Cranial computed tomography in disorders of complex carbohydrate metabolism and related storage diseases.

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Computed tomography (CT) was performed on 34 children with different disorders of complex carbohydrate metabolism and related storage diseases to obtain data on the degree of cerebral involvement. The main findings on CT were cerebral atrophy and hypodensity of the white matter. There was a great variability in these CT findings, even in siblings. Among the patients there were several in whom CT was normal, so a negative study does not exclude one of these disorders. These findings show that CT features such as cerebral atrophy or hypodensity are helpful in the evaluation of these disorders, though a diagnosis cannot be made on the basis of CT alone.

Patients with inborn errors of metabolism have a great variation in phenotype and symptomatology, which often makes diagnosis difficult. Recent computed tomographic (CT) reports have shown inconsistent results with different CT findings in such patients [1-9]. To test the diagnostic, differential diagnostic, and prognostic significance of the CT findings, and to obtain data of the degree of cerebral involvement in metabolic disorders, we investigated children with disorders of complex carbohydrate metabolism and related storage diseases.

Subjects and Methods

We examined 34 children with definite inborn errors by CT. Included in the series were 17 cases of mucopolysaccharidosis, two cases of oligosaccharidosis, three of mucolipidosis, nine of sphingolipidosis, and three of neuronal ceroid lipofuscinosis. In patients with defined enzyme defects the diagnoses were established biochemically. In patients suffering from neuronal ceroid lipofuscinosis, the diagnoses were verified histologically.

Clinical Description and Results

Mucopolysaccharidoses

The mucopolysaccharidoses are divided into six distinct entities and several subtypes based on biochemical, clinical, and genetic aspects [10]. The clinical symptomatology is due to the intralysosomal storage of glycosaminoglycans caused by the deficiency of a single lysosomal enzyme. Typical clinical features are dwarfism, organomegaly, joint contractures, thickened skin, skeletal abnormalities, and moderate to severe mental retardation. The involvement of neuronal tissue leads to progressive intellectual damage in most types. Recent investigations have shown a remarkable phenotypic variation even in those types that result from the same enzyme defect. On the other hand, clinically indistinguishable phenotypes originate from different enzyme defects, as seen in mucopolysaccharidosis III. Except for mucopolysaccharidosis II, the mode of inheritance is autosomal recessive [10-12].

There are other significant differences among the mucopolysaccharidoses that are obvious on CT [9]. These CT findings are partly (but not in every case) caused by the different ages of the patients or by different manifestations of the disease. Dilatation of the ventricles and of the subarachnoid spaces can occur. These findings become more distinct with increasing age. A density decrease of the white matter is possible in types IH, IIB, and VI. No disturbance of the blood-brain barrier is present. Therefore, no circumscribed enhancement after contrast injection is obvious on CT. Watts et al. [9] confirmed a normal blood-brain barrier with the aid of xenon investigations.

The variability of CT findings in mucopolysaccharidoses was shown in two sisters with mucopolysaccharidosis VI (fig. 1). CT findings of other types of mucopolysaccharidoses are represented in figures 2 and 3.

Oligosaccharidoses and Mucolipidoses

Some lysosomal storage disorders clinically resemble the sphingolipidoses and mucopolysaccharidoses. They formerly were designated as mucolipidoses [13]. After better understanding of the underlying biochemical defects they have been classified as the oligosaccharidoses (or glycoproteinoses), and as the mucolipidoses. In oligosaccharidoses one finds an elevated excretion of oligosaccharides in urine and a disturbance of glycoprotein catabolism; in mucolipidoses the catabolism of glycosaminoglycans, glycoproteins, and glycolipids is impaired [14]. The clinical picture of oligosaccharidoses and mucolipidoses results from the excessive and progressive intracellular accumulation of undegraded metabolites. Besides the typical symptoms like organomegaly, osteodysplasia, and dysmorphism, patients show a less or more pronounced mental deterioration. The inheritance is autosomal recessive.

Different CT changes can also be found in patients with oligosaccharidoses (mannosidosis, fucosidosis, aspartylglycosaminuria, sialidosis). In mannosidosis normal CT findings as well as dilatation of the subarachnoid spaces or density decrease of the white matter can occur. In fucosidosis, findings of a cortical and cerebellar

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Sphingolipidoses

Sphingolipidoses are inborn errors of lipid metabolism based on a lysosomal enzyme defect. In this group of disorders there is an abnormal intracellular accumulation of glycolipids, although the catabolism of glycosaminoglycans and glycolipids may be disturbed as well. Since glycolipids are essential in neuronal tissues, most of the patients develop a neurodegenerative symptomatology. Other leading symptoms are hepatosplenomegaly, ophthalmologic signs, and skeletal changes. There is a great clinical variability even within the same type of disorder. Except for the X-linked inherited Fabry disease (glycolipid lipidosis), the mode of inheritance is autosomal recessive [15].

In the group of sphingolipidoses, various CT findings are seen in metachromatic leukodystrophy. In this disease the ventricular system and the subarachnoid spaces may be quite normal, or there may be a significant dilatation of the ventricles and a distinct symmetrical density decrease of the white matter, especially in the frontal region. The CT changes are typical but not specific. Contrast enhancement does not occur [4].

In one of our female patients with metachromatic leukodystrophy, CT was performed at 10 and 15 years of age (fig. 4). Dilatation of the ventricles, cisterns, and sylvian fissures. Normal density of white matter.
Neuronal Cereoid Lipofuscinoses

Neuronal ceroid lipofuscinoses represent a group of neurodegenerative disorders in which four subtypes are differentiated by the age of onset. All patients except those with the adult type develop seizures, myoclonic jerks, dementia, and visual failure. In the infantile types the course is rapidly progressive; in the juvenile and adult type the course is slower. The mode of inheritance is autosomal recessive, although a dominant inheritance is described in some adult patients. The basic biochemical defect is unknown. Diagnosis is proven by analysis of electron microscopy of rectal or sural nerve biopsy. The leading histopathologic features are accumulation of typical ceroid lipofuscin in neuronal cells. The analysis of other tissues such as lymphocytes or fibroblasts seems to be as valid.

CT findings in neuronal ceroid lipofuscinosis vary [5, 8]. The ventricular system and the subarachnoid spaces may be normal or dilated. In general, cerebral atrophy is obvious without any correlation between the grade and the age of the process. The white matter has a normal density.

Discussion

CT findings must be considered in light of the clinical context. The different CT findings overlap considerably, restricting the differential diagnosis. The following conclusions can be made: (1) CT is an important diagnostic procedure in patients with suspected neurodegenerative diseases; (2) CT may alter the diagnostic workup of these patients, though a diagnosis cannot be made by CT alone; (3) CT follow-up may allow objective assessment of progression of the disease; and (4) if a definite biochemical diagnosis is not possible and a brain biopsy not available, CT may be contributory to the diagnosis.

REFERENCES