutions of test material are inoculated into mice and onset of disease noted. However, this procedure suffers from inaccuracy and is limited by the requirements for scores of mice and significant lengths of time. Recently, the use of highly susceptible cloned neural cell lines has provided what appears to be an assay that delivers a substantial reduction in both cost and time required to perform prion bioassays and may lend itself to high-throughput automation (67). Such assays may advance methodologies aimed at diagnostic assessment of the presence of...
the prion agent. However, it should be noted that these cell lines are currently reported only to be permissive to murine prions. It is to be expected that the spectrum of prion strains that can be assayed using this technology will expand.

Prion therapy

For all the promising approaches that are being explored (Table 3), no therapy for prion diseases is available as of yet. Many substances appear to possess prion-curing properties in vitro, including Congo red (68), amphotericin B, anthracyclines (69), sulfated polyanions (70), porphyrins (71), branched polyamines (72), -sheet breakers (73), the spice curcumin (74), and recently even small interfering RNAs (75). The majority of these molecules exert their biological effects by directly or indirectly interfering with conversion of PrPC to PrPSc, thereby also aiding clearance of PrPSc. Yet none of these compounds have proved very effective for actual therapy.

In a recent report, results obtained in mice have led to the theory that administration of cytidyl-guanyl oligodeoxynucleotides (CpG-ODNs), which stimulate the innate immune system via toll-like receptor 9 (TLR9) signaling receptors on a variety of immune cells, may represent an applicable treatment regimen to delay prion disease in humans (76). Here it was shown that the incubation period of prion disease was extended in mice multidose treated with CpG-ODNs for twenty days. It was concluded that stimulation of innate immunity accounts for this apparent anti-prion effect, possibly through induction of anti-PrP antibodies. However, this is difficult to reconcile with several studies indicating that immune deficiencies of various sorts inhibit prion pathogenesis (24, 25, 30, 77).

Do any serious candidates for prion therapeutic strategies exist? It is well established that expression of two PrPSc moieties that differ subtly from each other are able to inhibit prion replication (10). For example, humans heterozygous for a common PRNP polymorphism at codon 129 are largely protected from CJD (86). Although the precise molecular basis for this effect is unclear, it is possible that heterologous PrPC may exert inhibitory action on prion replication by sequestration. This has been addressed directly by fusion of PrPC to an immunoglobulin Fc domain (87), allowing fusion proteins either antagonizing soluble prion protein or depleting mature FDCs have been shown to efficiently prolong the incubation time of infected animals.

Is vaccination against prion disease possible?

Anti-PrP antibodies (30) and F(ab)2 fragments to PrP (80, 81) can suppress prion replication in cultured cells. However, the mammalian immune system is essentially tolerant to PrPC (82). Ablation of Prnp (39) renders mice highly susceptible to immunization with prions (22). Tolerance can be circumvented by transgenic expression of an immunoglobulin μ chain containing the epitope-interacting region of a high-affinity anti-PrP monoclonal antibody. This sufficed to block prion pathogenesis upon intraperitoneal prion inoculation (83). Passive immunization may be a useful strategy for prophylaxis of prion diseases, since it has been shown that passive transfer of anti-PrP monoclonal antibodies prior to prion inoculation (83). Passive immunization may be a useful strategy for prophylaxis of prion diseases, since it has been shown that passive transfer of anti-PrP monoclonal antibodies prior to the onset of clinical symptoms is able to delay the onset of prion disease in mice inoculated intraperitoneally (84). Unfortunately, several efforts aimed at active immunization strategies have met with little success due to the robust immune tolerance to PrPC. In this respect, it is certainly worth noting that extensive neuronal apoptosis in hippocampus and cerebellum has been shown following intracranial delivery of monoclonal antibodies reactive against a subset of PrP epitopes (85). The implications here are obvious; clearly, exhaustive in-vivo safety trials must be performed prior to the utilization of such strategies in humans.

Soluble prionostatic candidates

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Table 3

<table>
<thead>
<tr>
<th>Therapeutic approach</th>
<th>Target</th>
<th>Properties</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Polyanions</td>
<td>Possibly membrane-resident PrPSc</td>
<td>Efficient in cultured cells; relatively toxic in vivo</td>
<td>(70, 88)</td>
</tr>
<tr>
<td>Curcumin</td>
<td>Unknown FDCs</td>
<td>Efficacy in vivo unproven</td>
<td>(74)</td>
</tr>
<tr>
<td>Soluble lymphotxin receptor</td>
<td>Unknown</td>
<td>Effective in vivo, but only on peripheral pathogenesis; moderate untoward effects</td>
<td>(28, 29)</td>
</tr>
<tr>
<td>CpG oligodeoxynucleotides</td>
<td>FDCs, DCs; B cells and macrophages</td>
<td>Severely immunoclastic at doses effective in vivo</td>
<td>(76, 79)</td>
</tr>
<tr>
<td>Anti-PrP antibodies</td>
<td>PrPC</td>
<td>Effective in vivo only if administered in massive doses</td>
<td>(80, 83, 84)</td>
</tr>
<tr>
<td>Amyloidotrophic intercalators (e.g., Congo red)</td>
<td>PrPSc</td>
<td>Toxic; questionable efficacy in vivo</td>
<td>(89, 90)</td>
</tr>
<tr>
<td>Chemical or immunological sympathectomy</td>
<td>Peripheral nerves involved in neuroinvasion</td>
<td>Very efficacious, but unacceptable toxicity in vivo</td>
<td>(33)</td>
</tr>
<tr>
<td>Polyene antibiotics</td>
<td>Unknown</td>
<td>Low efficacy in vivo</td>
<td>(91, 92)</td>
</tr>
<tr>
<td>Chlorpromazine and quinacrine</td>
<td>Unknown</td>
<td>Questionable efficacy in vivo; hepatotoxicity</td>
<td>(93–95)</td>
</tr>
<tr>
<td>Soluble dimeric PrPC immunoadhesin</td>
<td>PrPSc</td>
<td>Effective as transgene, but efficacy upon injection unproven</td>
<td>(87)</td>
</tr>
</tbody>
</table>

While most substances investigated so far may possess prion-curing potential in vitro, no effective therapeutic substance has been identified so far for actual in-vivo therapy in humans. However, fusion proteins either antagonizing soluble prion protein or depleting mature FDCs have been shown to efficiently prolong the incubation time of infected animals.
Prion diseases continue to present a diagnostic and therapeutic challenge to clinicians and researchers worldwide. There are many aspects of prion biology that remain unclear; we still do not know the precise physical nature of the infectious agent, the molecular and biochemical mechanisms underlying associated neurodegeneration, or the physiological function of PrPSC. The diagnostic tools currently available for prion diseases are significantly less sensitive and satisfactory than those available for other infectious diseases. Additionally, there is a dearth of therapeutic intervention strategies available for these diseases. However, that said, the last decade or so of prion research has witnessed astounding advances in our knowledge and understanding of basic prion biology, and the field has attracted increasing numbers of researchers from diverse disciplines. Undoubtedly, this trend will trigger further important advances in prion science.

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