Autism is another unusual sequel to HSE (Gillberg IC. Dev Med Child Neurol 1991;33:920-924).

NEUROLOGIC COMPLICATIONS OF SMALLPOX VACCINATION

Smallpox and smallpox vaccination is reviewed from the Departments of Neurology, Yale University School of Medicine, New Haven, CT, and University of New Mexico School of Medicine, Albuquerque. Neurological complications of smallpox vaccination are a postvaccinial encephalopathy (PVE) in children under 2 years of age and postvaccinial encephalomyelitis (PVEM) in recipients over 2 years. Past experience showed that the mortality of PVEM was 10 to 50%. The neuropathology of PVEM suggests an immune-mediated illness. Many unanswered questions need to be addressed regarding the risks of PVEM and PVE after smallpox vaccination with newer vaccines, neuroimaging findings, the prevention of these complications, and optimal therapy. (Booss J, Davis LE. Smallpox and smallpox vaccination. Neurological implications. Neurology April (2 of 2) 2003;60:1241-1245). (Reprints: Dr John Booss, Neurology Service (200), VA Connecticut Health Care System, 950 Campbell Ave, West Haven, CT 06516).

COMMENT. The risks of the smallpox vaccine are reviewed in an editorial (Johnson RT. Neurology April (2 of 2) 2003;60:1228-1229). Rates of PVEM vary from 1 in 4000 to 1 in 80,000 after primary vaccination, and from 1 in 50,000 to 1 in 450,000 after revaccination. Long-term disabilities of survivors may reach 30%. A vaccination campaign involving a previously unvaccinated population less than 35 years of age would carry the highest risk of neurologic complications. In addition, approximately 10% of the US population has some immunodeficiency, resulting in a greater risk of cutaneous dissemination.

The outcome of acute disseminated encephalomyelitis (ADEM) in a long-term follow-up study of 84 pediatric patients is covered in Ped Neur Briefs Nov 2002;16:81-82. Childhood ADEM affects boys more frequently than girls, and recovery occurs in 90%. Residual disability in 10% is not related to MRI lesions at onset but is correlated with the occurrence of optic neuritis. MRI shows bilateral, asymmetrical involvement of white matter of frontal and parietal lobes, lesions in deep grey matter including the thalamus, and corpus callosum and periventricular demyelination. The location of pathology based on MRI findings was similar to that described in ADEM following smallpox vaccination (Greenfield JG. Neuropathology London, Edward Arnold Publishers, 1958;201-205). Turnbull and McIntosh (1926) published the first description of the encephalomyelitis following vaccination against smallpox in 7 cases in London. These authors emphasized involvement of the ventral half of the pons. In Holland about the same time, 139 cases of post-vaccinial encephalomyelitis with 41 deaths were reported (Bouman and Bok (1927). Case-reports of similar lesions to post-vaccinial ADEM soon appeared following varicella (1927), measles (1926), influenza (1930), and smallpox (1927). An increase in strength of the vaccine and primary vaccination of many older children and adolescents were considered responsible for the epidemic of post-vaccinial cases in Great Britain and Holland in the years 1923-6. Glanzmann (1927) was the first to suggest that post-vaccinial ADEM was due to an antigen-antibody reaction.
**Human monkeypox virus infection.** The recent outbreak of Human Monkeypox Virus disease in Midwestern States and the governmental decision to recommend smallpox vaccination to persons exposed to prairie dogs or Gambian giant rats, pets linked to the infection, provide neurologists with an additional reason to consider the implications of smallpox vaccination. Human monkeypox is a rare zoonotic viral disease that occurs primarily in the rain forest of central and west Africa. Person-to-person spread may occur. The incubation period is up to 21 days. Fatalities in Africa have ranged from one percent to 10% of cases. Most patients seen in the US have experienced a prodrome of fever, headaches, myalgia, chills and sweats, followed by a nonproductive cough in one-third. A papular rash developed at one to 10 days after onset of fever and it progressed to vesiculation, pustulation, umbilication and crusting. Lesions occurred on head, trunk and extremities, including palms and soles. In the US, among 22 cases contracted from pet prairie dogs no fatalities have been reported. (Conover CS, Illinois Department of Public Health, Springfield, IL, June 8, 2003).

**NEUROLOGIC COMPLICATIONS OF ANTHRAX**

The literature on neurologic complications of infection by *Bacillus anthracis* is reviewed at the Dent Neurologic Institute, Buffalo, NY. The major neurologic complication is a rapidly fatal hemorrhagic meningoencephalitis. The initial mode of entry of the bacillus is via the cutaneous or inhalation route. One of the earliest cases of hemorrhagic meningoencephalitis from anthrax, in a boy aged 15 years, was characterized by numerous focal bleeds in the surface grey matter but very few hemorrhages in the white matter (House SJ. *J Infect Dis* 1920;27:513-526). House also reported autopsies on 2 adults with anthrax-related hemorrhagic meningitis, contracted in a Chicago factory by handling “curled hair” from South America. Other cases have involved contaminated heroin from Afghanistan, Pakistan, and Iran, in drug users, contaminated beef in slaughter house workers, and contact with sheep in farmers. CSF findings in 2 children with anthrax meningitis showed low glucose, increased protein and leucocytes, and large gram-positive rods without endospores on gram stain. CT scan shows multifocal areas of intracerebral hemorrhage. Inhalation cases of anthrax meningoencephalitis, affecting half the cases autopsied in the Russian 1979 outbreak, are particularly severe. In a recent report of fatal inhalation anthrax due to bioterrorism, the patient developed a generalized seizure within hours of admission and died on the third hospital day (Bush LM et al. *New Engl J Med* 2001;345:1607-1610). Death usually occurs within a week in cases of anthrax with neurologic involvement. Further research is needed to develop more rapid PCR methods to detect the disease in suspected cases. Antibiotic treatment with ciprofloxacin or doxycycline must be started immediately when the diagnosis is first apparent. (Meyer MA. Neurologic complications of anthrax. A review of the literature. *Arch Neurol* April 2003;60:483-488). (Reprints: Michel A Meyer MD, Dent Neurologic Institute, 3980 Sheridan Dr, Amherst, NY 14226).

COMMENT. Anthrax should be considered in the differential diagnosis in a patient with fever, dark necrotic pustules on the extremities, acute neurologic deterioration, gram-positive rods in the CSF, and multifocal intracerebral hemorrhages on CT.