Complete LCAT deficiency

Complete LCAT deficiency is a disorder that primarily affects the eyes and kidneys.

In complete LCAT deficiency, the clear front surface of the eyes (the corneas) gradually becomes cloudy. The cloudiness, which generally first appears in early childhood, consists of small grayish dots of cholesterol (opacities) distributed across the corneas. Cholesterol is a waxy, fat-like substance that is produced in the body and obtained from foods that come from animals; it aids in many functions of the body but can become harmful in excessive amounts. As complete LCAT deficiency progresses, the corneal cloudiness worsens and can lead to severely impaired vision.

People with complete LCAT deficiency often have kidney disease that begins in adolescence or early adulthood. The kidney problems get worse over time and may eventually lead to kidney failure. Individuals with this disorder also usually have a condition known as hemolytic anemia, in which red blood cells are broken down (undergo hemolysis) prematurely, resulting in a shortage of red blood cells (anemia). Anemia can cause pale skin, weakness, fatigue, and more serious complications.

Other features of complete LCAT deficiency that occur in some affected individuals include enlargement of the liver (hepatomegaly), spleen (splenomegaly), or lymph nodes (lymphadenopathy) or an accumulation of fatty deposits on the artery walls (atherosclerosis).

Frequency

Complete LCAT deficiency is a rare disorder. Approximately 70 cases have been reported in the medical literature.

Causes

Complete LCAT deficiency is caused by mutations in the LCAT gene. This gene provides instructions for making an enzyme called lecithin-cholesterol acyltransferase (LCAT).

The LCAT enzyme plays a role in removing cholesterol from the blood and tissues by helping it attach to molecules called lipoproteins, which carry it to the liver. Once in the liver, the cholesterol is redistributed to other tissues or removed from the body. The enzyme has two major functions, called alpha- and beta-LCAT activity. Alpha-LCAT activity helps attach cholesterol to a lipoprotein called high-density lipoprotein (HDL). Beta-LCAT activity helps attach cholesterol to other lipoproteins called very low-density lipoprotein (VLDL) and low-density lipoprotein (LDL).

LCAT gene mutations that cause complete LCAT deficiency either prevent the production of LCAT or impair both alpha-LCAT and beta-LCAT activity, reducing the
enzyme’s ability to attach cholesterol to lipoproteins. Impairment of this mechanism for reducing cholesterol in the body leads to cholesterol deposits in the corneas, kidneys, and other tissues and organs. LCAT gene mutations that affect only alpha-LCAT activity cause a related disorder called fish-eye disease that affects only the corneas.

**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**

- familial LCAT deficiency
- familial lecithin-cholesterol acyltransferase deficiency
- FLD
- LCAT deficiency
- lecithin acyltransferase deficiency
- lecithin:cholesterol acyltransferase deficiency
- Norum disease
- Norum's disease

**Diagnosis & Management**

**Genetic Testing Information**

- What is genetic testing?  
  /primer/testing/genetictesting
- Genetic Testing Registry: Norum disease  

**Research Studies from ClinicalTrials.gov**

- ClinicalTrials.gov  
  https://clinicaltrials.gov/ct2/results?cond=%22Norum+disease%22+OR+%22Lecithin+acyltransferase+deficiency%22+OR+%22Lecithin%3Acholesterol+acyltransferase+deficiency%22

**Other Diagnosis and Management Resources**

- MedlinePlus Encyclopedia: Corneal Transplant  
  https://medlineplus.gov/ency/article/003008.htm
- National Heart, Lung, and Blood Institute: Hemolytic Anemia  
  https://www.nhlbi.nih.gov/health-topics/hemolytic-anemia
• National Institutes of Diabetes and Digestive and Kidney Diseases: Kidney Failure
  -- Choosing a Treatment That's Right for You
  choosing-treatment

• Oregon Health and Science University: Corneal Dystrophy
  https://www.ohsu.edu/casey-eye-institute/corneal-disease

**Additional Information & Resources**

**Health Information from MedlinePlus**

• Encyclopedia: Corneal Transplant
  https://medlineplus.gov/ency/article/003008.htm

• Encyclopedia: Hemolytic Anemia
  https://medlineplus.gov/ency/article/000571.htm

• Health Topic: Corneal Disorders
  https://medlineplus.gov/cornealdisorders.html

• Health Topic: Kidney Diseases
  https://medlineplus.gov/kidneydiseases.html

**Genetic and Rare Diseases Information Center**

• Familial LCAT deficiency
  https://rarediseases.info.nih.gov/diseases/4011/familial-lcat-deficiency

**Additional NIH Resources**

• National Eye Institute: Corneal Conditions
  corneal-conditions

• National Heart, Lung, and Blood Institute: Hemolytic Anemia
  https://www.nhlbi.nih.gov/health-topics/hemolytic-anemia

• National Institutes of Diabetes and Digestive and Kidney Diseases: Kidney Failure
  -- Choosing a Treatment That's Right for You
  choosing-treatment

**Educational Resources**

• MalaCards: familial lcat deficiency
  https://www.malacards.org/card/familial_lcat_deficiency

• Orphanet: Familial LCAT deficiency
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=79293
Patient Support and Advocacy Resources

• American Foundation for the Blind
  https://www.afb.org/
• National Kidney Foundation
  https://www.kidney.org/
• Royal National Institute of Blind People: Corneal Dystrophies
  https://www.rnib.org.uk/eye-health/eye-conditions/corneal-dystrophies

Scientific Articles on PubMed

• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28Lecithin+Acyltransferase+Deficiency%5BMAJR%5D%29+OR+%28Norum+disease%5BALL%5D%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+1800+days+22%5Bdp%5D

Catalog of Genes and Diseases from OMIM

• LECITHIN:CHEOLESTROL ACYLTRANSFERASE DEFICIENCY
  http://omim.org/entry/245900

Sources for This Summary

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