Optic nerve enlargement and leukodystrophy
An unusual finding of the infantile form of Krabbe disease

Antonio Milton Lima Garcia1, Norma Martins Menezes Morais1, Lygia Ohlweiler1, Maria Isabel Bragatti Winckler1, Josiane Ranzan3, Osvaldo Alfonso Pinto Artigalás2, Luise Lapagesse de Camargo Pinto2, Cristina Brinckmann Oliveira Netto3, Patrícia Ashton-Prolla4, Leonardo Vedolin5, Rudimar dos Santos Riesgo6, Newra Tellechea Rotta7

Leukodystrophies are a heterogeneous group of inherited neurological disorders characterized by progressive demyelination that causes loss of motor, sensory and intellectual functions leading to a fatal outcome. These disorders result from dysfunctions in myelin metabolism as a consequence of genetic enzymatic defects specific to each leukodystrophy subtype1. Krabbe disease (KD), also called globoid cell leukodystrophy (GCL), is inherited in an autosomal recessive pattern and has an estimated incidence of 1 in each 100,000/200,000 live births. It affects the peripheral and central nervous system (CNS)2.

We report on two cases of the infantile form of KD and describe MRI findings that suggest optic nerve enlargement (ONE).

CASES
Case 1
A 6-month-old white female patient was hospitalized due to respiratory infection symptoms. Her neuropsychological and motor development had been normal up to 3 months of age when she had bronchiolitis followed by progressive loss of developmental landmarks and slow progression of spasticity, but no seizures. The patient had a brother with a similar neurological condition that died at 6 months of age due to intestinal obstruction. Neurological examinations revealed generalized hypertonicity of the 4 limbs and decreased muscle mass, irritability and inconsolable crying, increased phasic-type myotatic reflexes and bilateral extensor plantar reflex. Cerebrospinal fluid was normal. Electroencephalogram showed signs of severe diffuse encephalopathy. Brain CT findings were suggestive of diffuse cerebral atrophy. MRI changes are described in Fig 1. Serum concentration of galactocerebrosidase was much lower than normal. The patient died due to respiratory complications.

Case 2
A 9-month-old white male patient presented with progressive loss of developmental landmarks, irritability, persistent crying, generalized hypertonicity and opisthotonos at seven months of age. The patient had dysphagia and fever, but no seizures. Perinatal and family histories were not relevant. Neurological examination showed excessive irritability, hypertonicity of the four limbs, hyperreflexia, bilateral extensor plantar reflex and normal fundoscopy findings. CSF showed discrete mononuclear pleocytosis and increase in
total protein content. Brain CT scan showed bilateral symmetrical hyperdensities located in the thalami, corona radiata and caudate nucleus. MRI findings are described in Fig 2. Serum B-galactocerebrosidase was not detected.

In both cases, the parents signed an informed consent for this publication.

DISCUSSION

KD is associated with a defective gene in the 14q31 chromosome, which leads to a reduction in the production of galactosylceramidase, a lysosomal enzyme responsible for the hydrolysis of galactosylceramide into ceramide and galactose. Consequently, galactocerebrosidase, as well as galactosylsphingosine, which causes cerebral toxicity, accumulate in macrophages. Their intracytoplasmic accumulation in macrophages and CSF cells leads to the formation of globoid cells, degeneration of oligodendrocytes, and, consequently, myelin deficiency and demyelination of white matter.

The onset of the infantile form of the disease, the most common subtype, occurs at about 6 months of age. Infants usually present with neurodegenerative symptoms that progress to spastic quadriplegia, tonic spasms, extreme irritability, inconsolable crying, secondary blindness, progressive optic atrophy, and death usually due to severe respiratory infection at about the second year of life. The juvenile and adult forms are characterized by insidious visual compromise, cognitive deterioration and gait disorders.

Brain CT scans in KD show characteristic hyperdensities in the thalami, the corona radiata, the body of the caudate nucleus, the cerebellum and the brainstem. Calcifications or globoid cell grouping may hypothetically explain the appearance of such images, but the real pathogenesis keep unexplained.

MRI shows signal abnormalities of similar distribution. Hypointense signs in T2-weighted images and hyperintense signs in T1-weighted images are seen in the early stages of the disease, but they later progress into hyperintense lesions on T2-weighted images, which suggest defective myelination and demyelination. In the cerebellum, the dentate nucleus and the white matter are typically affected. Clear cerebral and cerebellar atrophy can be seen in the late stages of the disease.

Recent case reports described MRI scans of the spinal cord of patients with KD and found, after contrast injection, diffuse enhancement at the nerve roots of the lumbar spine and low thoracic spine, which suggests the compromise of CNS at this level.

Histological findings reported in pioneering studies conducted in the last century showed optic nerve enlargement (ONE) in KD, particularly in its initial phase. ONE, which was found in the brain MRI scans of our two patients, was also described in recent studies that stressed...
the importance of these radiological findings. Hittmair et al. reported on the association of this finding with hypertrophy of other intracranial nerves. The following disorders should be included in the differential diagnosis of KD: optic nerve glioma with dural ectasia, frequently in type 1 neurofibromatosis; nerve sheath meningioma; granulomatous or histiocytic infiltration of optic nerves, leukemia, orbit pseudotumor, juvenile xanthogranuloma, post viral optic neuritis; optic nerve medulloepithelioma; and retinoblastoma with optic nerve compromise.

Spectroscopic findings in KD may include a low N-acetylaspartate (NAA) concentration, choline peaks (Cho), and an abnormally high Cho:NAA ratio in white matter. This reflects the presence of diffuse axonal degeneration and the proliferation of glial cells. Marked increases in the inositol (Ins)-to-NAA ratio is a sign of elevated astrocytosis in cerebral white matter.

Evaluation of the degree of anisotropy in the white matter of patients with KD using a new imaging method, diffusion tensor MR imaging (DTI), suggests that this method may detect changes in white matter earlier than MRI.

The diagnosis of KD is challenging because of the large number of demyelinating diseases and leukodystrophies, as well as the similarities between their clinical and imaging findings. Specific neuroimaging findings may help to establish this diagnosis. The detection of ONE on MRI scans, not found in other types of leukodystrophies, is an auxiliary tool in the differential diagnosis of KD.

In a patient with the clinical features of a metabolic disease, KD should be the main diagnostic hypothesis whenever there is evidence of ONE, particularly when associated with their typical intracranial neuroimaging findings.

REFERENCES