The human prion diseases, or transmissible spongiform encephalopathies, have captivated our imaginations since their discovery in the Fore linguistic group in Papua New Guinea in the 1950s. The mysterious and poorly understood “infectious protein” has become somewhat of a household name in many regions across the globe. From bovine spongiform encephalopathy (BSE), commonly identified as mad cow disease, to endocannibalism, media outlets have capitalized on these devastatingly fatal neurological conditions. Interestingly, since their discovery, there have been more than 492 incidents of iatrogenic transmission of prion diseases, largely resulting from prion-contaminated growth hormone and dura mater grafts. Although fewer than 9 cases of probable iatrogenic neurosurgical cases of Creutzfeldt-Jakob disease (CJD) have been reported worldwide, the likelihood of some missed cases and the potential for prion transmission by neurosurgery create considerable concern. Laboratory studies indicate that standard decontamination and sterilization procedures may be insufficient to completely remove infectivity from prion-contaminated instruments. In this unfortunate event, the instruments may transmit the prion disease to others. Much caution therefore should be taken in the absence of strong evidence against the presence of a prion disease in a neurosurgical patient. While the Centers for Disease Control and Prevention (CDC) and World Health Organization (WHO) have devised risk assessment and decontamination protocols for the prevention of iatrogenic transmission of the prion diseases, incidents of possible exposure to prions have unfortunately occurred in the United States. In this article, the authors outline the historical discoveries that led from kuru to the identification and isolation of the pathological prion proteins in addition to providing a brief description of human prion diseases and iatrogenic forms of CJD, a brief history of prion disease nosocomial transmission, and a summary of the CDC and WHO guidelines for prevention of prion disease transmission and decontamination of prion-contaminated neurosurgical instruments.
ons via the surgical instruments, as neural tissue presents the highest infectious burden of the disease.\(^8\)

Highly sensitive and specific diagnostic tests using cerebrospinal fluid and/or nasal brushings are becoming available at the National Prion Disease Pathology Surveillance Center (Cleveland, Ohio).\(^70,71\) Nevertheless, a definitive antemortem diagnosis of prion disease can only be made by tissue biopsy. The current difficulties in identifying prion-infected living patients constitute one of several challenges faced by institutions when determining whether specific prion decontamination measures should be taken during a neurosurgical procedure. In addition, the inability to identify patients who have been recently infected with the prion agent complicates responses to inadvertent exposures of surgical patients to potentially prion-contaminated instruments. This issue may arise when a patient who previously underwent a neurosurgical procedure receives a postoperative diagnosis of prion disease. Assessing the risk to those potentially exposed to contaminated instruments and making decisions related to informing such patients and preventing further exposures to the instruments can be difficult, time consuming, and costly. Fortunately, only a very small fraction of the total number of prion disease cases reported to date resulted from iatrogenic transmission; the social costs of iatrogenic transmission must also be considered; such costs include loss of public trust in medical personnel and institutions, investigative costs, and potential lawsuits.

This review seeks to provide an update on the existing information on the transmission of human prion diseases. A general background of the historical, epidemiological, pathobiological, and clinical aspects of the prion diseases

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### Table 1. Clinical and histopathological features of the human prion diseases

<table>
<thead>
<tr>
<th>Etiology &amp; Disease</th>
<th>Age at Onset (yrs)</th>
<th>Disease Duration (mos)</th>
<th>Presenting Clinical Sx/Signs</th>
<th>Neuropathology</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acquired</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Iatrogenic* CJD</td>
<td>Mean 58, SD 15</td>
<td>Mean 15.8, SD 9.2(^{16})</td>
<td>Typically w/ gait abnormalities &amp; ataxia</td>
<td>Spongiform degen, giosis, neuronal loss; ~68% of cases show amyloid plaques</td>
</tr>
<tr>
<td>Variant CJD</td>
<td>Median 26(^{11})</td>
<td>Median 14, range 6–39(^{11})</td>
<td>Psychiatric/behavioral Sx, paresthesia or dysthesia, delayed development of neurological signs</td>
<td>Numerous amyloid plaques surrounded by vacuoles (“florid plaques”), spongiform degen most evident in basal ganglia &amp; thalamus</td>
</tr>
<tr>
<td>Kuru</td>
<td>Range ~5 to &gt;50(^{24,59})</td>
<td>Range ~3–36(^{59})</td>
<td>Progressive cerebellar ataxia, no cognitive change</td>
<td>Kuru plaques (greatest frequency in cerebellum), neuronal loss, &amp; astrocyte hypertrophy</td>
</tr>
<tr>
<td>Inherited</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Familial CJD‡</td>
<td>Mean 58, range 35–66(^{37})</td>
<td>Mean 6, range 2–41(^{37})</td>
<td>Dementia, psychiatric changes &amp; ataxia; myoclonus; rarely gaze palsies &amp; neuropathy</td>
<td>Spongiform degen, giosis, &amp; neuronal loss w/ severity as function of disease duration</td>
</tr>
<tr>
<td>FFI</td>
<td>Mean 49, range 20–71(^{37})</td>
<td>Mean 11, SD 4 in 129 M/M; mean, 23, SD 19 in 129 M/V; range 6–33(^{37})</td>
<td>Sleep disturbances &amp; autonomic dysfunction</td>
<td>Neuronal loss &amp; mild giosis (predominantly in thalamus), rare spongiform degen or plaques</td>
</tr>
<tr>
<td>GSS§</td>
<td>Range 30–62(^{55})</td>
<td>Range 1–120(^{55})</td>
<td>Cerebellar abnormalities, dysthesia, hyporeflexia, proximal leg weakness</td>
<td>Variable; spongiform degen, giosis, &amp; kuru plaques vary in location &amp; severity, neurofibrillary tangles also evident</td>
</tr>
<tr>
<td>Sporadic</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sporadic CJD</td>
<td>Mean 64,(^{79}) range 19–91(^{55})</td>
<td>Mean 8,(^{79}) range 1–72(^{55})</td>
<td>Dementia, myoclonus, cerebellar dysfunction</td>
<td>Spongiform degen, giosis, neuronal loss; amyloid plaques in sCJD-MV2 subtype</td>
</tr>
<tr>
<td>Sporadic FFI</td>
<td>Median 46,(^{79}) SD13,(^{79}) range 24–74(^{55})</td>
<td>Median 24,(^{79}) SD 13,(^{79}) range 10–73(^{55})</td>
<td>Heterogeneous, including dementia &amp; ataxia; psychiatric &amp; visual Sx are less common; sleep disturbances at early stages of disease are often not investigated</td>
<td>Gliosis &amp; neuronal loss involving medial dorsal &amp; anterior ventral thalamic nuclei &amp; inferior olive; spongiform degeneration is minimal &amp; focal</td>
</tr>
<tr>
<td>VPSPr</td>
<td>Median 70,(^{79}) SD 9,(^{79}) range 48–87(^{55})</td>
<td>Median 24,(^{79}) SD 10,(^{79}) range 7–73(^{55})</td>
<td>Psychiatric signs, speech impairment, &amp; dementia</td>
<td>Spongiform degen w/ different size of vacuoles, microplaques in cerebellar molecular layer; round &amp; loose clusters of coarse PrP granules in cerebrum</td>
</tr>
</tbody>
</table>

**Notes:**
- Degen = degenerative; FFI = fatal familial insomnia; FI = fatal insomnia; GSS = Gerstmann-Schäussler-Scheinker syndrome; M/V = methionine/valine heterozygosity.
- Sx = symptoms; VPSPr = variably protease-sensitive prionopathy.
- * Dura mater graft–associated CJD.
- † Unicentric round plaque with a dense eosinophilic core and radiating spicules.
- ‡ Referred to fCJD.
- § Referred to GSS P102L-129M mutation.
- \(^{*}\) Referred to GSS P102L-129M mutation.

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D. J. Bonda et al.
is first provided, followed by descriptions of previous incidents and corresponding recommendations by the Centers for Disease Control and Prevention (CDC) and World Health Organization (WHO). This review is intended to disseminate information on how iatrogenic transmission of prion disease can be avoided in the hospital setting.

**Human Prion Diseases: Historical Perspective**

Human prion disease first came to the attention of the global scientific community during the 1950s when D. Carleton Gajdusek, a US physician and medical researcher, was called to investigate a “strange, encephalitis-like disease” occurring predominately among women and children of the Fore linguistic region of Papua New Guinea. Gajdusek and his colleague V. Zigas, a medical officer in the Fore tribe region of New Guinea, lived among the Fore and investigated this major medical problem. In 1957, Klatzo wrote: “[kuru] seems to be defined as a nervous muscular movement, etc. and which initiate as a startle response. If well and firmly supported, given passively by an examiner to head or upper extremities, suddenly sets off repetitive, irregular tremors or a choreiform pattern of movement. Rigidity is minimal, if at all present. It appears late. Instead, there is an increased tone to the muscles that are associated with attempts at maintaining posture and preventing the antigravity tremors which fight the slightest instability of standing, sitting, lying, head posture, etc. and which initiate as a startle response. If well and firmly supported, even in late cases, this “intermittent rigidity” subsides to complete relaxation.”

Gajdusek later noted the similarity of kuru to “heterofamilial degenerative disorders of the central nervous system,” whereas Igor Klatzo, neuropathologist at NIH, compared kuru to Creutzfeldt-Jakob disease (CJD), the latter being documented only 20 times and never observed in children. In 1957, Klatzo wrote: “[kuru] seems to be definitely a new condition without anything similar described in the literature. The closest condition that I can think of is that described by Jacob and Creutzfeldt.” Klato detected the vacuoles and the amyloidal kuru plaques in the brain of individuals with kuru, and, in 1959, he published a paper describing kuru’s histopathological features and the similarities between kuru and CJD. In 1966, Gajdusek, Gibbs, and Alpers experimentally transmitted kuru to chimpanzees, confirming that kuru was an infectious condition.

Two years later, CJD was shown to be transmissible. Gibbs and Gajdusek proposed the name of “subacute spongiform encephalopathy” for the entire class of scrapie-like diseases. The pathogenesis of the transmissible spongiform encephalopathies, however, remained a mystery, the solution to which would have an enormous impact on medicine.

**Kuru and Cannibalism**

Epidemiological and clinical research conducted by Zigas and Gajdusek revealed a complicated picture. According to their 1959 report, kuru had the following properties: 1) It was responsible for the death of approximately 1% of the population per year (up to 50% in certain areas). 2) It mainly affected adult females, with up to 25% of cases involving children. 3) It was restricted to the Fore people and some of their tribal neighbors. 4) Its histopathological hallmarks included neurological degeneration, myelin degeneration, astroglial and microglial proliferation, most predominately in the cerebellum and extrapyramidal system. 5) Cases of kuru had no identifiable relationship with cases of (presumably viral) infectious meningoencephalitis or with any toxic ingestion.

Gajdusek himself postulated a genetic predisposition to the spread of kuru; however, little evidence could be found to support such a hypothesis. Currently, however, we understand that heterozygosity at codon 129 of the prion protein (PrP) gene (PRNP) is protective (see below).

Remarkably, the kuru-affected people of Papua New Guinea were unique in their practice of “endocannibalism.” As Alpers describes, the people of the Fore region believed the “mortuary practice of consumption of the dead and incorporation of the body of the dead person into the bodies of living relatives, thus [helped] free the spirit of the dead.” In fact, the whole body of the deceased kin was eaten by female relatives and children of both sexes. Adult males, which included boys above age 7, rarely took part in this practice. The sex and age distribution of the kuru epidemic could thus be satisfactorily explained by the mortuary practices of the affected people. In fact, after the abandonment of the practice by the early 1960s (following the urging of the Australian administration), there was a sharp decline in cases that was characterized by the long incubation period of the disease.

The ensuing decades of clinical and basic research revealed the etiologic agent of kuru to be a small, self-aggregating protein unlike any infectious entity previously identified in biology. The agent, now known as “prion,” introduced to the scientific community principles of molecular catalytic activity that would influence the fields of molecular biology, genetics, virology, amyloidology, and aging. Such was the impact of the elucidation of the pathophysiology of kuru that D. Carleton Gajdusek was awarded the Nobel Prize in Physiology or Medicine in 1976 for his work on the transmission of the disease. Stanley Prusiner, whose work is described below, also received the Nobel Prize in 1997 for his work on the molecular identification and characterization of the prion protein. The detailed findings of the investigation into kuru and...
the discovery and development of the concept of transmissible prion proteins were summarized by Dr. Gajdusek and Dr. Prusiner, respectively, in *Fields' Fundamentals of Virology*, published in 1996,31,78

**Human Prion Diseases in the 21st Century**

The term “prion” was coined in 1982 by Stanley B. Prusiner to describe “proteinaceous infectious” particles responsible for scrapie in goats and sheep.77 Since their initial elucidation, much work has been done to understand and categorize these fatal neurodegenerative conditions. Prion diseases have different etiologies: they can arise sporadically, be genetically inherited, or acquired by infection.

The normal or cellular PrP (PrPc) is a glycosylphosphatidylinositol-anchored membrane glycoprotein that is largely expressed in neural and nonneural tissues and found primarily on the cell surface—plasma membrane of the central nervous system. PrPc is encoded by the PRNP gene on chromosome 20 and comprises 209 amino acids that fold to produce an α-helix–rich conformation that is soluble in buffers containing detergent and readily digestible by proteases (e.g., by the proteinase K).48,77,80 The function of PrPc is largely unknown.50,86

The abnormal or pathological form of PrP (PrPSc) displays a predominantly β-sheet conformation with a C-terminal region that is partially resistant to proteolytic degradation.7,22,35 The conformational conversion of PrPc to PrPSc seems to occur in a reaction whereby α-helical structures of PrPc refold into a β-sheets structure using a preexisting PrPSc as a template.1,27,40,49 Although the accumulation of the proteinase K–resistant PrPSc, a gold standard marker for prion disease, may not always be observed with standard detection procedures in a few human prion diseases, a central pathogenic event is the accumulation of PrPSc that is partially resistant to proteases.84

Polymorphism at codon 129 of the PRNP gene, encoding for either methionine (M) or valine (V), has been demonstrated to play a role in host susceptibility to phenotypic expression of sporadic, familial, and acquired or iatrogenic forms of prion disease.23,23,74 Codon 129 M/V heterozygosity seems to be protective against human prion diseases,93 and several studies have indicated a prominence of homozygosity for either methionine (129 M/M) or valine (129 V/V) in individuals with prion disease.23,52,63 Although the biophysical interplay between codon 129 and PrPSc conversion is incompletely understood, the epidemiological association is of considerable clinical predictive value.

There is a great deal of phenotypic heterogeneity in the prion diseases. The clinicopathological phenotype in CJD and other prion diseases is also influenced by the different types of the pathological PrPSc, identified as Type 1 and Type 2.66,72,25

**Creutzfeldt-Jakob Disease**

The most common human prion disease is Creutzfeldt-Jakob disease (CJD), with an estimated incidence of 1–1.5 cases per million people per year.8 Approximately 85% of all CJD cases are sporadic (sCJD) and considered to arise from somatic alteration in PrPc.20,35 A modern classification of sCJD into 5 distinct subtypes combines 2 types of PrPSc (Type 1 and Type 2) and 3 possible genotypes at codon 129 (129 M/M, 129 M/V, and 129 V/V). Each subtype of sCJD is characterized by a distinct clinical and histopathological phenotype.19,34 Genetic or familial CJD (fCJD) represents 5%–15% of all CJD cases89 and is associated with several pathogenic mutations in the PRNP gene. Patients with fCJD are usually younger than those with sCJD. Clinically, fCJD presents with rapidly progressive neurological and neuropsychiatric dysfunction, including dementia, visual abnormalities, muscle incoordination and myoclonus, and gait and speech abnormalities (Table 1). The rates of progression and symptoms at onset vary depending on the sCJD subtype. In about 85%–90% of cases, however, patients deteriorate rapidly, with death occurring within 12 months of the onset of illness.8 Variants CJD (vCJD) was initially reported in 1996 as a small case series of CJD-like illnesses in the United Kingdom that was epidemiologically linked to an outbreak of bovine spongiform encephalopathy (BSE). Affected patients exhibited early-onset disease (median age 28 years) with prominent behavioral changes at clinical presentation followed by neurological abnormalities, dementia, and myoclonus later in the course of the illness.91 Epidemiological and laboratory studies indicated that the same prion agent was responsible for BSE and vCJD.46,64 Iatrogenic transmission of vCJD has been linked to blood products (3 clinical cases and 2 subclinical cases)5,82 rendering iatrogenic transmission of prion diseases a greater potential problem.

**Fatal Insomnia**

Sporadic fatal insomnia (sFI) and fatal familial insomnia (FFI) are characterized by bilateral symmetric degeneration of the thalamus with marked gliosis and absence of minimal spongiform degeneration.58 Patients predominantly exhibit severe sleep disturbances, often with intractable insomnia, and autonomic dysfunction, characterized by hyperhidrosis, hyperthermia, tachycardia, and hypertension.90 It is most commonly associated with PRNP gene mutation at codon 178 (FFI), but sporadic cases have been identified lacking such mutation (sFI).90,73

**Kuru**

The clinical picture of kuru, as described above, is considerably distinct from that of classic CJD. Cases occurring 50 years after participation in ritual cannibalism have been reported,24 suggesting that measuring the full risk of person-to-person transmissions of prion disease can be challenging due to possibly decades-long incubation periods.

**Gerstmann-Sträussler-Scheinker Syndrome**

Gerstmann-Sträussler-Scheinker syndrome (GSS) is a slowly progressive hereditary cerebellar syndrome associated with PRNP point mutations at different codons.60 The incidence of GSS is estimated at approximately 1–10 cases per 100 million people per year.29 Typically reported neurological symptoms include cerebellar ataxia, gait abnormalities, dementia, dysarthria, ocular dysmetria, and hyporeflexia or areflexia in the lower extremities. As with
sCJD, however, the different mutations confer different clinical and histopathological phenotypes.

**Varially Protease-Sensitive Prionopathy**

Varially protease-sensitive prionopathy (VPSPr) is a recently identified prion disease affecting an estimated 2–3 people per 100 million per year. Clinical onset is characterized by psychosis, mood changes, speech impairment, and dementia, whereas progressive motor dysfunction is usually observed at later stages of the disease. About 30% of the affected individuals have a family history of dementia. The median age of disease onset, and the disease duration, is 70 years and 2 years, respectively. In VPSPr, disease prevalence in association with the 3 codon 129 genotypes of the PRNP gene (i.e., M/M, M/V, and V/V) is different from that of sCJD, suggesting a different role of codon 129 as a risk factor in the 2 conditions (VPSPr: 62% V/V, 26% M/V, and 12% M/M; sCJD: 19% V/V, 11% M/V, and 70% M/M). Other features of VPSPr are different sensitivity of PrP^Sc^ to proteolytic digestion with protease K (hence “varially protease-sensitive”), and poor transmission of the disease to transgenic mice expressing the human PrP<sup>Sc</sup>.69

**Iatrogenic CJD: Historical Context**

Iatrogenic CJD (iCJD) refers to the transmission of prions via inadvertent medical exposure. The first documented case of iCJD occurred via infected corneal transplant, and was described by Duffy in 1974. Three earlier iatrogenic prion transmissions by surgical instruments were suggested by examination of case notes of CJD cases described in a report by Nevins et al. in 1960. A fourth case of iatrogenic CJD attributed to a contaminated neurosurgical instrument was described by el Hachimi and colleagues in 1977; the case involved a 46-year-old man who had undergone cranial surgery in the same department 3 days after a cortical biopsy had confirmed CJD in a 59-year-old woman. In 1977, Christoph Bernoulli realized that a cortical electrode probe used in an elderly patient had transmitted CJD to 2 younger patients undergoing subsequent epilepsy surgery. Subsequent reports demonstrated the transmission of CJD to the frontal lobes of chimpanzees with the same electrodes, even after scrupulous attempts to clean them.

More than 492 (personal communication to L.B.S., 2015) cases of iCJD have been identified worldwide (Table 2). The majority of such cases resulted from the administration of prion-contaminated human growth hormone (hGH) and cadaveric dura mater graft. Other sources of iatrogenic transmission of the disease are contaminated blood, corneas, and neurosurgical instrumentation. Because of the latter risk, Drs. Paul Brown and Michael Farrell recently proposed routine use of prion diagnostic testing on all patients admitted with symptoms of either dementia or cerebellar disease and containment of infectivity commensurate with the degree of potential risk.16

**Cadaveric Dura Mater Transplantation**

The transmission of CJD by transplantation of commercially distributed cadaveric dura mater (dCJD) was first recognized in 1987. Since then, at least 238 dCJD cases (personal communication to L.B.S., 2015) have been identified worldwide, with over 60% occurring in Japan, reflecting primarily the frequent use of Lyodura. Lyodura was produced by the German manufacturer B. Braun Melssungen AG and was the major source of the outbreak. In 1987, the manufacturer reported that it changed its procedures to reduce the risk of prion contamination of its product. The mean incubation period for exposed dCJD patients has been estimated to be 12 years (range 1.2–30 years), although cases with even longer incubation periods are likely to occur. The predominant symptomatology has been atypical (slowly progressive and without characteristic electroencephalography [EEG] findings). In Japan, 2 groups of dCJD with distinct phenotypes and molecular features have been recently described.53

**Human Growth Hormone Administration**

The treatment of short stature with pituitary-derived human growth hormone (hGH) began in the 1950s and has been associated with more than 238 iCJD cases to date (personal communication to L.B.S., 2015). The majority of hGH-CJD cases have occurred in France, specifically among patients who received hGH treatment between December 1983 and July 1985 (119 cases in 1170

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**Note:**

The data presented in Table 2 is adapted and updated from Brown et al., Will & Matthews, and personal communication to L.B.S., 2015. The table illustrates the worldwide incidence of iatrogenic transmission of prion infectivity, with specific cases and outcomes such as surgical procedures, medical procedures, and blood transfusions. The table also highlights the clinical findings associated with these cases, including psychiatric, sensory, and dementia symptoms. Further details on the incidence and characteristics of these cases are provided in the text, along with the historical context and implications for future research and practice.
exposed patients). In the UK and the US, the numbers of cases are fewer (75 and 31, respectively). Except for 1 case of CJD associated with commercially derived hGH, 30 of the 31 patients in the US cases received hGH treatment through the government-supported National Hormone and Pituitary Program. In this program, no CJD cases have been identified among recipients who began their hormone treatment after 1977, the year when a highly selective column chromatography step was included in the purification protocol. The mean incubation time for hGH-CJD patients worldwide has been estimated to be 17 years (range 5–42 years), and the associated clinical picture is predominantly cerebellar, with dementia occurring late in the development should it appear.

Blood Transfusion

The first convincing evidence for blood transfusion transmission of a human prion disease was reported for vCJD in 2003. As indicated in Table 2, 5 cases of vCJD infection, including 3 involving patients who became ill with vCJD, have been documented among recipients of blood products from donors who subsequently developed vCJD. (Table 2). Blood transfusion

Other Sources

Although the vast majority of iCJD cases have occurred because of contaminated hGH or dura mater graft, other identified sources include corneal transplantation and contaminated stereotactic EEG depth electrodes. In four cases, patients underwent corneal transplantation or EEG with grafts from or instruments previously used on patients in patients subsequently discovered to have died of confirmed CJD. The latency period for corneal transplant cases ranged from 18 months to 30 years, whereas the latency periods for the cases ascribed to the contaminated EEG depth electrodes were 16 months and 20 months.

Transmission of Human Prions by Neurosurgical Instruments

Iatrogenic transmission of CJD via neurosurgical instrumentation is a worrisome, although rare, phenomenon. Four cases have been documented in the literature, with three having occurred in the UK and one in France during the 1950s. Exposed patients presented back to the hospital with onset of CJD between 1.4 and 2.2 years after their surgery. Although there have been no documented cases in the US, the unusual pathology of one CJD case in a US neurosurgeon has been reported to suggest iatrogenic rather than sporadic disease. In addition, several potential exposures have been described.

In the 15-year period (1998–2012), 19 incidents of suspected CJD exposure via contaminated surgical instruments in the US were reported to the CDC. Two cases involved ophthalmological procedures, whereas 17 involved intracranial neurological operations that were performed as a diagnostic workup in most of the cases. Operative personnel were not aware that their patient was infected with the CJD agent, so no recommended CJD-related protocols for instrument tracking/decontamination were followed. The contaminated instruments were cycled through the normal decontamination process and reused on subsequent patients. Several hospitals reported having multiple neurosurgical instrument sets, and in most of these hospitals (11 of 19) the originally contaminated set could not be identified. The potential nosocomial exposure to the CJD agent raises several concerns. Most important is the prevention of further exposure of patients to CJD. This requires identification of potentially contaminated instruments and their proper quarantine and sterilization. As demonstrated by these cases, instrument tracking can be very difficult when there are multiple instrument sets and when a large amount of time has passed since the initial neurosurgery. In some of the above-mentioned hospitals, the entire collection of sets had to be quarantined and decontaminated. Instruments used for neurosurgery in patients with dementia or cerebellar signs for whom there is not convincing evidence against a prion disease should be decontaminated using the sterilization protocols developed by WHO.

A second important consideration is patient notification. The hospital should try to identify the patients potentially exposed to the index instruments (another epidemiological obstacle) and decide if anyone, appropriately, should be notified. In this regard, several factors should be taken into account.

Risk Assessment and Mitigation

Given the difficulty of tracking instruments and connecting them to patients, it is crucial to identify patients who have, or are at risk for having, a prion disease before they undergo surgical procedures. Successful implementation of the appropriate precautions can effectively eliminate the risk of subsequent iatrogenic events. Suggestions for mitigation are provided in the WHO Infection Control Guidelines for the Transmissible Spongiform Encephalopathies. According to these guidelines, prior to the operation, efforts should be made to minimize the extent of contamination. The WHO specifically suggests that all staff directly involved in the procedure or the reprocessing or disposal of the contaminated items be made aware of recommended precautions; operative and relevant staff must be given enough time to obtain suitable instrumentation and equipment; specific protocols are followed; and staff is appropriately trained for these protocols. It is also suggested that the procedure in question be scheduled for the end of the day (i.e., as the last case) to ensure adequate time for decontamination. These procedures may have a visible effect on the surgical instrument to which the procedure is applied.

The WHO also lists basic protective measures to be taken within the operating room: 1) involve a minimum number of health care personnel in the operating room; 2) use single-use equipment whenever possible; 3) cover all nondisposable equipment (Fig. 1); 4) maintain a “one-way flow” of instruments; 5) dispose of single-use items via incineration when possible; and 6) clean all work surface areas according to guidelines.

Prion Protein Resilience and Decontamination

The most commonly used methods for disinfection and sterilization may not be adequate to remove all prion in-
Neurosurgical transmission of prion diseases

Conclusions

Iatrogenic forms of CJD represent a unique challenge to neurosurgeons. Although their incidence is rare, and the likelihood of encountering them in surgical practice is low, a missed diagnosis of prion disease in a neurosurgical patient can severely and negatively impact patients and associated hospitals. It is therefore important to remain vigilant during preoperative workup, especially when the preprocedural differential diagnosis could include the suspicion of a prion disease. Although intensive, the recommended precautions can reduce the potential risk of nosocomial prion infections and minimize the negative consequences.

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References


FIG. 1. Photographs showing the method used for protecting nondisposable operating room equipment. To minimize surface exposure to prion-related contaminants, all nondisposable instrumentation should be covered in a disposable protective material. Following the operation, these disposable items should be incinerated.


60. Liberski PP, Surewicz WK: Molecular genetics of Gerst-


Disclosures

The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

Author Contributions

Conception and design: Bonda. Acquisition of data: Cali, Bonda, Puoti, Cohen, Schonberger. Analysis and interpretation of data: Cali, Bonda, Schonberger. Drafting the article: Cali, Bonda, Schonberger. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Cali. Administrative/technical/material support: Cali, Schonberger. Study supervision: Cali.

Correspondence

Ignao Cali, Department of Pathology, Case Western Reserve University, 2085 Adelbert Rd., Cleveland, OH 44106. email: ixc20@case.edu.