Neuropathy, ataxia, and retinitis pigmentosa

Neuropathy, ataxia, and retinitis pigmentosa (NARP) is a condition that causes a variety of signs and symptoms that mainly affect the nervous system. The condition typically begins in childhood or early adulthood, and the signs and symptoms usually worsen over time. Most people with NARP experience numbness, tingling, or pain in the arms and legs (sensory neuropathy); muscle weakness; and problems with balance and coordination (ataxia). Many affected individuals also have vision loss caused by changes in the light-sensitive tissue that lines the back of the eye (the retina). In some cases, the vision loss results from a condition called retinitis pigmentosa. This eye disease causes the light-sensing cells of the retina gradually to deteriorate.

Learning disabilities and developmental delays are often seen in children with NARP, and older individuals with this condition may experience a loss of intellectual function (dementia). Other features of NARP include seizures, hearing loss, and abnormalities of the electrical signals that control the heartbeat (cardiac conduction defects). These signs and symptoms vary among affected individuals.

Frequency

The prevalence of NARP is unknown. This disorder is probably less common than a similar but more severe condition, Leigh syndrome, which affects about 1 in 40,000 people.

Causes

NARP results from mutations in the MT-ATP6 gene. This gene is contained in mitochondrial DNA, also known as mtDNA. Mitochondria are structures within cells that convert the energy from food into a form that cells can use. Although most DNA is packaged in chromosomes within the nucleus, mitochondria also have a small amount of their own DNA.

The MT-ATP6 gene provides instructions for making a protein that is essential for normal mitochondrial function. Through a series of chemical reactions, mitochondria use oxygen and simple sugars to create adenosine triphosphate (ATP), the cell's main energy source. The MT-ATP6 protein forms one part (subunit) of an enzyme called ATP synthase, which is responsible for the last step in ATP production. Mutations in the MT-ATP6 gene alter the structure or function of ATP synthase, reducing the ability of mitochondria to make ATP. It remains unclear how this disruption in mitochondrial energy production leads to muscle weakness, vision loss, and the other specific features of NARP.
Inheritance Pattern
This condition is inherited in a mitochondrial pattern, which is also known as maternal inheritance. This pattern of inheritance applies to genes contained in mtDNA. Because egg cells, but not sperm cells, contribute mitochondria to the developing embryo, children can inherit disorders resulting from mtDNA mutations only from their mother. These disorders can appear in every generation of a family and can affect both males and females, but fathers do not pass traits associated with changes in mtDNA to their children.

Most of the body's cells contain thousands of mitochondria, each with one or more copies of mtDNA. The severity of some mitochondrial disorders is associated with the percentage of mitochondria in each cell that has a particular genetic change. Most individuals with NARP have a specific *MT-ATP6* mutation in 70 percent to 90 percent of their mitochondria. When this mutation is present in a higher percentage of a person's mitochondria—more than 90 percent to 95 percent—it usually causes a more severe condition known as maternally inherited Leigh syndrome. Because these two conditions result from the same genetic changes and can occur in different members of a single family, and because some individuals with *MT-ATP6* gene mutations have related signs and symptoms that do not follow the specific patterns of these conditions, researchers believe that the conditions may be part of a spectrum of overlapping features rather than two distinct syndromes.

Other Names for This Condition
- NARP
- NARP syndrome
- neurogenic muscle weakness, ataxia, and retinitis pigmentosa
- neuropathy, ataxia, and retinitis pigmentosa

Diagnosis & Management

Genetic Testing Information
- What is genetic testing? /primer/testing/genetictesting

Research Studies from ClinicalTrials.gov
- ClinicalTrials.gov https://clinicaltrials.gov/ct2/results?cond=%22neuropathy%2C+ataxia%2C+and+retinitis+pigmentosa%22
Other Diagnosis and Management Resources

- GeneReview: Mitochondrial Disorders Overview
  https://www.ncbi.nlm.nih.gov/books/NBK1224
- GeneReview: Mitochondrial DNA-Associated Leigh Syndrome and NARP
  https://www.ncbi.nlm.nih.gov/books/NBK1173
- MedlinePlus Encyclopedia: Retinitis pigmentosa
  https://medlineplus.gov/ency/article/001029.htm

Additional Information & Resources

Health Information from MedlinePlus

- Encyclopedia: Retinitis pigmentosa
  https://medlineplus.gov/ency/article/001029.htm
- Health Topic: Cerebellar Disorders
  https://medlineplus.gov/cerebellardisorders.html
- Health Topic: Mitochondrial Diseases
  https://medlineplus.gov/mitochondrialdiseases.html
- Health Topic: Retinal Disorders
  https://medlineplus.gov/retinaldisorders.html

Genetic and Rare Diseases Information Center

- Neuropathy ataxia retinitis pigmentosa syndrome

Additional NIH Resources

- National Institute of Neurological Disorders and Stroke
  https://www.ninds.nih.gov/Disorders/All-Disorders/Mitochondrial-myopathy-Information-Page

Educational Resources

- Kennedy Krieger Institute
  https://www.kennedykrieger.org/patient-care/conditions/mitochondrial-disorders
- MalaCards: mitochondrial dna-associated leigh syndrome and narp
  https://www.malacards.org/card/mitochondrial_dna_associated_leigh_syndrome_and_narp
- Neuromuscular Disease Center, Washington University
  https://neuromuscular.wustl.edu/mitosyn.html#narp
- Orphanet: NARP syndrome
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=644
Patient Support and Advocacy Resources

- Children's Mitochondrial Disease Network (UK)
  http://www.cmdn.org.uk/
- Foundation Fighting Blindness: Retinitis Pigmentosa
  https://www.fightingblindness.org/diseases/retinitis-pigmentosa
- MitoAction
  https://www.mitoaction.org/
- Muscular Dystrophy Association: Facts About Mitochondrial Myopathies
- National Organization for Rare Disorders (NORD)
- Resource list from the University of Kansas Medical Center
  http://www.kumc.edu/gec/support/mitochon.html
- United Mitochondrial Disease Foundation
  https://www.umdf.org/

Clinical Information from GeneReviews

- Mitochondrial Disorders Overview
  https://www.ncbi.nlm.nih.gov/books/NBK1224
- Mitochondrial DNA-Associated Leigh Syndrome and NARP
  https://www.ncbi.nlm.nih.gov/books/NBK1173

Scientific Articles on PubMed

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28Mitochondrial+Diseases%5BMAJR%5D%29+AND+%28%28%5Bneuropathy%28+ataxia%29%2C+retinitis%29+AND+english%2B%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+3600+days%22+AND+%5B%5D+AND+%22last+3600+days%5B%22+AND+%5B%5D+AND+%22last+3600+days%5B%22+AND+%5B%5D

Catalog of Genes and Diseases from OMIM

- NEUROPATHY, ATAXIA, AND RETINITIS PIGMENTOSA
  http://omim.org/entry/551500

Medical Genetics Database from MedGen

- Neuropathy ataxia retinitis pigmentosa syndrome
Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/10396197 
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC1722923/

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/31187502

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16525806

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/20301352

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/9221962 
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2169628/

Reprinted from Genetics Home Reference: 

Reviewed: August 2019 
Published: December 10, 2019

Lister Hill National Center for Biomedical Communications
U.S. National Library of Medicine
National Institutes of Health
Department of Health & Human Services