conclusions that factors other than birth asphyxia may be more important in etiology (see Progress in Pediatric Neurology III, PNB Publ, 1997:p377). The reliability of dried neonatal blood spot samples for immunoassay of inflammatory cytokines may permit a correlation between prenatal or perinatal infection and the development of childhood neurobehavioral disorders, including ADHD, often associated with subtle neurologic abnormalities.

Criteria for inclusion of cases in CP Registers are discussed in an article from the University of Western Australia (Badawi N, Stanley F et al. What constitutes cerebral palsy? Dev Med Child Neurol Aug 1998;40:520-527). As generally accepted, CP is a motor impairment resulting from brain pathology that is non-progressive and is manifested in early childhood. Exclusion criteria are those motor disorders caused by neurodegenerative diseases, neuromuscular disorders, spinal neural-tube defects, and brain tumors. The MRI and other laboratory investigations have permitted an etiological approach to classification of cases of CP that was formerly regarded as a "waste-basket" diagnosis.

ACUTE TRANSVERSE MYELITIS: CAUSES AND TREATMENT

Clinical manifestations, laboratory findings, management, and course of nine children diagnosed with acute transverse myelitis (ATM) between 1993 and 1996 are reported from the Children's University Hospital, Wurzburg, Germany. Spinal cord functions, including sensation, motor activity, and sphincter control, were affected in differing degrees, and neurologic symptoms and signs were often preceded by non-specific fever, nausea, and muscle pain. The peak of the illness and paraplegia was generally seen within 10 days and no later than 4 weeks. The thoracic spinal cord was principally involved in 80% of cases. Initially, tone and reflexes were decreased, and later, paralyses were associated with spasticity and hyperreflexia. CSF pleocytosis and/or elevated protein levels occurred in 4 patients. Peripheral nerve conduction velocity was normal, but muscle action potential amplitudes were decreased. An infectious causative agent was found in only 2 cases, with increasing antibody titers against echovirus 25 in one and Borrelia in one other. A para-, postinfective, postvaccinal myelitis was suspected in 7 cases. A review of the literature found bacterial, parasitic, and systemic lupus collagen disease as rare causes of ATM. A 3-day high-dose IV steroid pulse therapy (20 mg/kg/day prednisone) offered the most promising response in therapy. MRI excluded tumors, abscess, and vascular malformation. Multiple sclerosis, Guillain-Barre syndrome, and spinal cord infarction were more difficult to exclude. Of 6 patients followed up, 1 had a good recovery and 5 had a fair outcome. (Knebusch M, Strassburg HM, Reiners K. Acute transverse myelitis in childhood: nine cases and review of the literature. Dev Med Child Neurol September 1998;40:631-639). (Respond. Hans M Strassburg MD, Universitats Kinderklinik, Josef-Schneider-Str. 2 D-97080 Wurzburg, Germany).

COMMENT. Diagnostic criteria for acute transverse myelitis include: 1) acute paraplegia and sphincter disturbance with maximum impairment within 4 weeks; 2) bilateral segmental sensory impairment; 3) exclusion of spinal cord compression or systemic neurologic disease; and 4) consistent MRI and laboratory EMG findings. Prognosis is variable and residual sequelae are common.

ASPERGILLUS MYELOPATHY

Three children, ages 14 - 15 years, who developed myelopathy as the first manifestation of invasive aspergillosis are reported from the Children's Hospital Los Angeles, University of Southern California. All were immunosuppressed because of chemotherapy and antibiotics for treatment of leukemia. Symptoms