Alpha-methylacyl-CoA racemase deficiency

Alpha-methylacyl-CoA racemase (AMACR) deficiency is a disorder that causes a variety of neurological problems that begin in adulthood and slowly get worse. People with AMACR deficiency may have a gradual loss in intellectual functioning (cognitive decline), seizures, and migraines. They may also have acute episodes of brain dysfunction (encephalopathy) similar to stroke, involving altered consciousness and areas of damage (lesions) in the brain. Other features of AMACR deficiency may include weakness and loss of sensation in the limbs due to nerve damage (sensorimotor neuropathy), muscle stiffness (spasticity), and difficulty coordinating movements (ataxia). Vision problems caused by deterioration of the light-sensitive layer at the back of the eye (the retina) can also occur in this disorder.

Frequency

AMACR deficiency is a rare disorder. Its prevalence is unknown. At least 10 cases have been described in the medical literature.

Causes

AMACR deficiency is caused by mutations in the AMACR gene. This gene provides instructions for making an enzyme called alpha-methylacyl-CoA racemase (AMACR).

The AMACR enzyme is found in the energy-producing centers in cells (mitochondria) and in cell structures called peroxisomes. Peroxisomes contain a variety of enzymes that break down many different substances, including fatty acids and certain toxic compounds. They are also important for the production (synthesis) of fats (lipids) used in digestion and in the nervous system. In peroxisomes, the AMACR enzyme plays a role in the breakdown of a fatty acid called pristanic acid, which comes from meat and dairy foods in the diet. In mitochondria, AMACR is thought to help further break down the molecules derived from pristanic acid.

Most individuals with AMACR deficiency have an AMACR gene mutation that results in a lack (deficiency) of functional enzyme. The enzyme deficiency leads to accumulation of pristanic acid in the blood. However, it is unclear how this accumulation is related to the specific signs and symptoms of AMACR deficiency.

Inheritance Pattern

This condition is inherited in an autosomal recessive pattern, which means both copies of the gene in each cell have mutations. The parents of an individual with an autosomal recessive condition each carry one copy of the mutated gene, but they typically do not show signs and symptoms of the condition.
Other Names for This Condition

• AMACR deficiency

Diagnosis & Management

Genetic Testing Information

• What is genetic testing?
  /primer/testing/genetictesting

• Genetic Testing Registry: Alpha-methylacyl-CoA racemase deficiency

Research Studies from ClinicalTrials.gov

• ClinicalTrials.gov
  https://clinicaltrials.gov/ct2/results?cond=%22alpha-methylacyl-CoA+racemase+deficiency%22+OR+%22Peroxisomal+Disorders%22

Other Diagnosis and Management Resources

• Kennedy Krieger Institute: Peroxisomal Diseases
  https://www.kennedykrieger.org/patient-care/centers-and-programs/genetics-laboratories/about/peroxisomal-diseases

Additional Information & Resources

Health Information from MedlinePlus

• Health Topic: Degenerative Nerve Diseases
  https://medlineplus.gov/degenerativenervediseases.html

• Health Topic: Genetic Brain Disorders
  https://medlineplus.gov/geneticbraindisorders.html

Educational Resources

• MalaCards: alpha-methylacyl-coa racemase deficiency
  https://www.malacards.org/card/alpha_methylacyl_coa_racemase_deficiency

• Merck Manual for Health Care Professionals: Peroxisomal Disorders
  https://www.merckmanuals.com/professional/pediatrics/inherited-disorders-of-metabolism/peroxisomal-disorders

Patient Support and Advocacy Resources

• Global Foundation for Peroxisomal Disorders
  https://www.thegfpd.org/

• Metabolic Support UK
  https://www.metabolicsupportuk.org/
Sources for This Summary


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