Mad Cow Disease

Mad Cow Disease, or bovine spongiform encephalopathy (BSE) is a brain disease of cattle first identified in the United Kingdom in the mid-1980s.

BSE is part of a group of diseases known as transmissible spongiform encephalopathies (TSEs). All of these diseases are characterized by distinctive changes in the brain, abnormal behavior and death.

What causes Mad Cow Disease (BSE)?
It is thought that this disease is caused and spread by an abnormal form of a protein called a “prion.” The abnormal protein triggers a chain-reaction causing other proteins in the brain to change to the abnormal form. Eventually these abnormal proteins accumulate in the brain, leading to the development of abnormal behavior and eventually death.

How do cattle get Mad Cow Disease (BSE)?
When cattle are slaughtered, portions of the animal not destined for human consumption may be used to produce a protein-rich byproduct that may be mixed with grain and fed back to livestock as feed. In the UK where the disease was first identified, it is thought that these by-products may have contained the prion protein, and that cattle became infected by consuming feed containing prion-contaminated animal protein. It is this reason that the feeding of animal proteins derived from cattle back to cattle has been banned in the U.S.

Are there other animal Transmissible Spongiform Encephalopathies (TSEs)?
Yes. There are other TSE diseases in animals and humans. Scrapie, one such prion-related disease, is a TSE that has been occurring in sheep and goats for at least 400 years. Despite close contact between people and sheep and goats, and the importance of lamb and mutton in many diets, there has been no evidence of human disease due to scrapie.

Chronic wasting disease (CWD) of deer and elk (cervids) is another TSE disease that has occurred in wild cervids in the Rocky Mountain states for many years and has recently been identified in Wisconsin. There are also TSEs in minks, cats and animals.

Are there TSEs which occur in humans?
Yes. There are a number of TSE diseases which occur in humans. One of the human diseases in this group is classic Creutzfeldt-Jakob disease (CJD). In the classic form of CJD, a chance (sporadic) mutation of normal brain protein into an abnormal protein occurs, which then leads to the formation of more abnormal protein.

CJD-like illness can also be familial, occurring in families where a genetic mutation favors the initial development of the abnormal protein. Also, there are cases of CJD that have followed transplantation of prion-infected cornea (eye) and other tissues, or the administration of hormones produced from human glands. More recently variant CJD (vCJD) has been identified. Variant CJD is a TSE disease in humans linked to eating meat products from cattle infected with Mad Cow Disease (BSE).

What are the symptoms of Creutzfeldt-Jakob Disease (CJD)?
Classic CJD is a human disease characterized by progressive dementia, leading to death. The disease may start with confusion or mental symptoms. Other symptoms are difficulty moving and muscle jerks. More and more signs and symptoms of brain damage and movement disorder follow until the patient goes into a coma and dies. Most of the cases of classic CJD have been diagnosed in people between the ages of 55 and 75.

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How long does it take to develop classic CJD?
Evidence suggests that it might take between 10 to 40 years to get sick with classic CJD once abnormal proteins have begun to develop. Classic CJD almost always occurs in people over 50 years old. Once someone gets classic CJD, they may live for six to 12 months.

How common is classic CJD?
Because classic CJD is caused by the chance, sporadic mutation of a normal brain protein into the prion, the risk of classic CJD is the same all over the world, about one per million persons per year.

What is variant-CJD (vCJD)?
Variant CJD is a disease in humans that has been linked to ingesting meat infected with the prion that causes mad cow disease (BSE). Variant CJD tends to occur in younger people (average age 29 rather than 65 for classic CJD) and has slightly different symptoms than classic CJD. People with vCJD tend to live slightly longer than people with classic CJD.

Variant CJD was first recognized in the UK in the mid-1990s, over ten years after the beginning of the BSE outbreak in cattle. Because of this and other similarities, vCJD is believed to be the result of consumption of prion-contaminated meat products. Muscle meat itself does not contain prions, but in processing of ground beef, sausage, and other meat products, contamination of the product with nervous tissues containing the infective prion may occur.

How common is vCJD, the human disease related to ingesting meat products infected with Mad Cow Disease (BSE)?
The risk of vCJD is thought to be related to exposure to animal products contaminated with the prion that causes BSE in cattle. Nearly all the cases of vCJD in humans have been in the UK or have been people with exposure to cattle meat products from the UK.

There have been 153 cases of vCJD reported in the world since the first case was identified in 1995, with 143 of these in the UK. Although many millions of people in the UK and around the world have been exposed to UK beef, and approximately 200,000 cases of BSE in cattle were reported and removed from the food supply over the past twenty years, the number of cases of vCJD remains low.

How do people get vCJD?
It is thought that a very small percentage of people who eat meat products infected with the BSE agent will get vCJD. Experiments done with cattle and other animals suggest that almost all of the infectivity is in tissue from the cattle's brain, spinal cord, and intestine, and that muscle meat itself is not infectious. Milk products from cattle are also not associated with any risk of infection with BSE.

What is being done in the U.S. to prevent BSE and vCJD?
The United States introduced a number of safeguards and surveillance efforts to lower the chances of BSE getting into the U.S. These included bans on use of certain animal proteins in animal feed, bans on UK beef products, and testing programs to quickly identify BSE in U.S. cattle.