FA2H gene
fatty acid 2-hydroxylase

Normal Function

The *FA2H* gene provides instructions for making an enzyme called fatty acid 2-hydroxylase. This enzyme modifies fatty acids, which are building blocks used to make fats (lipids). Specifically, fatty acid 2-hydroxylase adds a single oxygen atom to a hydrogen atom at a particular point on a fatty acid to create a 2-hydroxylated fatty acid. Certain 2-hydroxylated fatty acids are important in forming normal myelin; myelin is the protective covering that insulates nerves and ensures the rapid transmission of nerve impulses. The part of the brain and spinal cord that contains myelin is called white matter.

Health Conditions Related to Genetic Changes

Fatty acid hydroxylase-associated neurodegeneration

At least nine mutations in the *FA2H* gene have been identified in people with fatty acid hydroxylase-associated neurodegeneration (FAHN). FAHN is a progressive disorder of the nervous system characterized by problems with movement and vision that begin during childhood or adolescence and worsen with age. Brain scans of affected individuals show abnormal accumulation of iron in the brain, especially in a region that is involved in movement.

The *FA2H* gene mutations that cause FAHN reduce or eliminate the function of the fatty acid 2-hydroxylase enzyme. Reduction of this enzyme's function may result in abnormal myelin that is prone to deterioration (demyelination), leading to a loss of white matter (leukodystrophy). Leukodystrophy is likely involved in the development of the movement problems and other neurological abnormalities that occur in FAHN. Iron accumulation in the brain is probably also involved, although it is unclear how *FA2H* gene mutations lead to the buildup of iron.

People with *FA2H* gene mutations and some of the movement problems seen in FAHN were once classified as having a separate disorder called spastic paraplegia 35. People with mutations in this gene resulting in intellectual decline and optic nerve atrophy were said to have a disorder called *FA2H*-related leukodystrophy. However, these conditions are now generally considered to be forms of FAHN.
Chromosomal Location

Cytogenetic Location: 16q23.1, which is the long (q) arm of chromosome 16 at position 23.1

Molecular Location: base pairs 74,712,955 to 74,774,831 on chromosome 16 (Homo sapiens Updated Annotation Release 109.20190905, GRCh38.p13) (NCBI)

Credit: Genome Decoration Page/NCBI

Other Names for This Gene

- FA2H_HUMAN
- FAAH
- FAH1
- fatty acid alpha-hydroxylase
- fatty acid hydroxylase domain containing 1
- FAXDC1
- FLJ25287
- SCS7
- spastic paraplegia 35 (autosomal recessive)
- SPG35

Additional Information & Resources

Educational Resources

- Basic Neurochemistry: Molecular, Cellular and Medical Aspects (sixth edition, 1999): Characteristic Composition of Myelin
  https://www.ncbi.nlm.nih.gov/books/NBK28221/

Clinical Information from GeneReviews

- Fatty Acid Hydroxylase-Associated Neurodegeneration
  https://www.ncbi.nlm.nih.gov/books/NBK56080
**Scientific Articles on PubMed**

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28FA2H%5BTIAB%5D+OR+%28fatty+acid+2-hydroxylase%5BTIAB%5D%29%29+OR+%28%28FAH1%5BTIAB%5D%29+OR+%28SCS7%5BTIAB%5D%29+OR+%28SPG35%5BTIAB%5D%29+OR+%28fatty+acid+alpha-hydroxylase%5BTIAB%5D%29%29+AND+%28%28Genes%5BMH%5D%29+OR+%28Genetic+Phenomena%5BMH%5D%29%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+720+days%22%5Bdp%5D

**Catalog of Genes and Diseases from OMIM**

- FATTY ACID 2-HYDROXYLASE
  http://omim.org/entry/611026

**Research Resources**

- Atlas of Genetics and Cytogenetics in Oncology and Haematology
  http://atlasgeneticsoncology.org/Genes/GC_FA2H.html

- ClinVar

- HGNC Gene Symbol Report

- Monarch Initiative
  https://monarchinitiative.org/gene/NCBIGene:79152

- NCBI Gene

- UniProt
  https://www.uniprot.org/uniprot/Q7L5A8

**Sources for This Summary**

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15337768

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  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/21286947

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