XK gene
X-linked Kx blood group

Normal Function
The XK gene provides instructions for producing a protein that is found in various tissues of the body, particularly the brain, muscle, and heart. This protein is also present on the surface of red blood cells and carries a molecule known as the Kx blood group antigen. Blood group antigens are found on the surface of red blood cells and determine a number of blood types. When blood cells with unfamiliar antigens enter the bloodstream, the body’s immune system recognizes the cells as foreign and may trigger an immune reaction that destroys the foreign blood cells.

The function of the XK protein is unclear; researchers believe that it might play a role in transporting substances into and out of cells. On red blood cells, the XK protein attaches to another blood group protein, the Kell protein. The function of this blood group complex is unknown.

Health Conditions Related to Genetic Changes
McLeod neuroacanthocytosis syndrome

Approximately 30 mutations in the XK gene have been found to cause McLeod neuroacanthocytosis syndrome. Most of these mutations lead to the production of an abnormally short, nonfunctional XK protein or cause no XK protein to be produced at all. Missing or abnormal XK protein also affects another blood group antigen, the Kell protein, causing it to be less prevalent on the surface of red blood cells. How mutations in the XK gene cause the misshapen red blood cells and movement problems characteristic of McLeod neuroacanthocytosis syndrome is unknown.
Chromosomal Location

Cytogenetic Location: Xp21.1, which is the short (p) arm of the X chromosome at position 21.1

Molecular Location: base pairs 37,685,686 to 37,732,130 on the X chromosome (Homo sapiens Annotation Release 109, GRCh38.p12) (NCBI)

Credit: Genome Decoration Page/NCBI

Other Names for This Gene

- Kell blood group precursor (McLeod phenotype)
- KX
- Kx antigen
- membrane transport protein XK
- X-linked Kx blood group (McLeod syndrome)
- X1k
- XK-related protein 1
- XK, Kell blood group complex subunit (McLeod syndrome)
- XK_HUMAN
- XKR1

Additional Information & Resources

Educational Resources

Clinical Information from GeneReviews
Scientific Articles on PubMed

• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28XK%5BTIAB%5D%29+OR+%28XK+antigen%5BTIAB%5D%29+AND+%28%28Genes%5BMH%5D%29%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+2520+days%22%5Bdp%5D

Catalog of Genes and Diseases from OMIM

• KELL BLOOD GROUP PROTEIN, MCLEOD SYNDROME-ASSOCIATED
  http://omim.org/entry/314850

Research Resources

• HGNC Gene Symbol Report

• Monarch Initiative
  https://monarchinitiative.org/gene/NCBIGene:7504

• NCBI Gene

• UniProt
  https://www.uniprot.org/uniprot/P51811

Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17379193

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11761473

• OMIM: KELL BLOOD GROUP PROTEIN, MCLEOD SYNDROME-ASSOCIATED
  http://omim.org/entry/314850

• Lee S. The value of DNA analysis for antigens of the Kell and Kx blood group systems. Transfusion. 2007 Jul;47(1 Suppl):32S-9S. Review.
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17593284

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11961232

Reprinted from Genetics Home Reference: