Genetics
Home Reference

Your Guide to Understanding Genetic Conditions

GJB1 gene

gap junction protein beta 1

Normal Function

The GJB1 gene provides instructions for making a protein called connexin-32 (also known as gap junction beta 1). This protein is a member of the gap junction connexin family, which plays a role in cell communication by forming channels, or gap junctions, between cells. Gap junctions speed the transport of nutrients, charged particles (ions), and small molecules that carry communication signals between cells.

The connexin-32 protein is made in several tissues, including those of the liver, pancreas, kidney, and nervous system. In the nervous system, this protein is located in the cell membrane of specialized cells called Schwann cells and oligodendrocytes. Schwann cells are found in the peripheral nervous system, which consists of nerves connecting the brain and spinal cord (central nervous system) to muscles and sensory cells that detect sensations such as touch, pain, heat, and sound. Oligodendrocytes are located in the central nervous system.

The Schwann cells and oligodendrocytes surround nerves and are involved in the production and long-term maintenance of a fatty substance called myelin. Myelin forms a protective coating around certain nerve cells and ensures the smooth and rapid transmission of nerve impulses.

The connexin-32 protein forms channels through the myelin sheath, allowing efficient transport and communication between the outer myelin layers and the interior of the Schwann cell or oligodendrocyte.

Health Conditions Related to Genetic Changes

Charcot-Marie-Tooth disease

Researchers have identified approximately 300 GJB1 gene mutations in people with type X Charcot-Marie-Tooth disease, a disorder characterized by muscle weakness and sensory problems, especially in the hands and feet. A few of these mutations also cause hearing loss in individuals with this type of Charcot-Marie-Tooth disease. Most GJB1 gene mutations change single protein building blocks (amino acids) in the connexin-32 protein. Some GJB1 gene mutations result in a protein of abnormal size.

It is unclear how GJB1 gene mutations lead to the characteristic features of Charcot-Marie-Tooth disease, including a loss of myelin (demyelination) and the slowed transmission of nerve impulses in the peripheral nervous system. The altered protein may be degraded quickly or trapped inside the cell, preventing it from reaching the cell membrane to form gap junctions. In some cases, an altered protein reaches
the cell membrane but does not form properly functioning gap junctions. The loss of functional gap junctions probably impairs the normal activities of Schwann cells, such as myelin production. Malfunctioning gap junctions could also disrupt communication between Schwann cells and the underlying nerve cell, disturbing the transmission of nerve impulses.

In addition to the peripheral nervous system problems associated with this disorder, evidence of demyelination in the central nervous system has been reported in some people with Charcot-Marie-Tooth disease caused by \textit{GJB1} gene mutations. These central nervous system abnormalities did not generally cause any symptoms, but were identified by electrical testing of nerve impulses or imaging studies. Research suggests that another connexin protein whose function overlaps with that of connexin-32 helps compensate for the mutated connexin-32 protein in the oligodendrocytes of the central nervous system.

**Chromosomal Location**

Cytogenetic Location: Xq13.1, which is the long (q) arm of the X chromosome at position 13.1

Molecular Location: base pairs 71,215,212 to 71,225,215 on the X chromosome (Homo sapiens Annotation Release 109, GRCh38.p12) (NCBI)

Credit: Genome Decoration Page/NCBI

**Other Names for This Gene**

- CMTX
- CMTX1
- connexin 32
- CX32
- CXB1\_HUMAN
- gap junction protein, beta 1, 32kDa
- gap junction protein, beta 1, 32kDa (connexin 32, Charcot-Marie-Tooth neuropathy, X-linked)
Additional Information & Resources

Educational Resources

• Biochemistry (fifth edition, 2002): Gap Junctions Allow Ions and Small Molecules to Flow between Communicating Cells
  https://www.ncbi.nlm.nih.gov/books/NBK22492/

Clinical Information from GeneReviews

• Charcot-Marie-Tooth Neuropathy X Type 1
  https://www.ncbi.nlm.nih.gov/books/NBK1374

Scientific Articles on PubMed

• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28GJB1%5BTIAB%5D%29+OR+%28connexin+32%5BTIAB%5D%29+OR+%28cx+32%5BTIAB%5D%29%29+AND+%28Genes%5BMH%5D+OR+%28Genetic+Phenomena%5BMH%5D%29+OR+%28%5BMH%5D%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+1080+days%22+Bdp%5D

Catalog of Genes and Diseases from OMIM

• GAP JUNCTION PROTEIN, BETA-1
  http://omim.org/entry/304040

Research Resources

• Atlas of Genetics and Cytogenetics in Oncology and Haematology
  http://atlasgeneticsoncology.org/Genes/GC_GJB1.html

• ClinVar
  https://www.ncbi.nlm.nih.gov/clinvar?term=GJB1%5Bgene%5D

• Connexin-Deafness Homepage
  http://davinci.crg.es/deafness/

• HGNC Gene Family: Gap junction proteins
  https://www.genenames.org/cgi-bin/genefamilies/set/314

• HGNC Gene Symbol Report
  https://www.genenames.org/cgi-bin/gene_symbol_report?q=data/hgnc_data.php&hgnc_id=4283

• Inherited Peripheral Neuropathies Mutation Database
  http://www.molgen.ua.ac.be/CMTMutations/Mutations/Mutations.cfm?Context=3

• Monarch Initiative
  https://monarchinitiative.org/gene/NCBIGene:2705
Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/18949782


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