COMPOUND GALACTOSYLCERAMIDASE GENE (GALC) HETEROZYGOSITY IN A BOY WITH INFANTILE KRABBE DISEASE (KD)

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Abstract
Krabbe disease (KD) (globoid cell leukodystrophy) is a degenerative, lysosomal storage disease, caused by a severe loss of galactocerebrosidase (GALC) enzymatic activity. The inheritance is autosomal recessive. KD affects the white matter of the central and peripheral nervous systems. We present a 3 year old boy in whom the disease had an 'infantile' or 'classic' presentation, with spasticity, irritability, and developmental delay. In addition the boy showed progressive severe motor and mental deterioration, difficulties in swallowing and decerebration. Molecular analysis revealed that the child is a compound heterozygote: p.Asp187Val (c.560A>T) and p.Ile250Thr (c.749T>C). The father was the carrier of p.Asp187Val (c.560A>T), while the mother was the carrier of the p.Ile250Thr (c.749T>C) in exon 6 of the GALC gene. The clinical course in this compound heterozygote is severe and the patient passed away at the age of 3 years. Genotype-phenotype relations are discussed in this Macedonian patient with KD.

Key words: Krabbe disease, infantile form, compound heterozygote.

Introduction
The infantile or classic KD is a rapidly progressive disease with severe motor and mental deterioration, decerebration and death by age 2 years [1, 2]. However, there is a late onset form (late-infantile 6 months to 3 years, juvenile 3 to 8 years, and even adult-onset forms) in 10 to 15% of patients, with milder clinical course. Interestingly, even within families the disease severity is variable [3, 4].

We describe a compound heterozygote GALC patient with infantile form of KD.

Case report
This is a 12 month old boy, the first child of unrelated and healthy parents. The pregnancy and delivery (40th gestational week) were uneventful. Birth weight was 3370 g, birth length 52 cm, head circumference was not available, and the Apgar score was 9/10.

The parents noticed hypotony, the child was unable to hold his head or to seat at the age of 8 months. The feeding was difficult as the swallowing was affected. At the physical examination performed at the age of 8 months there was a significant hypotony, hyperextension of the axial muscles, the arms were in flexion, the hands clenched in fists, the legs were in a total extension with the plantar flexion, while the face was expressionless. No contact with people around him was possible.

The lumbar puncture showed normal cerebrospinal liquor without cells and with normal values of the glucose and the proteins. The electroencephalogram showed asymmetry of the background activity. The eye fundus examination was normal. The ultrasound examination of the brain, heart, kidneys was normal. The TORCH had normal values. The Amino acids in the blood were within the normal
range. The first MRI of the brain revealed enlarged subarachnoid spaces and secondary dilatation of the third and fourth ventricle. The bilateral symmetrical hyperintense areas were also present at the periventricular white matter.

The results of β-galactocerebrosidase activity measured in the blood leukocytes revealed low values (6 nmoles/17 h/mg proteins (normal value: 15 to 53).

The molecular genetic analysis was performed in Graz, Austria. A previously described mutation p.Asp187Val [5] was found in Exon 6 of the GALC-gene and a p.Ile250Thr [6] in Exon 8 on the same GALC-genes.

His spasticity, irritability, and developmental delay progressed. In addition, the boy showed progressive severe motor and mental deterioration, difficulties in swallowing and de cerebration. Ha passed away at the age of 3 years.

Discussion

The prevalence of KD is estimated to be about 1 in 100,000 (1.0 x 10⁻⁵) [4]: from 0.40 in the Czech Republic to 1.35 in the Netherlands. Higher frequency of the gene was found in Sicily [7], in the Druze population [8], in two inbred communities in Israel.

The infantile or classic form is the most severe with progressive and extreme irritability and spasticity, as well as motor and mental delay [9]. The described boy had this very lethal form. The late-onset form patients have a longer course of the disease and survival into childhood and adolescence [4, 10–14]. The late form of KB is predominant, found in 66% [15] with the oldest observed in 84 years old woman.

Severe compound heterozygotes have been described [6, 15–17]. The phenotype genotype relation is difficult to be established. Milder phenotype was found in the missense mutations G286D [4], I66M + I289V, G270D, and L618S [18]. The G41S mutation inferred a longer disease course [15]. The location of the mutations in the N- or C-terminus was more frequent in the adult-onset form, while the central domain mutations were predominantly found in the infantile form of KD [19]. About 45% of mutated alleles in patients of European origin have a 30 kb deletion covering exons 11–17 and inferred the classic infantile form of GLD. The late onset GLD was found with the c.857G>A (p.Gly286Asp in exon 8) mutation. There is an overlap of the measured GALC activity in blood or in skin fibroblasts between the control groups and the carriers, which renders the diagnosis challenging. Our patient was compound heterozygous with the genotype which proved to be severe: p.Asp187Val (c.560A>T) and p.Ile250Thr (c.749T>C).

There is not a causative treatment. Chaperones are promising, while the allogeneic hematopoietic stem cell transplantation is of uncertain effect [20]. The transplantation of umbilical-cord blood in patients with infantile KD was not successful [21].

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Compound galactosylceramidase gene (GALC) heterozygosity…