Creutzfeldt-Jakob disease (CJD) and variant CJD (vCJD)

**General introduction**

Creutzfeldt-Jakob disease is the prototype of a family of rare and fatal human degenerative conditions characterized by progressive brain dysfunction. CJD falls into four categories: sporadic, familial, iatrogenic and variant. Sporadic CJD (sCJD), thought to occur worldwide, is the most common form and represents about 85% of all CJD cases; incidence ranges between 0.5 and 1.5 cases per million inhabitants per year.

Variant CJD (vCJD), first recognized in the United Kingdom in 1996 has been linked to bovine spongiform encephalopathy, (BSE). Incidence is not currently monitored in many parts of the world and the size of the population exposed and susceptible to vCJD in the United Kingdom – and in other parts of the world – is not known. This and uncertainties relating to the potential length of the incubation period complicate predictions of the future number of vCJD cases. All those who have been identified with vCJD have died. No screening test is currently available.

**Causal agent and main modes of transmission**

**Causal agent:**
CJD is believed to be caused by a self-replicating host-encoded protein or prion protein (PrP), transmissible in the laboratory to many species, including wild and transgenic mice and non-human primates.

**Main modes of transmission:**

- For vCJD, the exact route of transmission from BSE to humans is uncertain, but is likely to be linked to contaminated foods of bovine origin. Probable secondary transmission of vCJD via blood transfusion has been reported. Incubation not known precisely, likely to be long.
- Mode of transmission for sCJD is not known; de novo spontaneous generation of self-replicating protein has been hypothesized. No clear evidence of risk from diet, previous surgery, blood transfusion, occupational or animal exposure, the possibility that sporadic CJD arises through other unrecognized environmental exposure cannot be dismissed.
- Familial CJD (fCJD) is an inherited condition and cases represent 20% to 15% of the total number of CJD cases. Gerstmann-Sträussler-Scheinker disease (GSS) and fatal familial insomnia (FFI) are very rare forms of fCJD.
- Iatrogenic CJD infection is inadvertently transmitted usually from a case with sCJD in the course of medical/surgical treatment, e.g. human pituitary hormone therapy, human *dura mater* grafts, corneal grafts or neurological instruments.

**Disease forms and recommended case definition**

**Sporadic CJD**

*Possible case*
- Progressive dementia; and
- EEG atypical or not carried out; and
- duration <2 years; and
- at least two out of the following clinical features: myoclonus, visual or cerebellar disturbance, pyramidal, extrapyramidal dysfunction, akinetic mutism.

*Probable case* (in the absence of an alternative diagnosis from routine investigation)
- Progressive dementia;
- at least two of the following four clinical features: myoclonus, visual or cerebellar disturbance, pyramidal/extrapyramidal dysfunction, akinetic mutism, with
  - a typical EEG (generalized triphasic periodic complexes at approximately one per second), whatever the clinical duration of the disease, and/or
  - a positive 14-3-3 assay in CSF and a clinical duration leading to death in <2 years.

*Definite case*
- Neuropathological confirmation (see end of CJD case classification below); and/or
- confirmation of protease-resistant prion protein (PrP) (immunocytochemistry or Western Blot); and/or

Excerpt from “WHO recommended standards and strategies for surveillance, prevention and control of communicable diseases”
• presence of scrapie-associated fibrils.

**Iatrogenic CJD**

*Probable*
- Progressive cerebellar syndrome in a recipient of human cadaver-derived pituitary hormone; or
- Probable CJD with a recognized iatrogenic risk.

*Definite*
- Definite CJD with a recognized iatrogenic risk.

**Familial CJD**

*Probable*
- Probable CJD plus confirmed or probable CJD in a first degree relative; and/or
- Neuropsychiatric disorder plus disease-specific prion protein gene (PRNP) mutation.

*Definite*
- Definite CJD with a recognized pathogenic PRNP mutation; and
- Definite or probable TSE in a first-degree relative

**Note.** For surveillance purposes, this definition includes Gerstmann-Sträussler-Scheinker (GSS) syndrome and fatal familial insomnia (FFI).

**Variant CJD**

Variant CJD cannot be diagnosed with certainty on clinical criteria alone; this requires neuropathological confirmation. The following combinations of signs, symptoms and clinical investigations serve to define possible, probable and definite vCJD:

(I)
- Progressive psychiatric disorder
- Clinical duration >6 months
- Routine investigations do not suggest an alternative diagnosis
- No history of potential iatrogenic exposure
- No evidence of a familial form of TSE (transmissible spongiform encephalopathies).

(II)
- Early psychiatric symptoms (depression, anxiety, apathy, withdrawal, delusions)
- Persistent painful sensory symptoms (pain and/or dysaesthesia)
- Ataxia
- Chorea / dystonia or myoclonus
- Dementia.

(III)
- EEG unknown or does not show the typical appearance of sporadic CJD (generalized triphasic periodic complexes at approximately one per second)
- Bilateral symmetrical pulvinar high signal on MRI brain scan (relative to other deep gray-matter nuclei).

(IV)
- Positive tonsil biopsy.

**Note.** Tonsil biopsy not recommended routinely nor in cases with EEG appearances typical of sporadic CJD, but useful in suspect cases where clinical features are compatible with vCJD and MRI does not show bilateral pulvinar high signal. Cerebral biopsy in living patients is to be discouraged unless its purpose is to arrive at an alternative diagnosis of a treatable disorder.

**vCJD case classification**

*Possible*
- A patient with the items under (I) above and at least 4 items under (II)
- EEG does not show the typical appearance of sporadic CJD

*Probable*
- A patient with the items under (I) and at least four items under (II)
- Bilateral pulvinar high signal on MRI brain scan
- EEG does not show the typical appearance of sporadic CJD although generalized periodic complexes may occasionally be seen at the later stages of the disease.

OR
- A patient with items under (I) and a positive tonsil biopsy.

*Definite*
- A patient with the items under (I) above
Neuropathological confirmation of vCJD.

Neuropathological criteria (CJD and other human transmissible spongiform encephalopathies)

**Creutzfeldt-Jakob disease:** sporadic, iatrogenic (recognized risk) or familial (same disease in first-degree relative or disease-associated PRNP mutation):
- spongiform encephalopathy in cerebral and/or cerebellar cortex and/or subcortical grey matter; and/or
- encephalopathy with prion protein (PrP) immunoreactivity (plaque and/or diffuse synaptic and/or patchy/perivacuolar types).

**Variant CJD**
- Spongiform encephalopathy with abundant PrP deposition, in particular multiple fibrillary PrP plaques surrounded by a halo of spongiform vacuoles (“florid” plaques, “daisy-like” plaques) and other PrP plaques, and amorphous pericellular and perivascular PrP deposits especially prominent in the cerebellar molecular layer.

**Gerstmann-Sträussler-Scheinker disease** (in family with dominantly inherited progressive ataxia and/or dementia and one of a variety of PRNP mutations)
- Encephalo(myelo)pathy with multicentric PrP plaques;
- Thalamic degeneration, variable spongiform change in cerebrum.

**Kuru**
Spongiform encephalopathy in the Fore population of Papua New Guinea.

**Surveillance**

**Rationale for surveillance**
Global surveillance of vCJD and other forms of CJD should lead to a better understanding of the disease, including potential causes of iatrogenic CJD, as well as the distribution of various hereditary forms. It should also provide information towards protection against the risks of disease.

**Recommended types of surveillance**
- One centre must be identified at central level to carry out surveillance.
- All definite, probable and possible cases must be notified by the appropriate health care professionals (usually physicians, neurologists, psychiatrists, neuropathologists) to the centre responsible for surveillance.
- Check death registrations in order to identify cases not detected by routine surveillance.

**Recommended minimum data elements**

**Case-based record**
- Subtype and classification of CJD.
- Age, sex, country of birth, geographical information, occupation.
- Date of onset, date of death.
- Vital status (alive, dead).

**Aggregated data reporting.** All reporting should be case-based.

**Recommended data analyses, presentation, reports**
- Number of cases by subtype, classification, occupational group, geographical area.
- Number of cases by year of death, by age at death.
- Sex ratio.

**Performance indicators for surveillance.** Time between onset of symptoms and reporting.

**Control activities**

**Case management**
Supportive case management (no specific treatment). The disease is always fatal.

**Prevention**
- Avoiding exposures to BSE-causing agent in food of bovine origin:

Excerpt from "WHO recommended standards and strategies for surveillance, prevention and control of communicable diseases"
BSE is a risk to animal and public health. It is transmissible to humans and food is considered the most likely source of exposure. Bovines, bovine products and by-products potentially carrying the BSE agent have been traded worldwide, giving this risk a global dimension, with possible repercussions on public health, animal health and trade. At this time protecting public health from exposure through food is primarily accomplished by preventing and eliminating BSE in livestock populations. For further information consult report of the Joint WHO/FAO/OIE technical consultation on BSE: public health, animal health and trade (see Bibliography below).

- Avoiding iatrogenic exposures:
  - Specific precautions should be taken in the management of persons with confirmed or suspected transmissible spongiform encephalopathy (TSEs) and their tissues.
  - The following persons have been regarded as “at risk” for developing TSE: recipients of human dura mater, of human cadaver-derived pituitary hormones especially human cadaver derived growth hormone, of cornea transplants, persons undergoing neurosurgery and members of families with heritable TSE.
  - When determining the risk of iatrogenic transmission, infectivity of a given tissue should be considered together with the route of exposure. Infectivity is found most often and in the highest concentration in the central nervous system. Precautions to be taken when performing certain interventions (dental, diagnostic, surgical procedures) and when handling instruments, cleaning and decontaminating instruments, work surface and wastes have been proposed.
  - Probable secondary transmission of vCJD via blood transfusion has been reported. The possibility of transfusion transmission of vCJD has important implications for public health and a range of measures have been taken in a number of countries, especially the United Kingdom, to minimize the risk. For further information see Bibliography below.

Epidemics

*Conditions under which epidemics may occur*

The most likely origin of bovine spongiform encephalopathy (BSE) was elucidated in the United Kingdom in 1988, namely the use of ruminant protein in cattle feedstuffs, recycling of BSE contaminated bovine offal being the driving force behind the spread of the epidemic. After the United Kingdom where it emerged, BSE was identified in the native cattle populations of several countries, mostly in Europe. See information at OIE (World Organisation for Animal Health) web site at [http://www.oie.int/eng/info/en_esbmonde.htm](http://www.oie.int/eng/info/en_esbmonde.htm).

As of October 2005, a total of 157 confirmed and probable cases of vCJD had been reported in the United Kingdom. A small number of vCJD cases had been notified in seven other countries. Food of bovine origin is considered the most likely source of human exposure to the BSE agent.

*Management of epidemics*

- All countries should determine the BSE risk status of their cattle population through the outcome of an annual risk assessment identifying all potential factors for BSE introduction, recycling and amplification. A BSE surveillance system fitting the estimated level of risk should be put into place.
- Whenever a risk of BSE is identified, countries must take immediate steps to define the specific risk material (SRM): all tissues that have been shown to contain infectivity should be removed and destroyed. If the BSE risk is considered higher other tissues that under certain conditions may carry infectivity should be added to the SRM list for removal and destruction. Additional precautions may be taken such as prohibiting cattle over a certain age from entering food or feed chains. WHO, FAO and OIE shall review this approach specifically in relation to public health issues.
- Countries should monitor the effective application of the regulatory measures that have been decided upon in particular the effectiveness of their ban on feeding ruminant tissues to ruminant, if in place.
- International trade in food products may disseminate tissues containing BSE. WHO, FAO and OIE should continue to work together to mitigate the risk of BSE agent further dissemination.

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Bibliography


Excerpt from “WHO recommended standards and strategies for surveillance, prevention and control of communicable diseases”