What is D-bifunctional protein deficiency?

D-bifunctional protein deficiency is an inherited disease characterized by neonatal low muscle tone, seizures, visual and hearing loss, developmental delays, and death usually by two years of age. Individuals with D-bifunctional protein deficiency have defects in the D-bifunctional protein that is involved in the breakdown of a very long-chain fatty acids (VLCFA). The symptoms of D-bifunctional protein deficiency are due to the toxic buildup of VLCFA, which causes damage in the cells, especially in the brain and liver. D-bifunctional protein deficiency is also known as peroxisomal bifunctional enzyme deficiency and pseudo-Zellweger syndrome.

What are the symptoms of D-bifunctional protein deficiency and what treatment is available?

Signs and symptoms of D-bifunctional protein deficiency are usually evident within the first month of life and may include:

- Hypotonia (low muscle tone)
- Seizures
- Hepatomegaly (enlarged liver) and liver disease
- Vision and hearing problems
- Unusual facial features
- Intellectual disability

There is no cure for D-bifunctional protein deficiency. Treatment is supportive care for symptoms. D-bifunctional protein deficiency is usually fatal by the age of two years, with survival rarely to later childhood.

How is D-bifunctional protein deficiency inherited?

D-bifunctional protein deficiency is an autosomal recessive disease caused by mutations in the HSD17B4 gene. An individual who inherits one HSD17B4 gene mutation is a carrier and is not expected to have related health problems. An individual who inherits two HSD17B4 gene mutations, one from each parent, is expected to be affected with D-bifunctional protein deficiency.

If both members of a couple are carriers of an HSD17B4 gene mutation, the risk for an affected child is 25% in each pregnancy; therefore, it is especially important that the reproductive partner of a carrier be offered testing.

Who is at risk for D-bifunctional protein deficiency?

D-bifunctional protein deficiency can occur in individuals of all races and ethnicities, and it is estimated to affect 1 in 100,000 individuals.

Having a relative who is a carrier or is affected can increase an individual’s risk of being a carrier. Consultation with a genetics health professional may be helpful in determining carrier risk and appropriate testing.

What does a positive test result mean?

If a gene mutation is identified, an individual should speak to a physician or genetics health professional about the implications of the result and appropriate testing for the reproductive partner and at-risk family members.

What does a negative test result mean?

A negative result reduces, but does not eliminate, the possibility that an individual carries a gene mutation. The likelihood of being a carrier is also influenced by family history, medical symptoms, and other relevant test results.
D-bifunctional protein deficiency

Where can I get more information?

Genetics and Rare Disease Information (GARD):
http://rarediseases.info.nih.gov/GARD/Condition/4539/Dbifunctional_protein_deficiency.aspx/ShowAll


References