Klinefelter syndrome

Klinefelter syndrome is a chromosomal condition that affects male physical and cognitive development. Its signs and symptoms vary among affected individuals.

Affected individuals typically have small testes that do not produce as much testosterone as usual. Testosterone is the hormone that directs male sexual development before birth and during puberty. A shortage of testosterone can lead to delayed or incomplete puberty, breast enlargement (gynecomastia), reduced facial and body hair, and an inability to have biological children (infertility). Some affected individuals also have genital differences including undescended testes (cryptorchidism), the opening of the urethra on the underside of the penis (hypospadias), or an unusually small penis (micropenis).

Older children and adults with Klinefelter syndrome tend to be taller than their peers. Compared with unaffected men, adults with Klinefelter syndrome have an increased risk of developing breast cancer and a chronic inflammatory disease called systemic lupus erythematosus. Their chance of developing these disorders is similar to that of women in the general population.

Children with Klinefelter syndrome may have learning disabilities and delayed speech and language development. They tend to be quiet, sensitive, and unassertive, but personality characteristics vary among affected individuals.

Frequency

Klinefelter syndrome affects 1 in 500 to 1,000 newborn males. Most variants of Klinefelter syndrome are much rarer, occurring in 1 in 50,000 or fewer newborns.

Researchers suspect that Klinefelter syndrome is underdiagnosed because the condition may not be identified in people with mild signs and symptoms. Additionally, the features of the condition vary and overlap significantly with those of other conditions.

Genetic Changes

Klinefelter syndrome is a condition related to the X and Y chromosomes (the sex chromosomes). People typically have two sex chromosomes in each cell: females have two X chromosomes (46,XX), and males have one X and one Y chromosome (46,XY). Most often, Klinefelter syndrome results from the presence of one extra copy of the X chromosome in each cell (47,XXY). Extra copies of genes on the X chromosome interfere with male sexual development, often preventing the testes from functioning normally and reducing the levels of testosterone. Most people with an extra
X chromosome have the features described above, although some have few or no associated signs and symptoms.

Some people with features of Klinefelter syndrome have more than one extra sex chromosome in each cell (for example, 48,XXXY or 49,XXXXY). These conditions, which are often called variants of Klinefelter syndrome, tend to cause more severe signs and symptoms than classic Klinefelter syndrome. In addition to affecting male sexual development, variants of Klinefelter syndrome are associated with intellectual disability, distinctive facial features, skeletal abnormalities, poor coordination, and severe problems with speech. As the number of extra sex chromosomes increases, so does the risk of these health problems.

Some people with features of Klinefelter syndrome have the extra X chromosome in only some of their cells; in these individuals, the condition is described as mosaic Klinefelter syndrome (46,XY/47,XXY). Individuals with mosaic Klinefelter syndrome may have milder signs and symptoms, depending on how many cells have an additional X chromosome.

Inheritance Pattern

Klinefelter syndrome and its variants are not inherited; these chromosomal changes usually occur as random events during the formation of reproductive cells (eggs and sperm) in a parent. An error in cell division called nondisjunction results in a reproductive cell with an abnormal number of chromosomes. For example, an egg or sperm cell may gain one or more extra copies of the X chromosome as a result of nondisjunction. If one of these atypical reproductive cells contributes to the genetic makeup of a child, the child will have one or more extra X chromosomes in each of the body's cells.

Mosaic 46,XY/47,XXY is also not inherited. It occurs as a random event during cell division early in fetal development. As a result, some of the body's cells have one X chromosome and one Y chromosome (46,XY), and other cells have an extra copy of the X chromosome (47,XXY).

Other Names for This Condition

- Klinefelter's syndrome
- XXY syndrome
- XXY trisomy

Diagnosis & Management

Genetic Testing

- Genetic Testing Registry: Klinefelter's syndrome, XXY
Other Diagnosis and Management Resources

- MedlinePlus Encyclopedia: Klinefelter Syndrome
  https://medlineplus.gov/ency/article/000382.htm
- MedlinePlus Encyclopedia: Testicular Failure
  https://medlineplus.gov/ency/article/000395.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: Klinefelter Syndrome
  https://medlineplus.gov/ency/article/000382.htm
- Encyclopedia: Testicular Failure
  https://medlineplus.gov/ency/article/000395.htm
- Health Topic: Klinefelter's Syndrome
  https://medlineplus.gov/klinefelterssyndrome.html

Genetic and Rare Diseases Information Center

- 47, XXY
  https://rarediseases.info.nih.gov/diseases/11920/47-xyy
- 48,XXYY syndrome
- Klinefelter syndrome
**Additional NIH Resources**

- Eunice Kennedy Shriver National Institute of Child Health and Human Development  
  https://www.nichd.nih.gov/health/topics/klinefelter

- National Human Genome Research Institute  
  https://www.genome.gov/19519068/

**Educational Resources**

- Centre for Genetics Education  

- Genetic Science Learning Center, University of Utah  
  http://learn.genetics.utah.edu/content/disorders/extraormissing/

- March of Dimes: Chromosomal Conditions  
  https://www.marchofdimes.org/baby/chromosomal-conditions.aspx

- Merck Manual Consumer Version  

- My46 Trait Profile  
  https://www.my46.org/trait-document?trait=Klinefelter%20syndrome&type=profile

- Orphanet: NON RARE IN EUROPE: Klinefelter syndrome  
  http://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=484

**Patient Support and Advocacy Resources**

- Accord Alliance  
  http://www.accordalliance.org/

- Association for X and Y Chromosome Variations: Tell Me About 47,XXY  
  https://genetic.org/variations/about-47xxy/

- National Organization for Rare Disorders (NORD)  
  https://rarediseases.org/rare-diseases/klinefelter-syndrome/

- Resource list from the University of Kansas Medical Center  
  http://www.kumc.edu/gec/support/chromoso.html#xxy

**ClinicalTrials.gov**

- ClinicalTrials.gov  
  https://clinicaltrials.gov/ct2/results?cond=%22klinefelter+syndrome%22
Scientific Articles on PubMed

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28Klinefelter+Syndrome%5BMAJR%29+%5D%29+AND+%28Klinefelter+syndrome%5BTIAB%5D%29+AND+english%5Bla%5D+AND+human%5Bmh%5D+AND+%22last+720+days%22%5Bdp%5D

MedGen

- Klinefelter Syndrome, Variants
- Klinefelter syndrome, XXY

Sources for This Summary


  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/21342258
  Free article on PubMed Central: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3314712/

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/11771918

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/17497714

  Citation on PubMed: https://www.ncbi.nlm.nih.gov/pubmed/16342850

Reprinted from Genetics Home Reference:

Reviewed: January 2013
Published: May 1, 2018

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