Hypokalemic periodic paralysis

Hypokalemic periodic paralysis is a condition that causes episodes of extreme muscle weakness typically beginning in childhood or adolescence. Most often, these episodes involve a temporary inability to move muscles in the arms and legs. Attacks cause severe weakness or paralysis that usually lasts from hours to days. Some people may have episodes almost every day, while others experience them weekly, monthly, or only rarely. Attacks can occur without warning or can be triggered by factors such as rest after exercise, a viral illness, or certain medications. Often, a large, carbohydrate-rich meal or vigorous exercise in the evening can trigger an attack upon waking the following morning. Although affected individuals usually regain their muscle strength between attacks, repeated episodes can lead to persistent muscle weakness later in life.

People with hypokalemic periodic paralysis have reduced levels of potassium in their blood (hypokalemia) during episodes of muscle weakness. Researchers are investigating how low potassium levels may be related to the muscle abnormalities in this condition.

Frequency

Although its exact prevalence is unknown, hypokalemic periodic paralysis is estimated to affect 1 in 100,000 people. Men tend to experience symptoms of this condition more often than women.

Causes

Mutations in the \textit{CACNA1S} or \textit{SCN4A} gene can cause hypokalemic periodic paralysis. These genes provide instructions for making proteins that play an essential role in muscles used for movement (skeletal muscles). For the body to move normally, these muscles must tense (contract) and relax in a coordinated way. Muscle contractions are triggered by the flow of certain positively charged atoms (ions) into muscle cells. The \textit{CACNA1S} and \textit{SCN4A} proteins form channels that control the flow of these ions. The channel formed by the \textit{CACNA1S} protein transports calcium ions into cells, while the channel formed by the \textit{SCN4A} protein transports sodium ions.

Mutations in the \textit{CACNA1S} or \textit{SCN4A} gene alter the usual structure and function of calcium or sodium channels. The altered channels cannot properly regulate the flow of ions into muscle cells, which reduces the ability of skeletal muscles to contract. Because muscle contraction is needed for movement, a disruption in normal ion transport leads to episodes of severe muscle weakness or paralysis.
A small percentage of people with the characteristic features of hypokalemic periodic paralysis do not have identified mutations in the *CACNA1S* or *SCN4A* gene. In these cases, the cause of the condition is unknown.

**Inheritance Pattern**

This condition is inherited in an autosomal dominant pattern, which means one copy of the altered gene in each cell is sufficient to cause the disorder.

**Other Names for This Condition**

- Familial Hypokalemic Periodic Paralysis
- HOKPP
- HypoKPP
- HypoPP
- Primary Hypokalemic Periodic Paralysis
- Westphall disease

**Diagnosis & Management**

**Formal Treatment/Management Guidelines**

- Cochrane Reviews: Treatment for Periodic Paralysis

**Genetic Testing Information**

- What is genetic testing?
  /primer/testing/genetictesting
- Genetic Testing Registry: Hypokalemic periodic paralysis
- Genetic Testing Registry: Hypokalemic periodic paralysis 1
- Genetic Testing Registry: Hypokalemic periodic paralysis, type 2

**Research Studies from ClinicalTrials.gov**

- ClinicalTrials.gov
  https://clinicaltrials.gov/ct2/results?cond=%22hypokalemic+periodic+paralysis%22
Other Diagnosis and Management Resources

• GeneReview: Hypokalemic Periodic Paralysis
  https://www.ncbi.nlm.nih.gov/books/NBK1338

• MedlinePlus Encyclopedia: Hypokalemic periodic paralysis
  https://medlineplus.gov/ency/article/000312.htm

Additional Information & Resources

Health Information from MedlinePlus

• Encyclopedia: Hypokalemic periodic paralysis
  https://medlineplus.gov/ency/article/000312.htm

• Health Topic: Muscle Disorders
  https://medlineplus.gov/muscledisorders.html

• Health Topic: Paralysis
  https://medlineplus.gov/paralysis.html

Genetic and Rare Diseases Information Center

• Hypokalemic periodic paralysis

Additional NIH Resources

• National Institute of Neurological Disorders and Stroke
  https://www.ninds.nih.gov/Disorders/All-Disorders/Familial-Periodic-Paralyses-Information-Page

Educational Resources

• MalaCards: hypokalemic periodic paralysis, type 1
  https://www.malacards.org/card/hypokalemic_periodic_paralysis_type_1_2

• MalaCards: hypokalemic periodic paralysis, type 2
  https://www.malacards.org/card/hypokalemic_periodic_paralysis_type_2_2

• Merck Manual Consumer Version

• Neuromuscular Disease Center, Washington University
  https://neuromuscular.wustl.edu/mtime/mepisodic.html#hopp

• Orphanet: Hypokalemic periodic paralysis
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=681
Patient Support and Advocacy Resources

• Muscular Dystrophy Association
  https://www.mda.org/disease/endocrine-myopathies

• Periodic Paralysis Association
  https://www.periodicparalysis.org/site/

• Periodic Paralysis International
  http://hkpp.org/

Clinical Information from GeneReviews

• Hypokalemic Periodic Paralysis
  https://www.ncbi.nlm.nih.gov/books/NBK1338

Scientific Articles on PubMed

• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28Hypokalemic+Periodic+Paralysis%5BMAJR%5D%29+AND+%28hypokalemic+periodic+paralysis%5BTIAB%5D%29+AND+english%5BLa%5D+AND+human%5Bmh%5D+AND+%22last+1800+days%22%5Bdp%5D

Catalog of Genes and Diseases from OMIM

• HYPOKALEMIC PERIODIC PARALYSIS, TYPE 1
  http://omim.org/entry/170400

Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/12117476

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/9132138

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17139526

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11898585

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/18426576
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2374768/


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