Anisocoria, ataxia, dysarthria, and confusional state were predominant manifestations. Beta activity in the EEG has been described previously with attacks of basilar migraine.

Familial hemiplegic migraine and autosomal dominant arteriopathy with leukoencephalopathy (CADASIL) is described from St Vincent's Hospital, Dublin, Ireland. (Hutchinson M et al. <u>Ann Neurol</u> Nov 1995;38:817-824). Four subjects with CADSIL (cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy) had a history of familial hemiplegic migraine dating back to childhood. The disorder typically presents in adulthood but the MRI may show evidence of leukoencephalopathy before symptoms develop. This family is the first with both hemiplegic migraine and migraine as presenting symptoms of CADSIL.

## INFECTIOUS DISEASES

## BENIGN NEUROLOGIC COMPLICATIONS OF PERTUSSIS

The neurologic complications of pertussis infection among 340 unvaccinated patients admitted to hospital between 1979-1994 are reported from the Pediatric Clinic of the University of Catania, Sicily. Fourteen (4.1%) developed neurologic complications: Seizures occurred in all cases, 4 with fever. and 3 with signs of acute encephalopathy, including obtundation and vomiting which lasted only 12 to 24 hours. None of the patients developed epilepsy, all attend regular schools in appropriate grades, and at 14-18 year follow-up, only one has a mild behavioral disorder as a possible sequel of encephalopathy. No serious neurologic complications or permanent sequelae were observed in this series of children hospitalized patients with pertussis: a 15-year Sicilian experience. <u>Child's Nerv System</u> June 1996;12:332-335). (Respond: Dr G Incorpora, Clinica Pediatrica, Universita di Catania, Viale Andrea Doria, 6, 1-95125 Catania, Italy).

COMMENT. The relatively mild and benign nature of the neurologic complications of pertussis infection reported in this study contrast with the severity and permanent sequelae of some reported cases of pertussis vaccine encephalopathy. Seizures were not associated with anoxic episodes and coughing bouts and were not complicated by epilepsy.

## CHANGING PATTERNS OF REYE'S SYNDROME

Trends in the clinical pattern of Reye's syndrome in the British Isles between 1982 and 1990, and their relation to the June 1986 warnings against the use of aspirin in children, were analysed at the PHLS Communicable Disease Surveillance Centre, London, and other Centres in the UK. Of 445 cases reported, 354 had confirmed diagnoses and received scores of severity ranging from non-classical "Reye-like" (low scorers) to classical Reye's syndrome (high scorers). Classical cases occurred more frequently in the 4 1/2 year period before June 1986 compared with the subsequent period of surveillance. After June 1986, non-classical cases declined by 50% and classical by 79%. Classical, high scorers had received aspirin more frequently and were older than low scorers. (Hardie RM et al. Changing clinical pattern of Reye's syndrome. <u>Arch Dis Child</u> May 1996;74:400-405). (Respond: Dr Susan Hall, Floor C, Stephenson Building, Children's Hospital, Western Bank, Sheffield S10 27H, UK).