Partington syndrome

Partington syndrome is a neurological disorder that causes intellectual disability along with a condition called focal dystonia that particularly affects movement of the hands. Partington syndrome usually occurs in males; when it occurs in females, the signs and symptoms are often less severe.

The intellectual disability associated with Partington syndrome usually ranges from mild to moderate. Some affected individuals have characteristics of autism spectrum disorders that affect communication and social interaction. Recurrent seizures (epilepsy) may also occur in Partington syndrome.

Focal dystonia of the hands is a feature that distinguishes Partington syndrome from other intellectual disability syndromes. Dystonias are a group of movement problems characterized by involuntary, sustained muscle contractions; tremors; and other uncontrolled movements. The term “focal” refers to a type of dystonia that affects a single part of the body, in this case the hands. In Partington syndrome, focal dystonia of the hands, which is called the Partington sign, begins in early childhood and gradually gets worse. This condition typically causes difficulty with grasping movements or using a pen or pencil.

People with Partington syndrome may also have dystonia affecting other parts of the body; dystonia affecting the muscles in the face and those involved in speech may cause impaired speech (dysarthria). People with this disorder may also have an awkward way of walking (gait). Signs and symptoms can vary widely, even within the same family.

Frequency

The prevalence of Partington syndrome is unknown. At least 20 cases have been described in the medical literature.

Causes

Partington syndrome is caused by mutations in the ARX gene. This gene provides instructions for producing a protein that regulates the activity of other genes. Within the developing brain, the ARX protein is involved with movement (migration) and communication of nerve cells (neurons). In particular, this protein regulates genes that play a role in the migration of specialized neurons (interneurons) to their proper location. Interneurons relay signals between other neurons.

The normal ARX protein contains four regions where a protein building block (amino acid) called alanine is repeated multiple times. These stretches of alanines are known as polyalanine tracts. The most common mutation that causes Partington syndrome,
a duplication of genetic material written as c.428_451dup, adds extra alanines to the second polyalanine tract in the ARX protein. This type of mutation is called a polyalanine repeat expansion. The expansion likely impairs ARX protein function and may disrupt normal interneuron migration in the developing brain, leading to the intellectual disability and dystonia characteristic of Partington syndrome.

Inheritance Pattern

This condition is inherited in an X-linked recessive pattern. The gene associated with this condition is located on the X chromosome, which is one of the two sex chromosomes. In males (who have only one X chromosome), one altered copy of the gene in each cell is sufficient to cause the condition. In females (who have two X chromosomes), a mutation would have to occur in both copies of the gene to cause the disorder. Because it is unlikely that females will have two altered copies of this gene, males are affected by X-linked recessive disorders much more frequently than females. Females with one altered copy of the gene may have some signs and symptoms related to the condition. A characteristic of X-linked inheritance is that fathers cannot pass X-linked traits to their sons.

Rarely, the genetic change that causes Partington syndrome is not inherited but occurs at some point during embryonic development. As cells continue to grow and divide, some of these cells will have the genetic change, and others will not (a situation known as mosaicism). The mosaic nature of these genetic changes lead to relatively mild features of Partington syndrome.

Other Names for This Condition

- MRX36
- Partington-Mulley syndrome
- Partington X-linked mental retardation syndrome
- PRTS
- X-linked intellectual deficit-dystonia-dysarthria
- X-linked mental retardation with dystonic movements, ataxia, and seizures

Diagnosis & Management

Genetic Testing Information

- What is genetic testing? /primer/testing/genetictesting
Other Diagnosis and Management Resources

- American Academy of Child and Adolescent Psychiatry: Services in School for Children with Special Needs

- American Academy of Pediatrics: What is a Developmental/Behavioral Pediatrician?

- Centers for Disease Control and Prevention: Developmental Screening Fact Sheet

Additional Information & Resources

Health Information from MedlinePlus

- Encyclopedia: Intellectual Disability
  https://medlineplus.gov/ency/article/001523.htm

- Health Topic: Developmental Disabilities
  https://medlineplus.gov/developmentaldisabilities.html

- Health Topic: Dystonia
  https://medlineplus.gov/dystonia.html

Genetic and Rare Diseases Information Center

- Partington syndrome

Additional NIH Resources

- Eunice Kennedy Shriver National Institute of Child Health and Human Development: Intellectual and Developmental Disabilities
  https://www.nichd.nih.gov/health/topics/idds/conditioninfo/default

- National Institute of Neurological Disorders and Stroke: Dystonias Fact Sheet
  https://www.ninds.nih.gov/Disorders/All-Disorders/Dystonias-Information-Page

Educational Resources

- Center for Parent Information and Resources: Intellectual Disability
  https://www.parentcenterhub.org/intellectual/

- Centers for Disease Control and Prevention: Intellectual Disability Fact Sheet
• MalaCards: partington x-linked mental retardation syndrome
  https://www.malacards.org/card/partington_x_linked_mental_retardation_syndrome

• Orphanet: Partington syndrome
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=94083

Patient Support and Advocacy Resources
• American Association on Intellectual and Developmental Disabilities (AAIDD)
  https://www.aaidd.org/

Scientific Articles on PubMed
• PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28partington+syndrome%5BTIAB%5D%29+AND+english%5Bla%5D+AND+human%5Bmh%5D

Catalog of Genes and Diseases from OMIM
• PARTINGTON X-LINKED MENTAL RETARDATION SYNDROME
  http://omim.org/entry/309510

Medical Genetics Database from MedGen
• Partington X-linked mental retardation syndrome

Sources for This Summary
  JF, Leheup B, Delobel B, Demeer B, Poirier K, Biancalana V, Pinoit JM, Julia S, Chelly J, Devys D,
  Mandel JL. ARX polyalanine expansions are highly implicated in familial cases of mental retardation

• Frints SG, Froyen G, Marynen P, Willekens D, Legius E, Fryns JP. Re-evaluation of MRX36 family
  after discovery of an ARX gene mutation reveals mild neurological features of Partington syndrome.

• Grønskov K, Diness B, Stahlhut M, Zilmer M, Tümer Z, Bisgaard AM, Brøndum-Nielsen K.
  Mosaicism for c.431_454dup in ARX causes a mild Partington syndrome phenotype. Eur J Med

• Partington MW, Turner G, Boyle J, Gécz J. Three new families with X-linked mental retardation

• Poirier K, Lacombe D, Gilbert-Dussardier B, Raynaud M, Desportes V, de Brouwer AP, Moraine C,
  Fryns JP, Ropers HH, Beldjord C, Chelly J, Bienvenu T. Screening of ARX in mental retardation
  families: Consequences for the strategy of molecular diagnosis. Neurogenetics. 2006 Mar;7(1):
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/14631200

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/20506206

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/15707237

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/12376946

Reprinted from Genetics Home Reference:

Reviewed: August 2017
Published: January 21, 2020

Lister Hill National Center for Biomedical Communications
U.S. National Library of Medicine
National Institutes of Health
Department of Health & Human Services