Spastic paraplegia type 15

Spastic paraplegia type 15 is part of a group of genetic disorders known as hereditary spastic paraplegias. These disorders are characterized by progressive muscle stiffness (spasticity) and the development of paralysis of the lower limbs (paraplegia). Spastic paraplegia type 15 is classified as a complex hereditary spastic paraplegia because it involves all four limbs as well as additional features, including abnormalities of the brain. In addition to the muscles and brain, spastic paraplegia type 15 affects the peripheral nervous system, which consists of nerves connecting the brain and spinal cord to muscles and sensory cells that detect sensations such as touch, pain, heat, and sound.

Spastic paraplegia type 15 usually becomes apparent in childhood or adolescence with the development of weak muscle tone (hypotonia), difficulty walking, or intellectual disability. In almost all affected individuals, the tissue connecting the left and right halves of the brain (corpus callosum) is abnormally thin and becomes thinner over time. Additionally, there is often a loss (atrophy) of nerve cells in several parts of the brain, including the cerebral cortex, which controls thinking and emotions, and the cerebellum, which coordinates movement.

People with this form of spastic paraplegia can have numbness, tingling, or pain in the arms and legs (sensory neuropathy); impairment of the nerves used for muscle movement (motor neuropathy); exaggerated reflexes (hyperreflexia) of the lower limbs; muscle wasting (amyotrophy); or reduced bladder control. Rarely, spastic paraplegia type 15 is associated with a group of movement abnormalities called parkinsonism, which includes tremors, rigidity, and unusually slow movement (bradykinesia). People with spastic paraplegia type 15 may have an eye condition called pigmentary maculopathy that often impairs vision. This condition results from the breakdown (degeneration) of tissue at the back of the eye called the macula, which is responsible for sharp central vision.

Most people with spastic paraplegia type 15 experience a decline in intellectual ability and an increase in muscle weakness and nerve abnormalities over time. As the condition progresses, many people require walking aids or wheelchair assistance in adulthood.

Frequency

Spastic paraplegia type 15 is a rare condition, although its exact prevalence is unknown.
Causes

Mutations in the ZFYVE26 gene cause spastic paraplegia type 15. This gene provides instructions for making a protein called spastizin. This protein is important in a process called autophagy, in which worn-out cell parts and unneeded proteins are recycled within cells. Specifically, spastizin is involved in the formation and maturation of sacs called autophagosomes (or autophagic vacuoles) that transport unneeded materials to be broken down. Spastizin also plays a role in the process by which dividing cells separate from one another (cytokinesis).

Many ZFYVE26 gene mutations that cause spastic paraplegia type 15 result in a shortened spastizin protein that is quickly broken down. As a result, functional autophagosomes are not produced, autophagy cannot occur, and recycling of materials within cells is decreased. An inability to break down unneeded materials, and the subsequent accumulation of these materials in cells, leads to cell dysfunction and often cell death. The loss of cells in the brain and other parts of the body is responsible for many of the features of spastic paraplegia type 15.

It is unclear whether a lack of spastizin protein interferes with normal cytokinesis and whether impaired cell division contributes to the signs and symptoms of spastic paraplegia type 15.

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

- autosomal recessive spastic paraplegia 15
- Kjellin syndrome
- spastic paraplegia and retinal degeneration
- SPG15

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=%22spastic+paraplegia+type+15%22+OR+%22SPG15%22+OR+%22Spastic+Paraplegia%22+OR+Hereditary%22

Other Diagnosis and Management Resources

- GeneReview: Hereditary Spastic Paraplegia Overview
  https://www.ncbi.nlm.nih.gov/books/NBK1509

- Spastic Paraplegia Foundation, Inc: Treatments and Therapies
  https://sp-foundation.org/understanding-pls-hsp/treatments.html

Additional Information & Resources

Health Information from MedlinePlus

- Health Topic: Neuromuscular Disorders
  https://medlineplus.gov/neuromusculardisorders.html

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

- Health Topic: Peripheral Nerve Disorders
  https://medlineplus.gov/peripheralnervedisorders.html

Genetic and Rare Diseases Information Center

- Spastic paraplegia 15

Additional NIH Resources

- National Institute of Neurological Disorders and Stroke: Hereditary Spastic Paraplegia Information Page
  https://www.ninds.nih.gov/Disorders/All-Disorders/Hereditary-spastic-paraplegia-Information-Page

Educational Resources


- Orphanet: Hereditary spastic paraplegia
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=685

- Washington University, St. Louis: Neuromuscular Disease Center
  https://neuromuscular.wustl.edu/spinal/fsp.html#spg15
Patient Support and Advocacy Resources

- National Organization for Rare Disorders (NORD): Hereditary Spastic Paraplegia
  https://rarediseases.org/rare-diseases/hereditary-spastic-paraplegia/
- RareConnect
  https://www.rareconnect.org/en/community/hereditary-spastic-paraplegia

Clinical Information from GeneReviews

- Hereditary Spastic Paraplegia Overview
  https://www.ncbi.nlm.nih.gov/books/NBK1509

Scientific Articles on PubMed

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28spg15%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

- SPASTIC PARAPLEGIA 15, AUTOSOMAL RECESSIVE
  http://omim.org/entry/270700

Sources for This Summary

- Boukhris A, Stevanin G, Feki I, Denis E, Elleuch N, Miladi MI, Truchetto J, Denora P, Belal S,
  Mhiri C, Brice A. Hereditary spastic paraplegia with mental impairment and thin corpus callosum

- Denora PS, Muglia M, Casali C, Truchetto J, Silvestri G, Messina D, Boukrhis A, Magariello A,
  Modoni A, Masiuollo M, Malandrini A, Morelli M, de Leva MF, Villanova M, Giugni E, Citrigno L,
  paraplegia with thinning of the corpus callosum and white matter abnormalities: further mutations
  and relative frequency in ZFYVE26/SPG15 in the Italian population. J Neurol Sci. 2009 Feb 15;

- Goizet C, Boukhris A, Maltete D, Guyant-Maréchal L, Truchetto J, Mundwiller E, Hanein S,
  G. SPG15 is the second most common cause of hereditary spastic paraplegia, including Kjellin syndrome.

  A, Stevanin G. Identification of the SPG15 gene, encoding spastizin, as a frequent cause of
  2008 Apr;82(4):992-1002. doi: 10.1016/j.ajhg.2008.03.004. Citation on PubMed:
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2427184/
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/24366652

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/24284334

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/24030950
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