Meier-Gorlin syndrome

Meier-Gorlin syndrome is a condition primarily characterized by short stature. It is considered a form of primordial dwarfism because the growth problems begin before birth (intrauterine growth retardation). After birth, affected individuals continue to grow at a slow rate. Other characteristic features of this condition are underdeveloped or missing kneecaps (patellae), small ears, and, often, an abnormally small head (microcephaly). Despite a small head size, most people with Meier-Gorlin syndrome have normal intellect.

Some people with Meier-Gorlin syndrome have other skeletal abnormalities, such as unusually narrow long bones in the arms and legs, a deformity of the knee joint that allows the knee to bend backwards (genu recurvatum), and slowed mineralization of bones (delayed bone age).

Most people with Meier-Gorlin syndrome have distinctive facial features. In addition to being abnormally small, the ears may be low-set or rotated backward. Additional features can include a small mouth (microstomia), an underdeveloped lower jaw (micrognathia), full lips, and a narrow nose with a high nasal bridge.

Abnormalities in sexual development may also occur in Meier-Gorlin syndrome. In some males with this condition, the testes are small or undescended (cryptorchidism). Affected females may have unusually small external genital folds (hypoplasia of the labia majora) and small breasts. Both males and females with this condition can have sparse or absent underarm (axillary) hair.

Additional features of Meier-Gorlin syndrome can include difficulty feeding and a lung condition known as pulmonary emphysema or other breathing problems.

Frequency

Meier-Gorlin syndrome is a rare condition; however, its prevalence is unknown.

Causes

Meier-Gorlin syndrome can be caused by mutations in one of several genes. Each of these genes, \textit{ORC1}, \textit{ORC4}, \textit{ORC6}, \textit{CDT1}, and \textit{CDC6}, provides instructions for making one of a group of proteins known as the pre-replication complex. This complex regulates initiation of the copying (replication) of DNA before cells divide. Specifically, the pre-replication complex attaches (binds) to certain regions of DNA known as origins of replication, allowing copying of the DNA to begin at that location. This tightly controlled process, called replication licensing, helps ensure that DNA replication occurs only once per cell division and is required for cells to divide.
Mutations in any one of these genes impair formation of the pre-replication complex and disrupt replication licensing; however, it is not clear how a reduction in replication licensing leads to Meier-Gorlin syndrome. Researchers speculate that such a reduction delays the cell division process, which impairs growth of the bones and other tissues during development. Some research suggests that some of the pre-replication complex proteins have additional functions, impairment of which may contribute to features of Meier-Gorlin syndrome, such as delayed development of the kneecaps and ears.

Inheritance Pattern

This condition is inherited in an autosomal recessive pattern, which means both copies of the gene in each cell have mutations. The parents of an individual with an autosomal recessive condition each carry one copy of the mutated gene, but they typically do not show signs and symptoms of the condition.

Other Names for This Condition

• ear, patella, short stature syndrome
• microtia, absent patellae, micrognathia syndrome

Diagnosis & Management

Genetic Testing Information

• What is genetic testing? /primer/testing/genetictesting

Additional Information & Resources

Health Information from MedlinePlus

• Encyclopedia: Microcephaly https://medlineplus.gov/ency/article/003272.htm
• Health Topic: Dwarfism https://medlineplus.gov/dwarfism.html
Genetic and Rare Diseases Information Center

- Meier-Gorlin syndrome

Additional NIH Resources

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

Educational Resources

- Boston Children's Hospital: Growth Problems
  http://www.childrenshospital.org/conditions-and-treatments/conditions/g/growth-problems
- MalaCards: meier-gorlin syndrome 1
  https://www.malacards.org/card/meier_gorlin_syndrome_1
- MalaCards: meier-gorlin syndrome 2
  https://www.malacards.org/card/meier_gorlin_syndrome_2
- MalaCards: meier-gorlin syndrome 3
  https://www.malacards.org/card/meier_gorlin_syndrome_3
- MalaCards: meier-gorlin syndrome 4
  https://www.malacards.org/card/meier_gorlin_syndrome_4
- MalaCards: meier-gorlin syndrome 5
  https://www.malacards.org/card/meier_gorlin_syndrome_5
- Orphanet: Ear-patella-short stature syndrome
  https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=2554
- Walking With Giants Foundation: Meier-Gorlin Syndrome
  https://www.walkingwithgiants.org/meier-gorlin-syndrome/

Patient Support and Advocacy Resources

- Little People of America
  https://www.lpaonline.org/
- Little People UK
  https://littlepeopleuk.org/
- National Organization for Rare Disorders (NORD): Ear, Patella, Short Stature Syndrome
- Potentials Foundation
  http://www.potentialsfoundation.org/
• The MAGIC Foundation
  https://www.magicfoundation.org/
• Walking With Giants Foundation
  https://www.walkingwithgiants.org/

**Scientific Articles on PubMed**

• PubMed
%5D%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last
+3600+days%22%5Bdp%5D

**Catalog of Genes and Diseases from OMIM**

• MEIER-GORLIN SYNDROME 1
  http://omim.org/entry/224690
• MEIER-GORLIN SYNDROME 2
  http://omim.org/entry/613800
• MEIER-GORLIN SYNDROME 3
  http://omim.org/entry/613803
• MEIER-GORLIN SYNDROME 4
  http://omim.org/entry/613804
• MEIER-GORLIN SYNDROME 5
  http://omim.org/entry/613805

**Sources for This Summary**

  M, Brown PA, van Bokhoven H, Dean J, Edrees AY, Feingold M, Fryer A, Hoefsloot LH, Kau N,
  Knoers NV, Mackenzie J, Opitz JM, Sarda P, Ross A, Temple IK, Toutain A, Wise CA, Wright M,
  Jackson AP. Mutations in the pre-replication complex cause Meier-Gorlin syndrome. Nat Genet.
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/21358632
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3068194/

  Sanna N, Bober M, Johnson D, Wise C, Jackson AP, O'Driscoll M, Jeggo PA. Mutations in ORC1,
  encoding the largest subunit of the origin recognition complex, cause microcephalic primordial
  ng.776. 
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/21358633

  Langlois S, Superneau DW, Parkash S, Ludman M, Skidmore DL, Samuels ME. Mutations in origin
  360-4. doi: 10.1038/ng.777. 
  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/21358631
Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/22333897
Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3355263/

Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/23023959


Reviewed: February 2014
Published: March 31, 2020

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