seizures (24%), neonatal (6%), generalized tonic-clonic (49%), and rare complex partial. Transmission of FC showed a maternal preponderance. The phenotypic expression of the genetic basis for focal sharp waves was remarkably variable. The clinical and EEG findings suggested a multifactorial pathogenesis for epilepsies characterized by benign focal sharp waves. (Doose H, Brigger-Heuer B, Neubauer B. Children with focal sharp waves: Clinical and genetic aspects. Epilepsia July 1997;38:788-796). (Reprints: Dr H Doose, Epilepsie-Zentrum, 24223 Raisdorf, Germany).

COMMENT. Focal sharp waves may be characteristic of a variety of different seizure patterns, including febrile convulsions, and not only that of rolandic epilepsy. In addition to epileptic syndromes, focal sharp waves may be found in children with learning disabilities such as dyslexia and dysphasia. The genetics of these syndromes appears to be multifactorial.

Genetics of febrile convulsions. Doose H and Maurer A have also reported on genetic factors in febrile convulsions. (Seizure risk in offspring of individuals with a history of febrile convulsions. Eur J Pediatr June 1997;156:476-481). The risk was 10%; only FC in 64% of affected offspring. Offspring of females with affected parents were at increased risk.

VAGUS NERVE STIMULATION FOR EPILEPSY
The Therapeutics and Technology Assessment Subcommittee of the American Academy of Neurology reports on vagus nerve stimulation (VNS) for epilepsy. A total of 104 articles and responses in 130 patients were reviewed. Data on VNS in children was sparse; 5 of 12 children with intractable partial, generalized, and mixed seizures, followed for 2 to 14 months in an open-label study, had a 90% reduction in seizure frequency. In a randomized prospective trial in 114 adult patients, high levels of stimulation showed a significant response compared to low stimulation. The degree of improvement was modest, with a mean partial seizure reduction of 31%, and 39% of patients having a >50% reduction in seizure frequency. Young patients with auras and idiopathic epilepsy had more than a 50% responder rate, while secondarily generalized seizures were not controlled. The mechanism of VNS was undetermined. Left-sided implantations were safer, since right VNS caused bradycardia. Electrodes around the vagus were connected to an infraclavicular generator pack. Serious complications were rare, but hoarseness, throat pain, and cough were common complaints during stimulation. The device is costly, and improvement in quality of life for the patient needs to be determined. (Fisher RS, Krauss GL, Ramsay E, Laxer K, Gates J. Assessment of vagus nerve stimulation for epilepsy: Report of the Therapeutics and Technology Assessment Subcommittee of the American Academy of Neurology. Neurology June 1997;49:293-297). (Reprints: Dr Robert S Fisher, Barrow Neurological Institute, 350 West Thomas Road, Phoenix, AZ 85013).

COMMENT. The panel considers vagal nerve stimulation a promising treatment for intractable partial seizures, but further controlled studies are recommended. Some predictors of a favorable response include age less than 34 years, idiopathic epilepsies, and early signs of effectiveness.

KETOGENIC DIET REVISITED IN THE LABORATORY
The effect of a ketogenic diet (KD) on kindled seizures and behavior in adult rats was studied at the Children's Hospital, Harvard Medical School, Boston, MA. In a kindling model, KD provided transient protection against
focal generation of seizures but not seizure spread. No change was noted in spatial learning and exploratory behavior tasks. (Hori A, Tandon P, Holmes GL, Stafstrom CE. Ketogenic diet: Effects on expression of kindled seizures and behavior in adult rats. Epilepsia July 1997;38:750-758). (Reprints: Dr CE Stafstrom, Division of Pediatric Neurology, #330, New England Medical Center, 750 Washington St, Boston, MA 02111).

COMMENT. The renewed interest in the clinical use of the ketogenic diet in the management of childhood epilepsy has stimulated further research in animal models. Some earlier studies of the effect of the KD on experimental seizures and seizure thresholds in laboratory animals showed that seizure susceptibility was not modified in normal animals, but an anticonvulsant effect of the KD was demonstrated in mice with seizure thresholds lowered by water intoxication and hypoelectrolytemia. (Millichap JG, Jones JD, Rudis BP. Mechanism of anticonvulsant action of the ketogenic diet. Amer J Dis Child 1962;104:506, and 1964;107:593-604). In animals and in children with absence epilepsy, the anticonvulsant effect of the KD was unrelated to diuresis, independent of acidosis and ketosis, similar to the effects of acetazolamide, and correlated most closely with a negative balance of sodium and potassium. Additional animal models were reported by Uhlemann ER, Neims AH, 1972; Appleton DB, DeVivo DC, 1974; and Nakazawa M et al, 1983.

Nordli DR Jr, and De Vivo DC provide an editorial commentary. (The ketogenic diet revisited: back to the future. Epilepsia July 1997;38:743-749). Further projected objectives of clinical trials and experimental models are discussed. We know that the ketogenic diet works, but how can we offer the diet or its specific ingredient in a more practical form? Clinical metabolic and laboratory studies should help to achieve this goal.

NEUROPATHIES

PERIPHERAL NEUROPATHY AND HIV INFECTION

The prevalence and patterns of peripheral neuropathy in 50 children with HIV infection were analyzed retrospectively in the EMG Section, National Institutes of Health, Bethesda, MD. Sensory complaints were the most common reason for referral for EMG. Of 19 children with neurological abnormalities, sensory loss or hypoactive reflexes, 10 had abnormal nerve conduction studies (NCS). In 7 children, NCSs were compatible with a distal symmetric sensory or sensorimotor axonal polyneuropathy (DSPN). All had advanced disease and had received antiretroviral agents; in one the neuropathy was linked to vincristine treatment. Three children had median nerve compression at the carpal tunnel; two had received AZT. Of 2 additional patients with abnormal NCS, one had a demyelinating neuropathy which improved after AZT, and one had a lumbosacral polyradiculopathy associated with varicella zoster involving legs and buttocks. Distal symmetric polyneuropathy affected older children. (Floeter MK, Civitello LA, Everett CR, Dambrosia J, Luciano CA. Peripheral neuropathy in children with HIV infection. Neurology July 1997;49:207-212). (Reprints: Dr MK Floeter, EMG Laboratory, NINDS, NIH, Bld 10 RM 5C101, 10 Center Drive MSC 1404, Bethesda, MD 20892).

COMMENT. The various patterns of peripheral neuropathy encountered in children with HIV resemble those described in adults. The most common type is a demyelinating sensory or sensorimotor polyneuropathy.