Clinical Group

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Case Report

Case Report of Clinically Probable Sporadic Creutzfeldt – Jakob Disease from A Tertiary Care Hospital in South India

Abstract

We are reporting an otherwise healthy lady of forty eight years suffering from rapidly progressive dementia, behavioural disturbances, myoclonus, ataxia and extrapyramidal features of eight months duration with characteristic abnormalities on brain imaging, electroencephalography with positive cerebrospinal fluid 14-3-3 protein, satisfying the revised World Health organization (WHO) criteria for diagnosis of “clinically probable Sporadic Creutzfeldt-Jakob disease (sCJD)”. An attempt was made to clinically differentiate the diagnostic possibilities.

Introduction

Sporadic Creutzfeldt-Jakob disease is the most common of the prion diseases affecting the human race [1]. Prion diseases, a group of uniformly fatal neurodegenerative diseases, caused by conformational change of PrPc (normal cellular form of prion protein) to abnormal PrPsc (abnormal scrapie of prion protein) which cumulates in various regions of the brain resulting in spongiform degeneration and gliosis [2]. In humans, prion protein is encoded by a gene PRNP located on short arm of chromosome 20. Sporadic CJD has been divided into approximately six molecular subtypes based on the genetic polymorphism at codon 129 in the prion gene (MM, MV, or VV) and the type of protease resistant Prion (type 1 and type 2). Sporadic CJD patient’s progress very rapidly starting with dementia, behavioral dyscontrol in the form of aggressive outbursts, restlessness, irritability, delirium, hallucinations, delusions, apathy, confusional spells followed by pyramidal, extrapyramidal, cerebellar symptoms including myoclonus.

The characteristic features of myoclonus includes-being generalized, fast at a frequency of 1 per second, stimulus sensitive and non-epileptic. Ocular symptoms sometimes coexist in the form of cortical blindness, distortion of seen objects and paralysis of convergence or upgaze. Our patient has multifocal generalized myoclonus on a background of dementia, behavioral disturbances, myoclonus and terminally into akinetic mute state.

As already mentioned, MRI brain can be normal in the earlier stage of the disease, however, the signal alterations...
Table 1: Differentiating features of Sporadic CJD from Variant CJD

<table>
<thead>
<tr>
<th>Clinical feature</th>
<th>Sporadic CJD</th>
<th>Variant CJD</th>
</tr>
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<tbody>
<tr>
<td>Average age at Clinical Onset</td>
<td>63 years</td>
<td>28 years</td>
</tr>
<tr>
<td>Length of survival from date of clinical onset</td>
<td>4 months</td>
<td>14 months</td>
</tr>
<tr>
<td>Early symptoms</td>
<td>Cognition, behavioural</td>
<td>Psychosis, depression</td>
</tr>
<tr>
<td>MRI Brain</td>
<td>Increased signal in caudate, putamen</td>
<td>Increased signal in Pulvinar region of Thalamus</td>
</tr>
<tr>
<td>EEG</td>
<td>Bi-phasic or triphasic complex</td>
<td>Non-specific or slow</td>
</tr>
<tr>
<td>CSF 14-3-3 Protein</td>
<td>Usually elevated</td>
<td>Usually not elevated</td>
</tr>
<tr>
<td>Histopathology of brain tissue</td>
<td>No amyloid plaques</td>
<td>100% florid plaques</td>
</tr>
<tr>
<td>IHC staining pattern of brain tissue</td>
<td>Punctate pattern</td>
<td>Widespread plaque staining pattern</td>
</tr>
<tr>
<td>IHC staining of tonsil/appendix</td>
<td>Negative</td>
<td>PrP/present in late stages</td>
</tr>
</tbody>
</table>

was present. She was swaying to either side while walking requiring one person support to walk. No preceding history of fever, headache, vomiting and seizures. No history of recent vaccination, dog bite or head injury. She consumes non-vegetarian diet.

She scored 12 over 28 Mini-mental status examination (Folsteine MMSE) scale suggestive of severe dementia. She was suspicious, anxious and fearful with hallucinations. She had insight about her problem but impaired judgement. Significant impairment in recent and immediate memory domains with remote memory being intact. Stimulus sensitive myoclonus was present with auditory startle. No cranial nerve involvement including eye movement abnormalities. She had lower limb rigidity with normal tone in the hands. She had normal power, preserved deep tendon jerks and plantars were flexors. She had postural tremors of both hands, generalised multifocal myoclonic jerks of the trunk and the limbs. Bilateral finger nose in coordination with impaired tandem gait was present.

She had rapidly progressive dementia to start with frontal lobes followed by temporal and parietal lobes with cerebellar ataxia, tremors and myoclonus. Differential diagnostic possibilities of Paraneoplastic encephalitis, autoimmune encephalitis, Hashimoto's encephalitis, Isolated CNS Vasculitis were considered (Table 2).

**Laboratory evaluation**

She had hemoglobin of 11 mg%, total leucocyte count of 8200 cells/cu.mm, ESR of 20 mm in 1st hour. Serum Vitamin B12 was within normal limits. Renal function tests were normal, liver profile was normal including plasma ammonia. Serum electrolytes including Calcium were within normal limits. Thyroid profile was normal including anti-thyroid peroxidase (anti–TPO) antibodies. Serum was negative for anti–nuclear antibodies (ANA), Human immunodeficiency virus (HIV), Venereal disease research laboratory test (VDRL) and Voltage gated potassium channel antibody level (VGKC).

Cerebrospinal fluid (CSF) analysis revealed glucose of 72 mg/dl, protein of 20 mg/dl with four lymphocytes. CSF for Gram stain and Acid fast bacillary stain was negative. HSV–DNA PCR was negative in CSF. CSF for 14–3–3 protein was positive 1.5 ng/ml (reference range <1.5 ng/ml). CSF for paraneoplastic panel was negative for anti–Hu, anti–Ma 2 and CV–2. Magnetic resonance imaging of the brain (Figures 1,2) showed more or less symmetrical T2 and FLAIR hyperintensities and reduction of ADC in striatal hyper intensity regions on DWI differentiates sporadic CJD from other rapidly progressive dementia with 98% sensitivity and 100% specificity [7].

**Case Report**

48 year old lady, mother of 2 children, uneducated, right handed, was brought by family members with eight months history of forgetfulness involving the household works, failure to recognize the familiar faces and naming them, associated with repeating the words uttered and delayed response to the question asked. She keeps on repeating the same task without interruption. Geographic disorientation and way finding difficulty was present since 1 month. Behavioral symptoms of 2 months duration comprising restlessness, irritable and agitated at times with emotional liability of 1 month duration. She also developed infrequent jerky movements of the trunk and the limbs and tremors of the hands more than legs, bilateral (left side more than right side). Occasional incontinence of urine precede the changes in EEG and CSF abnormalities. Among the sequences, T2, FLAIR, DWI, ADC mapping has been used to detect the abnormality. DWI sequence is more sensitive (92%) and specific (93%). However combination of DWI and FLAIR signal changes increased the sensitivity to 98% and specificity to 95%. DWI sequence is more sensitive in the diagnosis of CJD in its early stage independent of EEG changes. Two patterns of abnormalities were described like isolated cortical gyri hyperintensities, combined cortical and deep gray nuclear (basal ganglia) involvement and Isolated Basal ganglionic changes could have been Third pattern [4]. Most common type is pattern two. The pathological correlate of T2 and FLAIR hyperintensity is astroglisis, while vacuole formation which restricts the water diffusion in DWI imaging. Involvement of deep gray matter is associated with shorter disease course with rapidly progressing neurologic deterioration whereas absence of basal ganglia involvement correlates with delayed onset of dementia and longer disease course [5]. Combined DWI and FLAIR has higher sensitivity (91%) and specificity (95%) for CJD [6]. However, Vitali et al reaffirms that the pattern of FLAIR/DWI hyperintensity and reduction of ADC in striatal hyper intensity regions on DWI differentiates sporadic CJD from other rapidly progressive dementia with 98% sensitivity and 100% specificity [7].
Discussion

This patient was diagnosed as clinically probable CJD in view of rapidly progressive dementia, cerebellar ataxia and cortical myoclonus, disease duration of less than 2 years, EEG abnormality and MRI brain abnormality, which is compatible with the diagnosis of CJD.

The age at onset of the disease was relatively younger in our case as similar to the published cases by Mehndiratta et al. [8]. The female preponderance among the published case series was also reported by Mehndiratta et al. [8]. Our patient had the typical clinical manifestations like rapidly progressive dementia, myoclonus, ataxia, tremors and behavioural symptoms. Similar observations were noted in the case series by Velásquez-Pérez et al. [9] and González Duarte et al. [10]. Behavioural manifestations in CJD occur in 30% of patients at the onset and 57% at later stages of the disease [1]. Mahale et al. found 62% of their patients having behavioural manifestations [11]. Our patient did not have any family history, hence we presume this as sporadic CJD. Familial CJD was reported only in 2 cases by González Duarte et al [10], and single reports by Mehndiratta et al. [8] and Chandra et al. [12]. The time interval between the onset of the symptoms to the diagnosis was 8
months in our case while it ranges from 1-12 months in other case series [8,10]. As per the literature, the mean duration from onset of symptoms to the time of death was 6.6 months (Range: 3-14 months) [8,10]. The mean survival of CJD patients is 5 months and about 80% of patients succumb to disease after 12 months from onset [1]. Our patient is alive and dependent on care givers for her daily functioning.

EEG in our patient showed triphasic waves and periodic sharp wave complexes, however, it can be normal in the earlier part of the disease but classical features would be generalized periodic sharp waves, biphasic and triphasic waves at a rate of one per second. EEG has a sensitivity of 67% and specificity of 74 86% in the diagnosis of CJD [10]. Repeated EEG during the course of disease increases the probability of demonstrating characteristic EEG abnormality. Similar EEG findings is sometimes possible in Hashimotos encephalopathy and VGKC encephalitis but easily can be differentiated as illustrated in (Table 2).

Imaging study of the current patient revealed abnormality in both deep gray nuclei and cortical gyri. Similar to our observations, Mahale et al. reported six of their eight patients showing symmetrical basal ganglionic hyperintensities, which were showing restriction under DWI imaging, however cortical signal changes (parieto-occipital, frontal and temporal) did not show restriction [11]. Similar abnormalities were also reported by González Duarte et al. [10] in 5 of 7 patients showing only basal ganglionic abnormalities without cortical signal changes (Pattern 3). Recently, Biswas et al. [13] had shown 100% (10/10) of their patients showing Pattern two abnormalities (Basal ganglionic signal changes – 8 patients and parieto occipital signal changes – 2 patients). Magnetic resonance imaging of brain can be normal in the earliest stages of CJD while it may show only atrophy in the terminal stage of the illness with the classical signal alterations might have disappeared.

The detection of the 14-3-3 protein in CSF has been one of the markers for diagnosis of CJD [14]. This protein detection raises the accuracy in diagnosing sporadic CJD with sensitivity of 92% and specificity 80% [15]. But Zerr et al. refuted this observation as they found that imaging findings were equivalent to elevated levels of the 14-3-3 protein in the diagnosis of probable sporadic CJD [16]. Other biomarkers such as t-tau, p-tau, S-100, or neuron-specific enolase (NSE) in the CSF are required in addition to, protein 14-3-3 [15].

**Diagnosis**

The diagnosis is based on clinical profile of the onset and course of symptoms, EEG changes and typical imaging abnormalities. Several clinical criteria exist, but the better bedside one is the CDC’s Clinical criteria for diagnosis of CJD 2010 which is the Revised WHO criteria (Table 3) [17]. Of late, other than CSF 14-3-3 which is a surrogate marker for the diagnosis of CJD, several other markers like Neuron specific enolase (NSE), S-100 beta amyloid, p-tau, t-tau are available. Focusing the possibilities of early detection of prion diseases, recently, anti-prion screening test using real time quacking-induced conversion has been reported [18]. Though, olfactory epithelium is a known peripheral target of PrP deposition in sCJD [19], CSF and urine are also reliable source for early detection of sCJD [20], whereas, a new blood test has been recently identified with 100% sensitivity and specificity in vCJD [21]. Yield of different diagnostic tests is enlisted in the (Table 4).

**Conclusion**

High index of suspicion is required to identify the patients with classical manifestations so as to register to the national authority and to take necessary preventable action, if possible, and to treat symptomatically in view of lack of definite curative treatment at present for this fatal disease.

**References**


17. Revised WHO Definition of CJD Subtypes (Incorporating the New Definition of Probable Sporadic CJD as Per Section 3.1. Link: https://goo.gl/RNYMKO


