Cranio- and metaphyseal dysplasia

Cranio- and metaphyseal dysplasia is a rare condition characterized by progressive thickening of bones in the skull (cranium) and abnormalities at the ends of long bones in the limbs (metaphyseal dysplasia). Except in the most severe cases, the lifespan of people with cranio- and metaphyseal dysplasia is normal.

Bone overgrowth in the head causes many of the signs and symptoms of cranio- and metaphyseal dysplasia. Affected individuals typically have distinctive facial features such as a wide nasal bridge, a prominent forehead, wide-set eyes (hypertelorism), and a prominent jaw. Excessive new bone formation (hyperostosis) in the jaw can delay teething (dentition) or result in absent (non-erupting) teeth. Infants with this condition may have breathing or feeding problems caused by narrow nasal passages. In severe cases, abnormal bone growth can compress the nerves that emerge from the brain and extend to various areas of the head and neck (cranial nerves). Compression of the cranial nerves can lead to paralyzed facial muscles (facial nerve palsy), blindness, or deafness.

The x-rays of individuals with cranio- and metaphyseal dysplasia show unusually shaped long bones, particularly the large bones in the legs. The ends of these bones (metaphyses) are wider and appear less dense in people with this condition.

There are two types of cranio- and metaphyseal dysplasia, which are distinguished by their pattern of inheritance. They are known as the autosomal dominant and autosomal recessive types. Autosomal recessive cranio- and metaphyseal dysplasia is typically more severe than the autosomal dominant form.

Frequency

Cranio- and metaphyseal dysplasia is a very rare disorder; its incidence is unknown.

Genetic Changes

Mutations in the \textit{ANKH} gene cause autosomal dominant cranio- and metaphyseal dysplasia. The \textit{ANKH} gene provides instructions for making a protein that is present in bone and transports a molecule called pyrophosphate out of cells. Pyrophosphate helps regulate bone formation by preventing mineralization, the process by which minerals such as calcium and phosphorus are deposited in developing bones. The ANKH protein may have other, unknown functions.

Mutations in the \textit{ANKH} gene that cause autosomal dominant cranio- and metaphyseal dysplasia may decrease the ANKH protein’s ability to transport pyrophosphate out of cells. Reduced levels of pyrophosphate can increase bone mineralization, contributing to the bone overgrowth seen in cranio- and metaphyseal dysplasia. Why long bones are...
shaped differently and only the skull bones become thicker in people with this condition remains unclear.

The genetic cause of autosomal recessive craniometaphyseal dysplasia is unknown. Researchers believe that mutations in an unidentified gene on chromosome 6 may be responsible for the autosomal recessive form of this condition.

Inheritance Pattern

Craniometaphyseal dysplasia can have different inheritance patterns. In most cases this condition is inherited in an autosomal dominant pattern, which means one altered copy of the ANKH gene in each cell is sufficient to cause the disorder. Individuals with autosomal dominant craniometaphyseal dysplasia typically have one parent who also has the condition. Less often, cases result from new mutations in the gene and occur in people with no history of the disorder in their family.

Rarely, craniometaphyseal dysplasia is suspected to have autosomal recessive inheritance when unaffected parents have more than one child with the condition. Autosomal recessive disorders are caused by mutations in both copies of a gene in each cell. The parents of an individual with an autosomal recessive condition each carry one copy of a mutated gene, but they typically do not show signs and symptoms of the disorder.

Other Names for This Condition

- Autosomal dominant craniometaphyseal dysplasia
- Autosomal recessive craniometaphyseal dysplasia
- CMD
- CMDD
- CMDJ
- CMDR
- Craniometaphyseal dysplasia, Jackson type

Diagnosis & Management

Genetic Testing

Other Diagnosis and Management Resources

- GeneReview: Craniometaphyseal Dysplasia, Autosomal Dominant
  https://www.ncbi.nlm.nih.gov/books/NBK1461

- MedlinePlus Encyclopedia: Facial Paralysis
  https://medlineplus.gov/ency/article/003028.htm

General Information from MedlinePlus

- Diagnostic Tests
  https://medlineplus.gov/diagnostictests.html

- Drug Therapy
  https://medlineplus.gov/drugtherapy.html

- Genetic Counseling
  https://medlineplus.gov/geneticcounseling.html

- Palliative Care
  https://medlineplus.gov/palliativecare.html

- Surgery and Rehabilitation
  https://medlineplus.gov/surgeryandrehabilitation.html

Additional Information & Resources

MedlinePlus

- Encyclopedia: Facial Paralysis
  https://medlineplus.gov/ency/article/003028.htm

- Health Topic: Bone Diseases
  https://medlineplus.gov/bonediseases.html

- Health Topic: Craniofacial Abnormalities
  https://medlineplus.gov/craniofacialabnormalities.html

- Health Topic: Hearing Disorders and Deafness
  https://medlineplus.gov/hearingdisordersanddeafness.html

- Health Topic: Vision Impairment and Blindness
  https://medlineplus.gov/visionimpairmentandblindness.html

Genetic and Rare Diseases Information Center

- Craniometaphyseal dysplasia, autosomal dominant

- Craniometaphyseal dysplasia, autosomal recessive type
  https://rarediseases.info.nih.gov/diseases/1582/craniometaphyseal-dysplasia-autosomal-recessive-type
Educational Resources

- Disease InfoSearch: Craniometaphyseal Dysplasia, Autosomal Dominant
  http://www.diseaseinfosearch.org/Craniometaphyseal+Dysplasia%2C+Autosomal+Dominant/1975
- Disease InfoSearch: Craniometaphyseal Dysplasia, Autosomal Recessive Type
  http://www.diseaseinfosearch.org/Craniometaphyseal+Dysplasia%2C+Autosomal+Recessive+Type/1976
- MalaCards: craniometaphyseal dysplasia, autosomal dominant
  http://www.malacards.org/card/craniometaphyseal_dysplasia_autosomal_dominant
- MalaCards: craniometaphyseal dysplasia, autosomal recessive
  http://www.malacards.org/card/craniometaphyseal_dysplasia_autosomal_recessive
- Orphanet: Craniometaphyseal dysplasia
  http://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=1522

Patient Support and Advocacy Resources

- AboutFace International
  http://www.aboutface.ca/
- AmeriFace
  http://www.ameriface.org/
- Children's Craniofacial Association
  https://ccakids.org/
- International Skeletal Dysplasia Registry, UCLA
  http://ortho.ucla.edu/isdr
- National Organization for Rare Disorders (NORD)
  https://rarediseases.org/rare-diseases/craniometaphyseal-dysplasia/

GeneReviews

- Craniometaphyseal Dysplasia, Autosomal Dominant
  https://www.ncbi.nlm.nih.gov/books/NBK1461

ClinicalTrials.gov

- ClinicalTrials.gov
  https://clinicaltrials.gov/ct2/results?cond=%22craniometaphyseal+dysplasia%22

Scientific Articles on PubMed

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28%28craniometaphyseal+dysplasia%5BTIAB%5D%29+OR+%28autosomal+dominant+craniometaphyseal+dysplasia%5BTIAB%5D%29%29+AND+english%5BLa%5D+AND+human%5Bmh%5D+AND+%22last+3600+days%22%5Bdp%5D
CRANIOMETAPOPHYSEAL DYSPLASIA, AUTOSOMAL DOMINANT
http://omim.org/entry/123000

Sources for This Summary


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