TSC1 gene
TSC complex subunit 1

Normal Function

The *TSC1* gene provides instructions for producing a protein called hamartin, whose function is not fully understood. Within cells, hamartin interacts with a protein called tuberin, which is produced from the *TSC2* gene. These two proteins help control cell growth and size. Proteins that normally prevent cells from growing and dividing too fast or in an uncontrolled way are known as tumor suppressors. Hamartin and tuberin carry out their tumor suppressor function by interacting with and regulating a wide variety of other proteins.

Health Conditions Related to Genetic Changes

**Lymphangioleiomyomatosis**

Mutations in the *TSC1* gene can cause a disorder called lymphangioleiomyomatosis (LAM), although mutations in the *TSC2* gene appear to be responsible for most cases of this disorder. This destructive lung disease is caused by the abnormal overgrowth of smooth muscle-like tissue in the lungs. It occurs almost exclusively in women, causing coughing, shortness of breath, chest pain, and lung collapse.

LAM can occur alone (isolated or sporadic LAM) or in combination with a condition called tuberous sclerosis complex (described below). Researchers suggest that sporadic LAM can be caused by a random mutation in the *TSC1* gene that occurs very early in development. As a result, some of the body's cells have a normal version of the gene, while others have the mutated version. This situation is called mosaicism. When a mutation occurs in the other copy of the *TSC1* gene in certain cells during a woman's lifetime (a somatic mutation), she may develop LAM.

**Tuberous sclerosis complex**

More than 400 mutations in the *TSC1* gene have been identified in individuals with tuberous sclerosis complex, a condition characterized by developmental problems and the growth of noncancerous tumors in many parts of the body. Most of these mutations involve either small deletions or insertions of DNA in the *TSC1* gene. Some mutations create a premature stop signal in the instructions for making hamartin.

People with *TSC1*-related tuberous sclerosis complex are born with one mutated copy of the *TSC1* gene in each cell. This mutation prevents the cell from making functional hamartin from that copy of the gene. However, enough hamartin is usually produced from the other, normal copy of the *TSC1* gene to regulate cell growth.
effectively. For some types of tumors to develop, a second mutation involving the other copy of the gene must occur in certain cells during a person’s lifetime.

When both copies of the \( TSC1 \) gene are mutated in a particular cell, that cell cannot produce any functional hamartin. The loss of this protein allows the cell to grow and divide in an uncontrolled way to form a tumor. A shortage of hamartin also interferes with the normal development of certain cells. In people with \( TSC1 \)-related tuberous sclerosis complex, a second \( TSC1 \) gene mutation typically occurs in multiple cells over an affected person’s lifetime. The loss of hamartin in different types of cells disrupts normal development and leads to the growth of tumors in many different organs and tissues.

**Other disorders**

Inherited mutations in the \( TSC1 \) gene can cause a disorder known as focal cortical dysplasia of Taylor balloon cell type. This disorder involves malformations of the cerebrum, the large, frontal part of the brain that is responsible for thinking and learning. Focal cortical dysplasia causes severe recurrent seizures (epilepsy) in affected individuals.

**Chromosomal Location**

Cytogenetic Location: 9q34.13, which is the long (q) arm of chromosome 9 at position 34.13

Molecular Location: base pairs 132,891,349 to 132,945,269 on chromosome 9 (Homo sapiens Updated Annotation Release 109.20200228, GRCh38.p13) (NCBI)

Credit: Genome Decoration Page/NCBI

**Other Names for This Gene**

- hamartin
- KIAA0243
- TSC1_HUMAN
- tuberous sclerosis 1
Additional Information & Resources

Educational Resources

• Cancer Medicine (sixth edition, 2003): Tuberous Sclerosis
  https://www.ncbi.nlm.nih.gov/books/NBK13658/#A19617

Clinical Information from GeneReviews

• Tuberous Sclerosis Complex
  https://www.ncbi.nlm.nih.gov/books/NBK1220

Scientific Articles on PubMed

• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28TSC1%5BTIAB%5D%29+OR+%28hamartin%5BTIAB%5D%29%29+AND+%28%28Genes%5BMH%5D%29+OR+%28Genetic+Phenomena%5BMH%5D%29%29+AND+english%5BLan%5D+AND+human%5Bmh%5D+AND+%22last+720+days%22%5Bdp%5D

Catalog of Genes and Diseases from OMIM

• FOCAL CORTICAL DYSPLASIA, TYPE II
  http://omim.org/entry/607341

• TSC1 GENE
  http://omim.org/entry/605284

Research Resources

• Atlas of Genetics and Cytogenetics in Oncology and Haematology
  http://atlasgeneticsoncology.org/Genes/TSC1ID183.html

• ClinVar
  https://www.ncbi.nlm.nih.gov/clinvar?term=TSC1%5Bgene%5D

• HGNC Gene Symbol Report
  https://www.genenames.org/data/gene-symbol-report/#!/hgnc_id/HGNC:12362

• Monarch Initiative
  https://monarchinitiative.org/gene/NCBIGene:7248

• NCBI Gene
  https://www.ncbi.nlm.nih.gov/gene/7248

• UniProt
  https://www.uniprot.org/uniprot/Q92574
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