BIOGENIC AMINES IN RETT SYNDROME

Significant reductions in CSF metabolites of norepinephrine, dopamine, and serotonin are reported in 32 female patients with suspected Rett syndrome from the Baylor College of Medicine, Houston, Texas. CSF biopterin, an essential cofactor that may limit the synthesis of biogenic amines, was elevated in patients compared with controls. Diet, drugs, and nutritional status, that may affect monoamine metabolites, were thought to be unlikely explanations for these biochemical changes. (Zoghbi HY et al. Cerebrospinal fluid biogenic amines and biopterin in Rett syndrome. Ann Neurol Jan 1989;25:56-60).

COMMENT. Abnormal biochemical findings reported in Rett syndrome, including hyperammonemia, have not been substantiated. If specific and unexplained by anticonvulsant drugs or diet, this report is the first to suggest a metabolic disorder underlying the stereotypic hand movements and other neurologic signs of Rett syndrome. Abnormal CSF biogenic amines are also reported in Parkinson’s disease, Huntington’s chorea, and Lesch-Nyhan syndrome.

MENKES DISEASE WITH ‘RAGGED RED’ FIBERS

Subsarcolemmal aggregates of mitochondria (‘ragged red’ fibers) in skeletal muscle were found at autopsy in a 30-month-old male infant with Menkes kinky-hair disease reported from the New York Hospital-Cornell University Medical Center, New York. At birth, the infant had multiple depressed skull fractures and a cephalhematoma. At 5 days, he developed vertical nystagmus and staring episodes; at 2 months, generalized seizures; at 4 months, multifocal myoclonic twitches; and by 6 months, he had hypotonia, poor head control, and visual inattention. His height, weight, and head circumference were at the 3rd percentile. His hair was sparse, poorly pigmented, and showed pili torti (twisted), monilethrix (beadlike) and trichorrhexic nodosa (fractured nodes) on microscopic examination. The plasma copper was 16mcg/100ml (normal: approx 100mcg/100ml), and ceruloplasmin 12mg/dl (normal: 15-50mg/dl). He was treated with sodium valproate for seizures, and he had multiple episodes of vomiting, weight loss and dehydration, and respiratory infections. The immediate cause of death was bronchopneumonia. The brain weighed 500g (normal: 1100g) and showed diffuse cerebral and cerebellar atrophy, with focal polymicrogyria. Electronmicroscopy demonstrated numerous mitochondria within Purkinje cell cytoplasm. This report, “the first to describe ‘ragged red’ fibers in Menkes disease,” supports the concept that Menkes disease may be due in part to a mitochondrial enzyme deficiency. (Morgello S et al. Menkes kinky hair disease with ‘ragged red’ fibers. Dev Med Child Neurol Dec 1988;30:812-816).

COMMENT. Menkes kinky hair disease is an X-chromosome linked disorder of copper malabsorption characterized by low serum ceruloplasmin and copper levels, seizures, CNS degeneration, and pili torti (Menkes JH et al. Pediatrics 1962;29:764) (Menkes JH. Textbook of Child Neurology 3rd