NLRP3 gene
NLR family pyrin domain containing 3

Normal Function

The *NLRP3* gene (also known as *CIAS1*) provides instructions for making a protein called cryopyrin. Cryopyrin is a member of a family of proteins called nucleotide-binding domain and leucine-rich repeat containing (NLR) proteins, which are found in the fluid inside cells (cytoplasm). Cryopyrin is found mainly in white blood cells and in cartilage-forming cells (chondrocytes).

NLR proteins are involved in the immune system, helping to start and regulate the immune system's response to injury, toxins, or invasion by microorganisms. These proteins recognize specific molecules, become activated, and respond by helping to engage components of the immune system. Cryopyrin recognizes bacterial particles; chemicals such as asbestos, silica, and uric acid crystals; and compounds released by injured cells.

Once activated, groups of cryopyrin molecules assemble themselves along with other proteins into structures called inflammasomes, which are involved in the process of inflammation. Inflammation occurs when the immune system sends signaling molecules as well as white blood cells to a site of injury or disease to fight microbial invaders and facilitate tissue repair.

Health Conditions Related to Genetic Changes

**Familial cold autoinflammatory syndrome**

Several mutations in the *NLRP3* gene have been identified in people with familial cold autoinflammatory syndrome. These mutations are in a region of the gene known as exon 3. Researchers believe that the mutations cause cryopyrin to be hyperactive, leading to episodes of fever and inflammation that are usually triggered by exposure to cold.

**Muckle-Wells syndrome**

At least 10 mutations in exon 3 of the *NLRP3* gene have been identified in people with Muckle-Wells syndrome. These mutations are believed to cause hyperactive cryopyrin, resulting in episodes of fever and inflammation, as well as the hearing loss and kidney problems that occur in Muckle-Wells syndrome.

**Neonatal onset multisystem inflammatory disease**

About 30 mutations in the *NLRP3* gene have been identified in people with neonatal onset multisystem inflammatory disease (NOMID). Almost all of these mutations
are found in exon 3. The mutations likely cause cryopyrin to be hyperactive, leading to an inappropriate inflammatory response that results in episodes of fever and widespread inflammatory damage to the body’s cells and tissues. It is unclear why some mutations in exon 3 cause the severe symptoms of NOMID, some cause the less serious familial cold autoinflammatory syndrome, and others cause Muckle-Wells syndrome, which is intermediate in severity.

**Chromosomal Location**

Cytogenetic Location: 1q44, which is the long (q) arm of chromosome 1 at position 44

Molecular Location: base pairs 247,416,156 to 247,449,108 on chromosome 1 (Homo sapiens Annotation Release 109, GRCh38.p12) (NCBI)

Credit: Genome Decoration Page/NCBI

**Other Names for This Gene**

- AGTAVPRL
- AII
- AII/AVP
- AII/AVP receptor-like
- angiotensin/vasopressin receptor AII/AVP-like
- AVP
- C1orf7
- CIAS1
- CLR1.1
- cryopyrin
- FCAS
- FCU
- FLJ95925
- MWS
• NACHT domain-, leucine-rich repeat-, and PYD-containing protein 3
• NACHT, LRR and PYD containing protein 3
• NALP3
• NALP3_HUMAN
• NLR family, pyrin domain containing 3
• nucleotide-binding oligomerization domain, leucine rich repeat and pyrin domain containing 3
• PYPAF1
• PYRIN-containing APAF1-like protein 1

Additional Information & Resources

Educational Resources
• Immunobiology (fifth edition, 2001): Innate Immunity
  https://www.ncbi.nlm.nih.gov/books/NBK10769/

Scientific Articles on PubMed
• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28NLRP3%5BTIAB%5D%29+OR+%28%28AII/AVP%5BTIAB%5D%29+OR+%28CIAS1%5BTIAB%5D%29+OR+%28cryopyrin%5BTIAB%5D%29+OR+%28FCAS%5BTIAB%5D%29+OR+%28FCU%5BTIAB%5D%29+OR+%28NALP3%5BTIAB%5D%29+OR+%28PYPAF1%5BTIAB%5D%29+OR+%28PYRIN-containing+APAF1-like+protein+1%5BTIAB%5D+OR+%28Genetic+Phenomena%5BMH%5D%29+AND+%28Genes%5BMH%5D%29+AND+english+AND+human+AND+%22last+360+days%22

Catalog of Genes and Diseases from OMIM
• NLR FAMILY, PYRIN DOMAIN-CONTAINING 3
  http://omim.org/entry/606416

Research Resources
• Atlas of Genetics and Cytogenetics in Oncology and Haematology
  http://atlasgeneticsoncology.org/Genes/GC_NLRP3.html
• ClinVar
• HGNC Gene Symbol Report
• Monarch Initiative
  https://monarchinitiative.org/gene/NCBIGene:114548
Sources for This Summary

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17393462
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4321998/

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/18172447

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/18347298

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16407890

- OMIM: NLR FAMILY, PYRIN DOMAIN-CONTAINING 3
  http://omim.org/entry/606416

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16546100

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15300846

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15720239

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