Hypomyelination and congenital cataract

Hypomyelination and congenital cataract is an inherited condition that affects the nervous system and the eyes. This disease is one of a group of genetic disorders called leukoencephalopathies. Leukoencephalopathies involve abnormalities of the brain’s white matter. White matter consists of nerve fibers covered by a fatty substance called myelin. Myelin insulates nerve fibers and promotes the rapid transmission of nerve impulses. Hypomyelination and congenital cataract is caused by a reduced ability to form myelin (hypomyelination). Additionally, people with this disorder are typically born with a clouding of the lens (cataract) in both eyes.

People with this condition usually have normal development throughout the first year of life. Development slows around the age of 1. Most affected children learn to walk between the ages of 1 and 2, although they usually need some type of support. Over time they experience muscle weakness and wasting (atrophy) in their legs, and many affected people eventually require wheelchair assistance. Weakness in the muscles of the trunk and a progressive abnormal curvature of the spine (scoliosis) further impair walking in some individuals. Most people with hypomyelination and congenital cataract have reduced sensation in their arms and legs (peripheral neuropathy). In addition, affected individuals typically have speech difficulties (dysarthria) and mild to moderate intellectual disability.

Frequency

The prevalence of hypomyelination and congenital cataract is unknown.

Causes

Mutations in the FAM126A gene cause hypomyelination and congenital cataract. The FAM126A gene provides instructions for making a protein called hyccin, the function of which is not completely understood. Based on the features of hypomyelination and congenital cataract, researchers presume that hyccin is involved in the formation of myelin throughout the nervous system. Hyccin is also active in the lens of the eye, the heart, and the kidneys. It is unclear how mutations in the FAM126A gene cause cataracts.

Most FAM126A gene mutations that cause hypomyelination and congenital cataract prevent the production of hyccin. People who cannot produce any hyccin have problems forming myelin, leading to the signs and symptoms of this condition.

People who have mutations that allow some protein production tend to have milder symptoms than those who produce no protein. These individuals typically retain the ability to walk longer, although they still need support, and they usually do not have peripheral neuropathy.
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

- HCC

Diagnosis & Management

Genetic Testing Information

- What is genetic testing?
  https://primer/testing/genetictesting
- Genetic Testing Registry: Hypomyelination and Congenital Cataract

Other Diagnosis and Management Resources

- GeneReview: Hypomyelination and Congenital Cataract
  https://www.ncbi.nlm.nih.gov/books/NBK2587
- MedlinePlus Encyclopedia: Congenital Cataract
  https://medlineplus.gov/ency/article/001615.htm
- MedlinePlus Encyclopedia: Muscle Atrophy
  https://medlineplus.gov/ency/article/003188.htm

Additional Information & Resources

Health Information from MedlinePlus

- Encyclopedia: Congenital Cataract
  https://medlineplus.gov/ency/article/001615.htm
- Encyclopedia: Muscle Atrophy
  https://medlineplus.gov/ency/article/003188.htm
- Health Topic: Leukodystrophies
  https://medlineplus.gov/leukodystrophies.html
- Health Topic: Peripheral Nerve Disorders
  https://medlineplus.gov/peripheralnervedisorders.html
- Health Topic: Scoliosis
  https://medlineplus.gov/scoliosis.html
Genetic and Rare Diseases Information Center

- Hypomyelination and congenital cataract
  https://rarediseases.info.nih.gov/diseases/11980/hypomyelination-and-congenital-cataract

Additional NIH Resources

- National Eye Institute: Cataract
- National Institute of Neurological Disorders and Stroke: Peripheral Neuropathy Fact Sheet
  https://www.ninds.nih.gov/Disorders/All-Disorders/Peripheral-Neuropathy-Information-Page

Educational Resources

- Boston Children's Hospital: Cataracts in Children
  http://www.childrenshospital.org/conditions-and-treatments/conditions/c/cataracts
- Boston Children's Hospital: Nervous System Disorders in Children
  http://www.childrenshospital.org/conditions-and-treatments/conditions/n/nervous-system-disorders
- Merck Manual Home Edition for Patients and Caregivers: Cataract
  https://www.merckmanuals.com/home/eye-disorders/cataract/cataract
- Orphanet: Hypomyelination-congenital cataract syndrome
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=85163

Patient Support and Advocacy Resources

- Prevent Blindness America: Cataract
  https://www.preventblindness.org/cataract
- The Foundation Fighting Blindness
  https://www.fightingblindness.ca/
- The Myelin Project
  https://www.myelin.org/
- University of Kansas Medical Center Resource List: Developmental Delay/Mental Retardation
  http://www.kumc.edu/gec/support/devdelay.html

Clinical Information from GeneReviews

- Hypomyelination and Congenital Cataract
  https://www.ncbi.nlm.nih.gov/books/NBK2587
Scientific Articles on PubMed

• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28hypomyelination%5BTIAB
%5D%29+AND+%28congenital+cataract%29%29+AND+english%5Bla%5D+AND
+human%5Bmh%5D+AND+%22last+3600+days%22%5Bdp%5D

Catalog of Genes and Diseases from OMIM

• LEUKODYSTROPHY, HYPOMYELINATING, 5
  http://omim.org/entry/610532

Medical Genetics Database from MedGen

• Hypomyelination and Congenital Cataract

Sources for This Summary

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  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17683097

  Hypomyelination and congenital cataract: neuroimaging features of a novel inherited white matter
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17974614

• Ugur SA, Tolun A. A deletion in DRCTNNB1A associated with hypomyelination and juvenile onset
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17928815

  MS, Lisanti MP, Minetti C. Deficiency of hyccin, a newly identified membrane protein, causes
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16951682

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