What is Alpha-Mannosidosis?
Alpha-mannosidosis is an inherited disease characterized by developmental delays, facial and skeletal abnormalities, hearing loss, and immune deficiency. It is caused by abnormalities in the enzyme alpha-mannosidase that breaks down small sugar molecules called oligosaccharides. Symptoms associated with alpha-mannosidosis are due to the toxic build-up of oligosaccharides and the progressive destruction of cells, particularly in the central nervous system.

What are the symptoms of Alpha-Mannosidosis and what treatment is available?
Alpha-mannosidosis is a disease that varies in severity and age at presentation, even within families. Symptoms may include:
- Progressive myopathy (muscle weakness and pain)
- Ataxia (difficulty controlling movements)
- Developmental delays
- Mental retardation
- Hearing loss
- Distinctive facial characteristics (large head, prominent forehead, low hairline, large ears, protruding jaw, and widely spaced teeth)
- Scoliosis (abnormal curvature of the spine)
- Joint and bone abnormalities (types 2 and 3)
- Frequent infections
- Psychiatric disease (confusion, anxiety, depression, hallucinations)

There have been three types of alpha-mannosidosis described. Most individuals experience symptoms before age 10, with a slow progression (type 2). Some individuals present after age 10, show milder symptoms, no skeletal abnormalities and even slower progression (type 1). A severe form may also occur, however, with onset during infancy, a rapid progression, and early death (type 3).

There is no cure for alpha-mannosidosis. Many individuals live to more than 50 years of age (types 1 & 2); however, the long-term prognosis is poor with the majority being wheelchair-dependent. Treatment is supportive and includes antibiotics to fight infections, hearing aids, and surgery if needed for bone abnormalities. Bone marrow transplant may be beneficial in some individuals if alpha-mannosidosis is diagnosed before the onset of neurological symptoms.

How is Alpha-Mannosidosis inherited?
Alpha-mannosidosis is an autosomal recessive disease caused by mutations in the MAN2B1 gene. An individual who inherits one copy of a MAN2B1 gene mutation is a “carrier” and is not expected to have health related problems. An individual who inherits two mutations in this gene, one from each parent, is expected to be affected with alpha-mannosidosis.

If both members of a couple are carriers, 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 Alpha-Mannosidosis?
Alpha-mannosidosis can occur in individuals of all races and ethnicities. The incidence of alpha-mannosidosis is approximately 1 in 500,000, with a calculated carrier frequency of approximately 1 in 350.
Having a relative who is a carrier or who is affected can also 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 a 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 medical tests.

**Where can I get more information?**

- ISMRD: The International Advocate for Glycoprotein Storage Diseases: [http://www.ismrd.org/the_diseases/alpha_mannosidosis](http://www.ismrd.org/the_diseases/alpha_mannosidosis)
- CLIMB: Children Living with Inherited Metabolic diseases: [http://www.climb.org.uk/IMD/Mike/MannosidosisAlpha.pdf](http://www.climb.org.uk/IMD/Mike/MannosidosisAlpha.pdf)

**References**