Gyrate atrophy of the choroid and retina

Gyrate atrophy of the choroid and retina, which is often shortened to gyrate atrophy, is an inherited disorder characterized by progressive vision loss. People with this disorder have an ongoing loss of cells (atrophy) in the retina, which is the specialized light-sensitive tissue that lines the back of the eye, and in a nearby tissue layer called the choroid. During childhood, they begin experiencing nearsightedness (myopia), difficulty seeing in low light (night blindness), and loss of side (peripheral) vision. Over time, their field of vision continues to narrow, resulting in tunnel vision. Many people with gyrate atrophy also develop clouding of the lens of the eyes (cataracts). These progressive vision changes lead to blindness by about the age of 50.

Most people with gyrate atrophy have no symptoms other than vision loss, but some have additional features of the disorder. Occasionally, newborns with gyrate atrophy develop excess ammonia in the blood (hyperammonemia), which may lead to poor feeding, vomiting, seizures, or coma. Neonatal hyperammonemia associated with gyrate atrophy generally responds quickly to treatment and does not recur after the newborn period.

Gyrate atrophy usually does not affect intelligence; however, abnormalities may be observed in brain imaging or other neurological testing. In some cases, mild to moderate intellectual disability is associated with gyrate atrophy.

Gyrate atrophy may also cause disturbances in the nerves connecting the brain and spinal cord to muscles and sensory cells (peripheral nervous system). In some people with the disorder these abnormalities lead to numbness, tingling, or pain in the hands or feet, while in others they are detectable only by electrical testing of the nerve impulses.

In some people with gyrate atrophy, a particular type of muscle fibers (type II fibers) break down over time. While this muscle abnormality usually causes no symptoms, it may result in mild weakness.

Frequency

More than 150 individuals with gyrate atrophy have been identified; approximately one third are from Finland.

Genetic Changes

Mutations in the OAT gene cause gyrate atrophy. The OAT gene provides instructions for making the enzyme ornithine aminotransferase. This enzyme is active in the energy-producing centers of cells (mitochondria), where it helps break down a molecule called ornithine. Ornithine is involved in the urea cycle, which processes excess nitrogen (in the form of ammonia) that is generated when protein is broken down by the body.
In addition to its role in the urea cycle, ornithine participates in several reactions that help ensure the proper balance of protein building blocks (amino acids) in the body. This balance is important because a specific sequence of amino acids is required to build each of the many different proteins needed for the body's functions. The ornithine aminotransferase enzyme helps convert ornithine into another molecule called pyrroline-5-carboxylate (P5C). P5C can be converted into the amino acids glutamate and proline.

OAT gene mutations that cause gyrate atrophy result in a reduced amount of functional ornithine aminotransferase enzyme. A shortage of this enzyme impedes the conversion of ornithine into P5C. As a result, excess ornithine accumulates in the blood (hyperornithinemia), and less P5C than normal is produced. It is not clear how these changes result in the specific signs and symptoms of gyrate atrophy. Researchers have suggested that a deficiency of P5C may interfere with the function of the retina. It has also been proposed that excess ornithine may suppress the production of a molecule called creatine. Creatine is needed for many tissues in the body to store and use energy properly. It is involved in providing energy for muscle contraction, and it is also important in nervous system functioning.

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

- gyrate atrophy
- HOGA
- hyperornithinemia with gyrate atrophy of choroid and retina
- OAT deficiency
- OKT deficiency
- ornithine aminotransferase deficiency
- ornithine-delta-aminotransferase deficiency
- ornithine keto acid aminotransferase deficiency
- Ornithinemia with gyrate atrophy

Diagnosis & Management

Genetic Testing

- Genetic Testing Registry: Ornithine aminotransferase deficiency
Other Diagnosis and Management Resources

- Baby's First Test  
  http://www.babysfirsttest.org/newborn-screening/conditions/hyperornithine-with-gyrate-deficiency

General Information from MedlinePlus

- Diagnostic Tests  
  https://medlineplus.gov/diagnostictests.html
- Drug Therapy  
  https://medlineplus.gov/drugtherapy.html
- Genetic Counseling  
  https://medlineplus.gov/geneticcounseling.html
- Palliative Care  
  https://medlineplus.gov/palliativecare.html
- Surgery and Rehabilitation  
  https://medlineplus.gov/surgeryandrehabilitation.html

Additional Information & Resources

MedlinePlus

- Health Topic: Eye Diseases  
  https://medlineplus.gov/eyediseases.html

Genetic and Rare Diseases Information Center

- Gyrate atrophy of choroid and retina  

Additional NIH Resources

- National Eye Institute: Low Vision  
  https://nei.nih.gov/LowVision/

Educational Resources

- Disease InfoSearch: Ornithine aminotransferase deficiency  
  http://www.diseaseinfosearch.org/Ornithine+aminotransferase+deficiency/5411
- MalaCards: gyrate atrophy of choroid and retina  
  http://www.malacards.org/card/gyrate_atrophy_of_choroid_and_retina
- Orphanet: Gyrate atrophy of choroid and retina  
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=414
Patient Support and Advocacy Resources

- American Foundation for the Blind
  https://www.afb.org/default.aspx
- Children Living with Inherited Metabolic Diseases
  http://www.climb.org.uk/
- Foundation Fighting Blindness
  http://www.blindness.org/glossary/gyrate-atrophy-ga
- Retina International
  http://www.retina-international.org/patients/rare-conditions/gyrate-atrophy

ClinicalTrials.gov

- ClinicalTrials.gov
  https://clinicaltrials.gov/ct2/results?cond=%22Gyrate+Atrophy%22+OR+%22gyrate+atrophy+of+the+choroid+and+retina%22

Scientific Articles on PubMed

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28gyrate+atrophy+of+the+choroid+and+retina%5BTIAB%5D%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+3600+days%22%5Bdp%5D

OMIM

- GYRATE ATROPHY OF CHOROID AND RETINA
  http://omim.org/entry/258870

MedGen

- Gyrate atrophy of choroid and retina

Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16151897

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17088329
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2077941/
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/10354199

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15249361

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11831916

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/10617919

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/3339136
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC329615/

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/12221166

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11297489

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15159649

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11521006

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/10604138

Reprinted from Genetics Home Reference:

Reviewed: August 2009
Published: July 17, 2018