Creutzfeldt-Jakob disease: updated diagnostic criteria, treatment algorithm, and the utility of brain biopsy

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Creutzfeldt-Jakob disease (CJD) is a rare neurodegenerative condition with a rapid disease course and a mortality rate of 100%. Several forms of the disease have been described, and the most common is the sporadic type. The most challenging aspect of this disease is its diagnosis—the gold standard for definitive diagnosis is considered to be histopathological confirmation—but newer tests are providing means for an antemortem diagnosis in ways less invasive than brain biopsy. Imaging studies, electroencephalography, and biomarkers are used in conjunction with the clinical picture to try to make the diagnosis of CJD without brain tissue samples, and all of these are reviewed in this article. The current diagnostic criteria are limited; test sensitivity and specificity varies with the genetics of the disease as well as the clinical stage. Physicians may be unsure of all diagnostic testing available, and may order outdated tests or prematurely request a brain biopsy when the diagnostic workup is incomplete. The authors review CJD, discuss the role of brain biopsy in this patient population, provide a diagnostic pathway for the patient presenting with rapidly progressive dementia, and propose newer diagnostic criteria.

http://thejns.org/doi/abs10.3171/2015.8.FOCUS15328

KEY WORDS Creutzfeldt-Jakob disease; prion; brain biopsy; diagnostic criteria; diagnostic algorithm; National (Nationwide) Inpatient Sample

The term Creutzfeldt-Jakob disease (CJD) was first used in 1922 by Spielmeyer to name the condition described by 2 German physicians. The 6 original subjects (a case report by Creutzfeld and a series of 5 patients by Jakob) were a heterogeneous group of patients with unusual neuropathological findings associated with other disorders, and years later only 2 of the cases were confirmed with modern techniques to actually be CJD. In 1960 the typical clinical picture, electroencephalography (EEG) findings, and classic spongiform changes in the neuropil were recognized as the cardinal features of the disease.

Now there are diagnostic criteria to guide the physician in the workup of a patient with suspected CJD; physical examination and tests are used to provide a definitive, probable, or possible diagnosis. A brain biopsy is able to give pathological confirmation of the disease and is considered the gold standard for diagnosis. However, research has progressed to the point where newer tests should be considered in the diagnostic criteria and may be more sensitive in disease confirmation than tissue obtained from surgery.

Creutzfeldt-Jakob disease is classified as familial, sporadic, or acquired. Regardless of the type, the disease has a rapid clinical course that is uniformly fatal. There are some consistencies on physical examination, radiographic studies, and EEG, but the most common form is sporadic CJD (sCJD), and it follows a theme of heterogeneity. The infectious agent is the abnormal scrapie form (PrPSc) of the host-encoded cellular prion protein (PrPC) that causes a posttranslational modification of PrPC into the disease form, accumulating in the brain and causing neurodegeneration. Familial CJD (fCJD), or the genetic type, is due to a mutation in the gene encoding PrPC, whereas the sporadic form is thought to originate after a somatic mutation or a stochastic protein alteration. Creutzfeldt-Jakob disease is

ABBREVIATIONS ADC = apparent diffusion coefficient; BSE = bovine spongiform encephalopathy; CDI = conformation-dependent immunocassay; CJD, fCJD, sCJD, vCJD = Creutzfeldt-Jakob disease, familial CJD, sporadic CJD, variant CJD; DWI = diffusion-weighted imaging; EEG = electroencephalography; FFI = fatal familial insomnia; GSS = Gerstmann-Sträussler-Scheinker; NIS = National (Nationwide) Inpatient Sample; PSWCs = periodic sharp wave complexes; RT-QuIC = real-time quaking-induced conversion; UK = United Kingdom.

SUBMITTED July 1, 2015. ACCEPTED August 6, 2015.

INCLUDE WHEN CITING DOI: 10.3171/2015.8.FOCUS15328.
also transmissible by iatrogenic causes or by ingesting beef with bovine spongiform encephalopathy (BSE or “mad cow disease”), leading to variant CJD (vCJD).25

Creutzfeld-Jakob disease is rare, and is often a diagnostic challenge for physicians facing a rapidly progressing dementia. As of now, definitive diagnosis is provided with brain biopsy, but more often than not the results are inconclusive because not all areas of the brain will show the classic histological changes in CJD, even if the disease is present. Surgeons target areas that appear the most abnormal on imaging studies, but this is most often in deep-seated subcortical structures. The WHO criteria, as well as more updated suggested criteria, are outdated because they do not use newer tests that provide a less invasive method for definitive diagnosis. Given these issues, a newer approach to diagnosis is needed. In this paper we briefly review the disease, assess the current tools used in making the diagnosis, discuss when a brain biopsy should be performed, propose changes to diagnostic criteria, and provide an algorithm for the workup of the patient with a rapidly progressive dementia of unknown origin.

Pathophysiology

To understand some of the clinical findings of the disease, one must first review the pathophysiology of how a healthy protein becomes abnormal and destructive to the brain. PrPC is found in lipid rafts on the cell surface of normal brains. The function of the protein is unknown—prion protein (PRNP) knockout mice completely lacking the protein do not show any obvious abnormalities, and have normal brain development.7

The central pathological event is the formation of the abnormal PrPSc from the wild-type, cellular form of PrPC. This is hypothesized to occur in a pathway where PrPSc serves as the template for PrPC to fold abnormally into the pathogenic conformation. This is an autocatalytic process that is poorly understood, but the change in protein shape is the hallmark of the pathology.44

Both forms have an identical amino acid sequence (primary structure), but the posttranslational changes cause the PrPC (40% alpha helix) to refold into a form with 45% beta-sheet composition.44 This makes the protein not only highly insoluble, but also resistant to proteinase digestion. The subsequent multimerization accumulates, spreads throughout the brain parenchyma, and induces the classic spongiform change (vacuolation of gray matter) by microglial activation and neuronal loss, leading to progressive neurodegeneration and astroglisis over time.

Genetics

PrPC is encoded by the prion protein (PRNP) gene on human chromosome 20.26 All familial forms of the disease are characterized by a mutation in this gene. PRNP has a normal genetic polymorphism at codon 129, where either methionine (M) or valine (V) may be encoded. This is very important because the genotype imparts genetic susceptibility in all types of prion disease. The most dramatic example of this is that all cases of vCJD to date are in individuals homozygous for methionine (MM).16

Parchi et al.45 classified sCJD into 6 different molecular strains based on the genotype and biochemical properties. When PrPSc is cleaved by proteinase K under defined conditions, there are 2 truncated forms that result: the 21-kD fragment (Type 1) and one that is 19 kD (Type 2). The importance of this classification into MM1, MM2, MV1, MV2, VV1, and VV2 is that the different strains strongly correlate with different pathologically distinct phenotypes, which is summarized in Table 1. The fact that PrPSc has the same amino acid sequence but different pathological and clinical presentation implies that the conformational variant is what determines the phenotypic or molecular “strain.”

The changes in protein structure are investigated by a conformation-dependent immunoassay (CDI). This test identifies PrPSc by exposing specific epitopes of the protein that are unmasked with progressive denaturation, and antibodies specific to these areas bind and elicit a positive result. It is 100% specific for the disease, and recent data show sensitivity that is at least equal to other diagnostic tests.

Forms of CJD

Sporadic CJD

By far the most common form of the disease is the sporadic type—sCJD—which occurs at an incidence of 1 case in 1 million per annum, and accounts for 85% of CJD cases. Researchers speculate that it results from a spontaneous neurodegenerative illness, and the hypothesis is that it results from either a somatic mutation in the gene or a random structural change in the PrP protein causing formation of PrPSc. Onset usually occurs in the 7th decade of life, and the median time to death is 5 months, with 90% of patients dead by 1 year.25 Unlike vCJD, the clinical and pathological findings are more heterogeneous, and this is probably due to the different molecular phenotypes present.

Familial CJD

Familial CJD is a result of known mutations of PRNP. This accounts for approximately 10% of all cases of prion diseases35 and is historically split into the 3 phenotypic categories of Gerstmann-Sträussler-Scheinker (GSS) syndrome, familial fatal insomnia (FFI), and fCJD. There are more than 50 mutations described, and the disease is transmitted in an autosomal dominant pattern with high penetrance, and with an incidence that increases with age. fCJD has similar clinical, radiographic, and test findings as sCJD, whereas GSS and FFI are well-described variants.35 A comparison of the familial types with sporadic and variant CJD is provided in Table 2.

Iatrogenic CJD

Creutzfeld-Jakob disease can be acquired iatrogenically; several outbreaks over the last few decades were associated with intracerebral electrodes, corneal transplantation, dura mater grafts, and growth hormone injections. The first case report, in 1974, was in a patient who received a corneal transplant from an infected cadaver.15 Dura mater grafts with contaminated material led to more than 60 cases of CJD, with incubation periods lasting be-
The clinical and pathological findings differ from other types of CJD (Table 2), in that the early course consists of mostly psychiatric symptoms before ataxia begins at approximately the 6-month mark. The median age of onset is much younger than in the sporadic or familial types, and the mean survival is also longer at 14 months. It is unclear if the survival increase is due to the age differences, but recent epidemiological studies do suggest that exposure alone is not sufficient to explain the higher incidence, and that age is an important risk factor for contracting the disease.4 Again, genetics play a crucial role; all confirmed cases were homozygous for methionine (MM) at the polymorphic codon 129 of the PRNP gene.

As with peripheral exposure to the infectious protein in acquired CJD, the incubation period is longer in vCJD given the parenteral route of infection. Most patients who developed the disease were exposed in the late 1980s, and the peak incidence of vCJD was in the early 2000s, giving an incubation period of 11–12 years. Another distinct feature is that PrPSc accumulates in germinal follicles and lymph nodes, and has been pathologically confirmed after tonsillectomy. This deposition of the abnormal protein in the lymphoreticular system is not seen in other human prion diseases.21

### Table 1. Genetic subtypes of sCJD and typical features

<table>
<thead>
<tr>
<th>Features</th>
<th>MM1/MV1</th>
<th>VV2</th>
<th>MV2</th>
<th>MM2</th>
<th>VV1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age at onset</td>
<td>70 yrs</td>
<td>65 yrs</td>
<td>60 yrs</td>
<td>67 yrs</td>
<td>44 yrs</td>
</tr>
<tr>
<td>Duration of illness</td>
<td>4 mos</td>
<td>6 mos</td>
<td>18 mos</td>
<td>14 mos</td>
<td>21 mos</td>
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<td>MRI findings</td>
<td>70% MRI hyperintensity in basal ganglia or cortex</td>
<td>70% hyperintensity in basal ganglia, 45% in thalamus</td>
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<td>EEG findings</td>
<td>PSWCs in 80%</td>
<td>PSWCs in 10%</td>
<td>Similar to VV2</td>
<td>PSWCs in 42%</td>
<td>PSWCs negative</td>
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<tr>
<td>14–3-3 status</td>
<td>95% positive</td>
<td>80% positive</td>
<td>Similar to VV2</td>
<td>91% positive</td>
<td>Positive in nearly all cases</td>
</tr>
<tr>
<td>Percentage of sCJD cases</td>
<td>60%–70%</td>
<td>Approximately 15%</td>
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<td>Approximately 5%</td>
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### Table 2. Findings in different types of CJD

<table>
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Between 1 and 14 years,10 pooled cadaveric growth hormone was used for injections and led to CJD 5–30 years after the injections. The clinical symptoms are similar to sCJD, as are the MRI and EEG findings.6 Iatrogenic CJD also has a clinical picture that is similar to sCJD.

It is clear that the incubation time is reflective of the inoculation site. Those with contaminated electrodes placed directly in the brain had short incubation periods of 16–28 months, whereas peripheral injections of growth hormone took anywhere from 5 to 30 years for the symptoms to begin.52 There are 3 cases of probable transmission of CJD to individuals who received blood transfusions from a donor with vCJD, which is why there is a ban on donors who lived in the United Kingdom (UK) during the epidemic of BSE.30

### Variant CJD

Variant CJD was first described in 1996 and is the result of eating food contaminated with BSE, which was a major problem in the late 1980s through the early 1990s in the UK.56 Since the original 1996 report, there have been a total of 229 confirmed cases worldwide, with the most being found in the UK. There have been 4 total cases in the US, with 3 of the 4 having a history of residence in the UK.32

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Pathological investigation shows the characteristic spongiform change with gliosis in the brain, predominantly in the basal ganglia and cerebellum. The most severe damage is in the pulvinar, which correlates with the characteristic high signal abnormality on T2 MRI studies that is seen in 75% of cases. In contrast to the familial and sporadic types, the EEG study is usually negative for any periodic sharp wave forms.21

**Molecular Strains**

The most common strain is the MM1/MV1. These two are combined because there are no differences in the pathological features, and there is similarity between the clinical symptoms. Between 60% and 70% of all CJD cases are grouped here, with 95% of these being MM1. These patients fall under the “classic” CJD criteria, with myoclonus, an age of onset in the 7th decade of life, and a short mean disease duration of 4 months. The most common presentation is cognitive impairment, but prominent visual signs (Heidenhain’s variant), cerebellar ataxia, and psychiatric symptoms can be present as well. The main difference between MM1 and MV1 is that in patients with MV1, ataxia is the more prominent neurological finding over cognitive impairment, especially early in the disease course. EEG is positive for periodic sharp wave complexes (PSWCs) in the first 3 months (80%), and 14–3-3 protein sensitivity is 95%.41 MRI shows changes in the striatum and cerebral cortex in 70% of cases.26 VV2 is the next most common; it is seen in approximately 15% of cases. There is a similar age of onset at 65 years and a slightly longer clinical course of 6 months, but EEG is nonspecific and rarely shows PSWCs (less than 10% of cases). MRI again shows hyperintensity in the striatum and thalamus, and the sensitivity of the 14–3-3 protein assay is 80%.41 This group can be distinguished on examination because of the predominance of rapidly progressive ataxia in the absence of myoclonus, which does not appear until late.

MV2 is present in 9% of the patient population and is distinguished from all other subtypes because of the significantly longer disease course. There is a mean age of onset at 60 years, but the duration of the disease is, on average, 18 months. This slower progression rate is in conjunction with prominent ataxia as well as the more typical cognitive decline, myoclonus, and psychiatric signs. The EEG and 14–3-3 results are similar to those in VV2, but the MRI shows thalamic signal most frequently in the pulvinar, leading to the described “pulvinar sign.”27

MM2 also has a longer disease course (14 months), and is seen in 2%–8% of cases. The main difference between MM2 and MV2 is that MM2 has cognitive impairment in MM2, as opposed to the predominant ataxia in the latter. EEG shows PSWCs in 42%, and MRI has increased cortical signal in 25%, usually seen in the temporal lobe and with little basal ganglia involvement.26,42

The final group, the VV1, is the most uncommon, being seen in just 1% of cases. This is an early-onset subtype, with the reported age of onset being 40–44 years, and a disease duration of 21 months. A prominent, slowly progressive frontotemporal dementia with psychiatric changes is described; MRI shows hyperintensity in the cortex without much basal ganglia involvement, EEG does not show PSWCs, but the test for 14–3-3 protein is almost always positive.36,42

**Clinical Signs**

The hallmark of CJD is rapidly progressive dementia of unknown origin. Also, numerous atypical neurological examination findings are commonly seen: myoclonus, visual changes leading to cortical blindness, ataxia, and usually an akinetic mutism in the last stages of the disease. Myoclonus is the most common sign, but there are atypical findings such as sleep disturbances, chorea, psychiatric symptoms, and peripheral neuropathy.19 The clinical examination can highlight the heterogeneity of CJD phenotypes, but the common factor is progressive decline in neurological status. The most common physical examination findings stratified by phenotypic strain are listed in Table 1.

**Diagnostic Tests**

In 1998 the WHO published diagnostic criteria for CJD (Table 3), with the diagnosis relying on clinical examination, EEG, and CSF findings. This is somewhat outdated in contemporary medicine because it does not take into account MRI findings, genetic testing, or modern laboratory tests for confirmation of the diagnosis. EEG and testing for 14–3-3 protein in the CSF are included in the criteria, and these studies are commonly done in the early stages of the diagnostic workup.

**EEG Findings**

Periodic sharp wave complexes (PSWCs) are found in the EEG recordings of approximately two-thirds of patients with sCJD, and have therefore been incorporated into probable sCJD diagnostic criteria by the WHO.55 The typical appearance for sCJD is that of a 1/second periodic triphasic sharp wave complex. The simple sharp waves can be classic triphasic, biphasic, or mixed. EEG-related spikes are independent from the traditional clinical findings of myoclonic jerking and are more likely to be related

**TABLE 3. The 1998 WHO diagnostic criteria for CJD**

<table>
<thead>
<tr>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rapidly progressive dementia</td>
<td>Myoclonus</td>
<td>PSWCs on EEG during an illness of any duration</td>
</tr>
<tr>
<td>Visual or cerebellar signs</td>
<td>Pyramidal/extrapyramidal signs</td>
<td>Positive 14–3-3 CSF assay in patients w/ disease duration of &lt;2 yrs</td>
</tr>
<tr>
<td>Akinetic mutism</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Probable CJD = (A) + at least 2 of (B) + positive result on at least 1 of (C) criteria. Possible CJD = (A) + at least 2 of (B) + no supportive EEG findings.
to the fusion of dendritic membranes in neurons. This subsequently results in electronic coupling through synchro-
nization in the corpus callosum.31

Historically, PSWCs for CJD were described as typi-
cally presenting in a bilateral frontal midline. However, there is growing evidence to support a more temporal EEG diagnostic template among patients diagnosed with sCJD. A lateralized PSWC distribution represents the pro-
dromal stage of disease onset. This localization will then progress to the typical more global bifrontal findings as the disease accelerates.54 This has dramatic effects on the clinical use of EEG findings as diagnostic criteria during initial disease onset.

Although lateralized PSWCs are described early in the disease course, nonspecific diffuse slowing background EEG patterns or other nondescriptive findings for primary cerebral function are most common. The sensitivity of EEG is generally low in sCJD, but when positive it is of value, giving a very high specificity. Heinemann et al. looked at PSWCs at a CJD surveillance unit in Germany, and found 37.5% sensitivity and 100% specificity in their cohort of 26 patients.20 When combined with clinical pre-
sentation, examination, and symptoms, the positive pre-
dictive value of diagnosis reached 99%.

On average, there is a 3.7-month delay from disease on-
set to the manifestation of PSWC findings on EEG. This is temporarily limited by the fact that there is only a mean 8-week survival after development of PSWCs with CJD. Even more important is that not all patients with sCJD develop PSWCs; these complexes are typically found in patients who have the MV1 and MM1 genotypes and not in those with valine homozygous variant at codon 129.58 The false-positive EEG findings were found in patients with Alzheimer disease and vascular dementia—patients who can easily be differentiated based on their profoundly different clinical disease presentation.54

It is vital to recognize that EEG patterns associated with CJD are sensitive to disease timing, benzodiazepines, and external stimulation. The PSWCs are likely to present while patients are awake and to be exacerbated with sleep deprivation. Benzodiazepines mask PSWC findings on EEG for patients with sCJD. Seizures, especially status epilepticus treated with benzodiazepines, are typically found in 15% of patients with CJD and may cloud the utility of EEG in diagnosis.

MRI Findings

Initial studies investigating the usefulness of the MRI in the diagnosis of CJD used a T2 imaging pattern consistently.3 The technological advancements in MRI enabled physicians to use FLAIR, diffusion-weighted imaging (DWI), and apparent diffusion coefficient (ADC), improving both the negative and positive predictive value.6 The term “pulvinar sign,” or bilateral FLAIR hyperintensity of the pulvinar area, was coined after reviewing 86 patients with vCJD. In that study 90% of the MR images were positive for the sign.13 However, recent comparative studies reliably concluded that both DWI and FLAIR were comparable in detecting hyperintensities in basal ganglia, although DWI was clearly superior in detection of similar changes in the neocortex.24

Now the anatomical areas of diagnosis have expanded from the traditional basal ganglia/thalamic areas to frontal, parietal, visual, temporal, limbic, and hippocampal cortices, which also correlate to the different molecular types of CJD. The pathophysiology of CJD with respect to imaging pattern is unique. Astroglialis has been linked to high-intensity signal patterns on T2/FLAIR sequences. Restricted diffusion of water leading to hyperintensity signals in DWI is attributed to the vacuole formation and deposition of prion protein in CJD, and is accounted as the principal neuropathological finding in the latter.37

Hyperintensity on DWI and ADC studies also correlate with the symptoms and clinical course of CJD. In patients with hyperintensity of basal ganglia on DWI, there is a shorter disease duration and higher incidence of myoc-
lonus. Shorter time between symptom onset and akinetic mutism is strongly correlated with the patients having hyperintense lesions in the occipital cortex on DWI.37

Diffusion tensor imaging was recently incorporated to study the pathogenesis of CJD in relation to white matter. Significant reductions of fractional anisotropy in patients with CJD in distinct and functionally relevant white matter pathways (including corticospinal, internal capsule, external capsule, fornix, and posterior thalamic radiation) correlate with progressive leukoencephalopathy and, more importantly, provide diagnostic criteria early in the disease course.28

The disease stage also has a significant impact on the nature of the MRI pattern. DWI is considered superior to any other MRI sequence in the early stages of CJD.53 Hy-
perintensity decreases in the later stages of the disease, and the only findings may be cortical atrophy.31 This is crucial information for the clinician to remember when the possibility of CJD is in the differential diagnosis, because the stage of the disease in relation to the positive predictive value of diagnostic criteria must be considered.

Patterns of FLAIR/DWI hyperintensity and restricted diffusion have also been shown to differentiate sporadic CJD from other causes of rapidly progressive dementia. Vitali et al. studied these MRI sequences across 90 pa-
tients, and found gray matter hyperintensities in all cases of sCJD, but never confined to just limbic regions.33 In all sCJD cases with basal ganglia or thalamic DWI hyperin-
tensities, there was associated restricted diffusion in the ADC map. These ADC findings were not seen in any non-
prion-related cases, where isolated limbic hyperintensities (FLAIR > DWI) were common. Combined MRI sensitivity and specificity for sCJD was 96% and 93%, respec-
tively, and these investigators concluded that the pattern of FLAIR/DWI hyperintensity and restricted diffusion can differentiate sCJD from other forms of progressive dementia.

Biomarkers in CSF

Several CSF biomarkers were proposed to play a sig-
nificant role in the diagnosis of sCJD. The most commonly studied is 14–3-3 protein, which is a surrogate marker for CJD and appears after neuronal destruction. Although there is still some debate in the literature, several studies conclude that the sensitivity of positive 14–3-3 protein in CSF for classic sCJD is 92%—96%.56 Tau protein is also
released with damage to neurons, and in classic sCJD has a sensitivity of 81% and a specificity of 85%; when tested in combination with 14–3–3 protein the positive predictive value is 91%. This is in contrast to the lower sensitivity of 53% in vCJD. The timing of CSF testing in relation to disease stage is crucial as well. Both 14–3–3 protein and tau protein are biomarkers for rapid neuronal destruction, and so the total concentrations increase as CJD progresses. When looking at the sensitivities and specificities of these tests in the course of the disease, Chohan et al. found a lower sensitivity and specificity of 64% and 71%, respectively, in the first third of the disease duration compared with 91% and 83% in the final third stage. The clinical picture is a necessary context when sending off for these studies. In the absence of clinical presentation, 14–3–3 detection has little to no value, because these results can be positive in Alzheimer disease or other disorders resulting in neuronal damage. Given the nonspecificity of these biomarkers, they should only be used to give confidence in the diagnosis with the right clinical picture, and never used to rule out the disease.

The problem with using these biomarkers is that they are not specific for CJD, because they are also found in many other diseases of the brain in which the pathology is neuronal loss. Positive 14–3–3 results are seen in viral encephalitides, recent stroke, subarachnoid hemorrhage, hypoxic brain damage, metabolic encephalopathy, glioblastoma, carcinomatous meningitis, paraneoplastic disorders, and corticobasal degeneration.

Real-time quaking-induced conversion (RT-QuIC) is a recently described laboratory technique that provides definitive diagnosis of CJD from CSF samples by detecting PrPSc. The RT-QuIC technique uses recombinant PrP as a substrate that is seeded with PrPSc and undergoes intermittent automated shaking, which amplifies the small amount of protein present into an accumulation of amyloid fibrils. Thioflavin T binds to these fibrils, and emits fluorescence that is detected in real time! The test has 80%–90% sensitivity, but is better than the test for 14–3–3 protein because it is 100% specific. There is also no difference in detection among the genetic subtypes.

As this methodology improves and becomes widely available, the test for 14–3–3 protein will become obsolete. The samples used in the test are becoming easier to obtain as well. Orru et al. used RT-QuIC with nasal brushings and showed a sensitivity of 97% and specificity of 100%, with positive results seen within 30 hours. These impressive results may be due to the suspicion that prions are shed in the olfactory nerves. This method is even less invasive than lumbar puncture, which only had 77% sensitivity when the CSF was tested in the same patients. Therefore, this modern technique should be part of the standard, initial testing for CJD, and part of updated diagnostic criteria for the disease.

Treatment

Despite all the advances in the understanding of this disease, the prognosis remains grim—CJD is a terminal disease. The mainstay of treatment is symptomatic and supportive, for example, using clonazepam for the treatment of myoclonus. Otto et al. showed a statistically significant improvement in cognitive function in a group of 28 patients with CJD treated with flupirtine, but this is the only study in the literature to report any symptom improvement with the use of medication. Future targets of therapy involve preventing the conversion of PrPSc to PrPSc.

Epidemiology

To evaluate the age-related incidence of CJD in the US, we queried the National (Nationwide) Inpatient Sample (NIS) to identify age-related frequency of inpatient hospitalization for CJD (ICD-9-CM diagnosis code 046.1, 046.11, and 046.19) between 2002 and 2011. Using discharge-level weights (DISCWT), 4428 cases were identified as CJD. The highest proportion of cases were observed in the 65- to 69-year-old group (820; 18.5%), followed by 70–74 years (629; 14.2%) and 75–79 years (625; 14.1%), followed by a decline (Fig. 1). Although limitations of the NIS are well known, including coding errors, these numbers approximately represent the epidemiology of inpatient admissions for CJD.

A comparison of incurred mean hospital charges for the inpatient stay for those undergoing brain biopsy/surgery found that costs were significantly higher than in those without any surgical intervention (p < 0.0001) for the years 2003–2008 and 2010–2011 (Fig. 2).

Discussion

Creutzfeldt-Jakob disease started out as a description of unusual neuropathological and clinical findings associated with other diseases in a series of 6 patients; over the last 90 years research has led to a well-described clinical and pathological entity, with typical imaging characteristics. The laboratory study of the disease also led to the development of new research techniques. More importantly, the study of prions demonstrated a novel disease pathogenesis devoid of DNA, dependent on pathological protein accumulation for propagation, and opened new fields of research in molecular and cellular biology, biochemistry, and biophysics that led to a Nobel Prize.

Need for New Diagnostic Criteria?

Despite the wealth of new information at present, most clinicians are at a disadvantage in diagnosing the disease, given that the WHO guidelines are outdated. A physician faced with the challenging task of diagnosing a patient with progressive neurological decline may not be aware of all the information at hand and may request a brain biopsy without a thorough diagnostic workup. Likewise, neurosurgeons may proceed with a biopsy that, based on the literature, has a low probability of being diagnostic, and an even lower chance of making a difference in the patient’s treatment plan or disease course. Newer diagnostic criteria do include hyperintensities on MRI as one of the tests that can be positive to make the case for probable CJD, but this still does not take into account more recent advances in research on the disease. Criteria for a diagnosis of CJD that includes newer confirmatory tests and incorporates
phenotypic strain provide a less invasive means of definitive diagnosis and will include the atypical patient. An updated diagnostic criteria and algorithm should provide confidence in the diagnosis, while avoiding unnecessary costs, waste of resources, and the potential morbidity associated with a biopsy.

Proposal for a Diagnostic Algorithm

The best approach to a suspected case of CJD is with a critical appraisal of the information known about the disease. We now have a classification of the different subtypes of disease present, and the goal in classifying prion diseases is to associate phenotype with genotype to aid diagnosis. If the most common imaging, clinical, and laboratory findings are studied in the context of the genotype, the clinician will have less doubt about the diagnosis. This is one more factor to be taken into account in updated diagnostic criteria. An updated list of criteria (Table 4) is proposed that takes into account CDI and RT-QuIC as part of the definitive diagnostics, removes the stipulation that the disease be present for less than 2 years (there are many reports in the literature of slowly progressive variants with a clinical course longer than 24 months), and includes specific phenotypes of the less common molecular strains.

An algorithm for the initial workup is provided in Fig. 3. The RT-QuIC technique is the most sensitive and specific antemortem diagnostic test, and should be the first one performed in the workup of a patient with suspected CJD, because it is the only study needed to diagnose the disease. A positive result will save money and resources by negating the need for any other studies. Because this test may not be available to many facilities yet, the proposed diagnostic pathway is for those without access to RT-QuIC, or for whom the results are negative.

This algorithm places diagnostic tests in order of sensitivity and timing of the disease course. The 14–3-3 protein is tested first, because CSF will be sent off to diagnose potentially treatable disorders. The routine CSF studies of glucose, total protein, white blood cell count, total cell count, and oligoclonal IgG in patients with CJD are generally unremarkable (Stoeck et al.20). This is of the utmost importance, because most other diseases in the differential diagnosis will usually have some abnormal result. Vasculitis, encephalitis, autoimmune disorders, and demyelination all usually have either CSF pleocytosis or elevated
protein levels, and should be ruled out in the initial round of testing. MRI is more sensitive than EEG and should be done next if there are negative results on the 14–3-3 assay, and then EEG is last because it is the least sensitive test of the three. An MRI study that shows the described CJD findings in combination with the usual symptomatology points toward CJD, and PSWCs are highly specific as well, so a positive test provides confidence in the diagnosis of probable CJD.

An examination finding in conjunction with genetic testing increases the diagnostic assuredness of the physician. Cerebellar signs or visual changes are atypical examination findings that are typical in CJD, and so a good history of the present illness is vital. In the patient with akinesis mutism and myoclonus, CJD should be at the top of the differential diagnosis. Those presenting with ataxia as the predominant symptom and MV1 or VV2 subtypes strengthen the probability of CJD. A long disease course with prominent ataxia and an MV2 genotype should give confidence in the CJD diagnosis, whereas a patient with temporal lobe hyperintensity on MRI, without ataxia, and an MM2 genotype also should be considered a probable case of CJD. On the other hand, a young patient (< 50 years) is likely to have the VV1 type.

**Utility of Biopsy**

The gold standard for diagnosis is pathological confirmation from a brain biopsy, but the frequency of a positively diagnostic biopsy is surprisingly low. Bai et al. performed a meta-analysis of 20 studies that described brain biopsy in cryptogenic neurological disease, and found an average diagnostic success rate of 54%, with a morbidity of 9%. Twenty-six biopsies at a CJD surveillance unit in Germany, where all suspected cases of CJD are referred, showed that 42% of the biopsies were nondiagnostic. All patients with CJD lacked inflammatory CSF numbers, and there was a statistically significant difference in the presence of myoclonus in CJD patients (82%) versus those with another disease (27%), again highlighting the necessity of accounting for all clinical information.

**Tissue Confirmatory Tests**

The reason for the low diagnostic yield of brain biopsy is the methodology used in the pathological sections. Histopathological investigation must show vacuolation, which may be missed on specimens. There needs to be ample amyloid deposition present for a positive immunohistochemical test for PrP. The CDI is a newer method that can detect the disease with a better diagnostic success rate than the current tests. Safar et al. used the CDI to examine 8 different regions of the brain in 28 patients who died of CJD. These investigators found 100% accuracy and detection of PrPSc in all regions tested in all patients, and no false positives in the control group. This is dramatically better than the diagnostic sensitivity of 17% and 22% for histological and immunohistochemical tests, respectively, in the comparison group. Another advantage of the CDI is that any part of the brain may be tested; it does not need to be an area with maximal change on imaging studies, and so a nondominant frontal lobe tissue sample will suffice.

**Indication for Biopsy**

The argument against biopsy in CJD is that not only are the diagnostic success rates low, but more importantly—will this lead to any change in treatment plan or disease course? In a study of 90 patients with rapidly progressive dementia, Schott et al. showed a 57% diagnostic rate, but only 11% of these diagnoses resulted in a change of treatment. Some will argue that because a biopsy will provide no change in the treatment, it only is adding costs and wasting resources that can be used elsewhere. Figure 2 shows a recent trend in hospital costs associated with those undergoing a surgical procedure that provides no benefit. It is important for treating or consulting physicians to know that the RT-QuIC test exists and that getting brain tissue is unnecessarily invasive and has a lower diagnostic sensitivity.

Proponents of brain biopsy argue that the main justification for biopsy is to diagnose a potentially treatable disease or confirm the diagnosis of CJD in the presence...
of atypical features. These include CNS vasculitis, encephalitis (viral or autoimmune), malignancy, demyelination, nonviral CNS infections, and sarcoidosis. Schuette et al. showed a rate of change of 8% in the treatment plan in a series of 135 consecutive brain biopsies, and in only 4% did the findings lead to an improvement in the disease course.47 Chitravas et al. looked at 1106 patients who underwent a brain biopsy for dementia and found that only 6% of the patients in the study had a treatable disorder (i.e., not CJD) and that two-thirds of these cases did not meet WHO criteria for CJD.11

If all patient information is taken into account, then the likelihood of missing a treatable disease on brain biopsy is very low. Neoplasms are often the most common conditions found in these studies, but MRI would easily differentiate between the two, because prion diseases do not present with mass lesions or gadolinium uptake. History, physical examination, MRI, and routine CSF studies are ample and necessary to separate treatable conditions (that may need a brain biopsy) from prion disease, which can be confirmed in less invasive ways.

Brain biopsy should only be done after all noninvasive diagnostic avenues are exhausted, and do not point to a cause. Because a biopsy for dementia is rare, there is no way to standardize when to perform the procedure. All information must be taken into account: the prominent clinical signs, patient’s age, duration of disease, imaging studies, EEG results, and RT-QuIC assay of either nasal brushing or CSF. In the patient with atypical features, no findings on imaging studies or EEG, and negative CSF studies (including ruling out the autoimmune and viral encephalitides), a biopsy is indicated and the CDI must be the diagnostic test used.

Conclusions

Creutzfeldt-Jakob disease is a fatal neurological condi-
tion that often perplexes the treating physician. Given its rarity, published diagnostic criteria aid in the approach to diagnosis, but these are unfortunately outdated and do not incorporate modern research laboratory methods. The proposed updated diagnostic criteria and treatment algorithm are based on the latest research, and help guide the workup in a disease that can be difficult to diagnose. Knowledge of different strain phenotypes is crucial to making an accurate diagnosis, and a brain biopsy need be performed only rarely in the atypical patient, because RT-QuIC appears to be the best test to provide a confirmed diagnosis.

Acknowledgments

We thank Drs. Osama Ahmed and Tammy Maiti for their support.

References

4. Boëlle PY, Cesbron JY, Valleron AJ: Epidemiological evidence of higher susceptibility to vCJD in the young. BMC Infect Dis 4:26, 2004

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

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Conception and design: Manix, Henry, Guthikonda. Acquisition of data: Manix, Kalakoti, Thakur, Menger. Analysis and interpretation of data: Manix, Kalakoti, Thakur, Menger. Drafting the article: Manix, Kalakoti, Henry. Critically revising the article: Manix, Henry, Thakur, Menger, Guthikonda. Reviewed submitted version of manuscript: Manix, Kalakoti, Henry, Guthikonda. Administrative/technical/material support: Nanda, Kalakoti. Study supervision: Nanda, Guthikonda.

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