Ocular albinism

Ocular albinism is a genetic condition that primarily affects the eyes. This condition reduces the coloring (pigmentation) of the iris, which is the colored part of the eye, and the retina, which is the light-sensitive tissue at the back of the eye. Pigmentation in the eye is essential for normal vision.

Ocular albinism is characterized by severely impaired sharpness of vision (visual acuity) and problems with combining vision from both eyes to perceive depth (stereoscopic vision). Although the vision loss is permanent, it does not worsen over time. Other eye abnormalities associated with this condition include rapid, involuntary eye movements (nystagmus); eyes that do not look in the same direction (strabismus); and increased sensitivity to light (photophobia). Many affected individuals also have abnormalities involving the optic nerves, which carry visual information from the eye to the brain.

Unlike some other forms of albinism, ocular albinism does not significantly affect the color of the skin and hair. People with this condition may have a somewhat lighter complexion than other members of their family, but these differences are usually minor.

The most common form of ocular albinism is known as the Nettleship-Falls type or type 1. Other forms of ocular albinism are much rarer and may be associated with additional signs and symptoms, such as hearing loss.

Frequency

The most common form of this disorder, ocular albinism type 1, affects at least 1 in 60,000 males. The classic signs and symptoms of this condition are much less common in females.

Causes

Ocular albinism type 1 results from mutations in the GPR143 gene. This gene provides instructions for making a protein that plays a role in pigmentation of the eyes and skin. It helps control the growth of melanosomes, which are cellular structures that produce and store a pigment called melanin. Melanin is the substance that gives skin, hair, and eyes their color. In the retina, this pigment also plays a role in normal vision.

Most mutations in the GPR143 gene alter the size or shape of the GPR143 protein. Many of these genetic changes prevent the protein from reaching melanosomes to control their growth. In other cases, the protein reaches melanosomes normally but mutations disrupt the protein's function. As a result of these changes, melanosomes in skin cells and the retina can grow abnormally large. Researchers are uncertain how these giant melanosomes are related to vision loss and other eye abnormalities in people with ocular albinism.
Rare cases of ocular albinism are not caused by mutations in the \textit{GPR143} gene. In these cases, the genetic cause of the condition is often unknown.

\textbf{Inheritance Pattern}

Ocular albinism type 1 is inherited in an X-linked pattern. A condition is considered X-linked if the mutated gene that causes the disorder is located on the X chromosome, one of the two sex chromosomes. In males (who have only one X chromosome), one altered copy of the \textit{GPR143} gene in each cell is sufficient to cause the characteristic features of ocular albinism. Because females have two copies of the X chromosome, women with only one copy of a \textit{GPR143} mutation in each cell usually do not experience vision loss or other significant eye abnormalities. They may have mild changes in retinal pigmentation that can be detected during an eye examination.

\textbf{Other Names for This Condition}

- albinism, ocular
- OA
- XLOA

\textbf{Diagnosis & Management}

\textbf{Genetic Testing Information}

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

\textbf{Research Studies from ClinicalTrials.gov}

- ClinicalTrials.gov https://clinicaltrials.gov/ct2/results?cond=%22ocular+albinism%22

\textbf{Other Diagnosis and Management Resources}

Additional Information & Resources

Health Information from MedlinePlus
- Encyclopedia: Albinism
  https://medlineplus.gov/ency/article/001479.htm
- Health Topic: Eye Diseases
  https://medlineplus.gov/eyediseases.html
- Health Topic: Vision Impairment and Blindness
  https://medlineplus.gov/visionimpairmentandblindness.html

Genetic and Rare Diseases Information Center
- Albinism ocular late onset sensorineural deafness
  https://rarediseases.info.nih.gov/diseases/592/albinism-ocular-late-onset-sensorineural-deafness
- Ocular albinism type 1
  https://rarediseases.info.nih.gov/diseases/8471/ocular-albinism-type-1

Educational Resources
- MalaCards: ocular albinism
  https://www.malacards.org/card/ocular_albinism
- Orphanet: Ocular albinism
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=284804
- Orphanet: X-linked recessive ocular albinism
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=54

Patient Support and Advocacy Resources
- National Organization for Rare Disorders
  https://rarediseases.org/rare-diseases/ocular-albinism/
- Resource list from the University of Kansas Medical Center
  http://www.kumc.edu/gec/support/albinism.html
- The Vision of Children
  https://www.visionofchildren.org/

Clinical Information from GeneReviews
- Ocular Albinism, X-Linked
  https://www.ncbi.nlm.nih.gov/books/NBK1343
Scientific Articles on PubMed

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28Albinism,+Ocular%5BMAJR%5D%29+AND+%28ocular+albinism%5BTIAB%5D%29+AND+english%5BLa%5D+AND+human%5Bmh%5D+AND+%22last+3600+days%22%5Bdp%5D

Catalog of Genes and Diseases from OMIM

- ALBINISM, OCULAR, TYPE I
  http://omim.org/entry/300500

- ALBINISM, OCULAR, WITH LATE-ONSET SENSORINEURAL DEAFNESS
  http://omim.org/entry/300650

- ALBINISM, OCULOCUTANEOUS, TYPE IB
  http://omim.org/entry/606952

Medical Genetics Database from MedGen

- Albinism, ocular, with late-onset sensorineural deafness

- Albinism, ocular, with sensorineural deafness

- Ocular albinism, type I

- Ocular albinism, type II

Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/12868035

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16303920

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/20301517
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11793467

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15090016

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