کارگاه‌های آموزشی مرکز اطلاعات علمی

مقاله نویسی علوم انسانی

اصول تنظیم قراردادها

آموزش مهارت های کاربردی در تدوین و چاپ مقاله
Case Report

Creutzfeldt-Jakob Disease Presenting with Confusion and Visual Disturbance

Masoud Nikanfar MD*, Mehdi Farhoudi MD**, Monireh Halimi MD***, Fereidoon Ashrafian-Bonab MD***, Kaveh Mehrvar MD†

Creutzfeldt-Jakob disease is increasingly being reported in the last three decades as a result of increased awareness for the disease. Various studies have reported an annual incidence of 0.5 – 1.5 cases of Creutzfeldt-Jakob disease per million of general population. However, in our country, like other developing countries, the disease is still under-reported. Herewith, we described our clinical experience with an autopsy-proven case of Creutzfeldt-Jakob disease.

Keywords: Confusion • Creutzfeldt-Jakob disease (CJD) • vision

Introduction

Prions can cause neurodegenerative diseases that have long incubation periods and progress inexorably once the clinical symptoms appear. So far, five human prion diseases have been recognized including Kuru, Creutzfeldt-Jakob disease (CJD), variant Creutzfeldt-Jakob disease (vCJD also known as new variant CJD), Gerstmann-Strassler-Scheinker (GSS) syndrome, and fatal familial insomnia (FFI).1, 2 Bovine spongiform encephalopathy (BSE), one of prion infections affecting animals, was responsible for a more widespread public attention with its possible link to vCJD.3, 4

These human prion diseases share certain common neuropathologic features including neuronal loss, proliferation of glial cells, absence of an inflammatory response, and the presence of small vacuoles within the neutrophils, which produces a spongiform appearance. The current theory is that prion diseases are associated with the accumulation of an abnormal form of a host cell protein, the so-called “prion protein” (PrP).5

This paper describes the clinical experience with an autopsy-proven case of CJD managed at the Department of Neurology of Razi Hospital, Tabriz, North-West of Iran.

Case Report

A 71-year-old woman referred to our hospital on May 3, 2004 with a one and a half-month history of visual symptoms accompanied by transient confusion and disorientation and dystonic posture in her right arm.

Her family members reported that she had transient attacks of agitation, hallucinations, and confusional state with a duration of about 15 minutes. After each attack, she became normal without any memories of attacks. Gradually, the frequency of these attacks increased. Meanwhile, she complained of transient visual problems such as macropsy, micropsy, blurred vision, and color vision disturbance. She had also transient dystonic posture in her right arm. Before admission, she also developed a mild left-sided hemiparesis. She was visited by a psychiatrist who prescribed haloperidol and clonazepam with no beneficial effect; her condition became worse progressively.

On admission, she had mild confusion, and headache, and vertigo. General examination was normal. In neurologic examination, she had a left hemiparesis, left central hemifacial weakness, no
plantar reflex in the left side, and an action tremor in her right arm.

All biochemical laboratory tests of plasma and urine were normal. Cerebrospinal fluid analysis was normal. In brain computed tomography (CT) and magnetic resonance imaging (MRI), multiple lacunar infarctions and senile atrophy were reported. In her first electroencephalogram (EEG), generalized slow sharp waves appeared in all montages (Figure 1A).

The patient’s general condition and level of consciousness progressively deteriorated during hospitalization. After 15 days, she developed myoclonic seizures — first in her right arm and then generalized. In the next EEG, there was progressive slowness and periodic sharp waves. In her last EEG, slow and disorganized background activity that was interrupted by repetitive discharges of large sharp waves in all montages — about one cycle per second — were recorded (Figure 1B, C). She was intubated and finally on June 9, 2004, she died of sepsis.

In pathologic study performed two days after death, the spongiform encephalopathy was documented. Histopathologic examination revealed no Kuru plaque in Congo red and PAS staining. No inflammatory infiltration was present (Figure 2).

**Discussion**

CJD, although rare, is the most frequent prion disease of humans. Sporadic (sCJD), familial (fCJD), iatrogenic (iCJD), and recently, variant (vCJD) forms of CJD are all recognized. Approximately, one case of CJD occurs per 1,000,000 population per year with a worldwide distribution. The mean age of onset is 65 years. Patients with vCJD and iCJD tend to be much younger.

There is no gender predilection for CJD. The vast majority of CJD cases (85 – 95%) are sporadic in whom there is no known infectious source and no evidence of the disease in the prior or subsequent generations of the patient’s family.

Rapidly progressive mental deterioration and myoclonic seizures are the two cardinal clinical manifestations of CJD. However, a number of variants or subtypes of the disease have been defined based upon focal neurologic findings. Examples of these include forms with mainly visual, cerebellar, thalamic, and striatal features.

Some clinical findings, although compatible

---

**Figure 1.** A) Generalized slow sharp waves. B) Progressive slowness and periodic sharp wave; C) Slow and disorganized background activity that interrupted by repetitive discharges of large sharp waves about one cycle per second.
with CJD, should raise the suspicion of an alternative diagnosis. These include cranial nerve abnormalities, sensory abnormalities, and involvement of the peripheral nervous system.

Five percent of CJD cases are familial. The only clearly established route of transmission is iatrogenic — during transplantation of corneas or dural grafts from infected individuals, after implantation of infected depth electrodes, and injection of human growth hormone or gonadotropins with cadaveric sources. The transmissible agent is resistant to boiling, treatment with formalin and alcohol, and ultraviolet radiation, but can be inactivated by autoclaving at 132 °C and 15 PSI for one hour or by immersion in 5% sodium hypochlorite (household bleach) for one hour.11

The WHO criteria for probable sporadic CJD include progressive dementia, and at least two of the following criteria: myoclonus seizure, visual or cerebellar disturbance, pyramidal/extrapyramidal dysfunction, akinetic mutism, typical EEG, and/or CSF positive for 14-3-3 protein with a clinical duration to death of <2 years; routine investigations should not suggest an alternative diagnosis for definite diagnosis. In addition to the above criteria, one of the two followings should be found: characteristic pathologic changes in brain or positive western blot for confirming the prion protein PrPSC.12

The role of MRI and especially diffusion-weighted MRI (DWI) is becoming increasingly important in clinical diagnosis of CJD.

Patients with CJD may have abnormal patterns of hyperintensity in the basal ganglia on MRI or DWI. DWI is more sensitive than T2-weighted and flair MRI for the detection of CJD-related lesions, especially in the early weeks of the disease.13

A characteristic EEG pattern of periodic synchronous bi- or tri-phasic sharp wave complexes (PSWC) is observed in 67 – 95% of patients with sCJD at sometime in the course of illness. However, the absence of PSWSC does not exclude the diagnosis of sCJD. In these cases, serial EEG is recommended.

Abnormal CSF protein, especially the 14-3-3 protein serves as a marker for CJD. The protein 14-3-3 has been advanced as a sensitive and specific diagnostic test for sCJD.14

Routine laboratory studies are normal in CJD with occasional exceptions of liver function tests.15 Elevated CSF protein level may occur in about 40% of patients.

In differential diagnosis of CJD, Wipple disease, lithium intoxication, Hashimoto encephalopathy, angiocentric lymphoma, carcinomatous meningitis, major psychosis, unusual form of Alzheimer disease, Lewy-body disease, subacute sclerosing panencephalitis, limbic-brainstem-cerebellar encephalitis, and cerebral lipidosis should be considered.11

A definitive diagnosis of CJD requires brain biopsy or autopsy which can detect the characteristic changes in the brain tissue caused by the disease. Because these procedures are expensive and hazardous, they are usually not performed.

The difficulties of diagnosing CJD might have led to rare reports of the disease. There are only two published case reports from Iran.16, 17 Since the disease progresses rapidly, the patient may die before a diagnosis can be made. Furthermore, some physician may not even consider the possibility of CJD because the disease is rare and the clinical symptoms can often be attributed to other ailments. Consequently, CJD may be mistaken for a variety of psychological illnesses because the behavioral changes are seen in 30% of patients at the onset of disease and in 57% of patients in later stages. The disease is fatal within one year in 90% of patients. Our patient had a rapid course and died after two and a half months.

In conclusion, our experience with this patient showed that in any adult patient with psychotic symptoms such as delirium hallucinations, etc., CJD must be considered. In such patients, EEG is an important diagnostic test and must be done.
References


کارگاه‌های آموزشی مرکز اطلاعات علمی

مقاله نویسی علوم انسانی

اصول تنظیم قراردادها

آموزش مهارت های کاربردی در تدوین و چاپ مقاله