Prekallikrein deficiency

Prekallikrein deficiency is a blood condition that usually causes no health problems. In people with this condition, blood tests show a prolonged activated partial thromboplastin time (PTT), a result that is typically associated with bleeding problems; however, bleeding problems generally do not occur in prekallikrein deficiency. The condition is usually discovered when blood tests are done for other reasons.

A few people with prekallikrein deficiency have experienced health problems related to blood clotting such as heart attack, stroke, a clot in the deep veins of the arms or legs (deep vein thrombosis), nosebleeds, or excessive bleeding after surgery. However, these are common problems in the general population, and most affected individuals have other risk factors for developing them, so it is unclear whether their occurrence is related to prekallikrein deficiency.

Frequency

The prevalence of prekallikrein deficiency is unknown. Approximately 80 affected individuals in about 30 families have been described in the medical literature. Because prekallikrein deficiency usually does not cause any symptoms, researchers suspect that most people with the condition are never diagnosed.

Causes

Prekallikrein deficiency is caused by mutations in the KLKB1 gene, which provides instructions for making a protein called prekallikrein. This protein, when converted to an active form called plasma kallikrein in the blood, is involved in the early stages of blood clotting. Plasma kallikrein plays a role in a process called the intrinsic coagulation pathway (also called the contact activation pathway). This pathway turns on (activates) proteins that are needed later in the clotting process. Blood clots protect the body after an injury by sealing off damaged blood vessels and preventing further blood loss.

The KLKB1 gene mutations that cause prekallikrein deficiency reduce or eliminate functional plasma kallikrein, which likely impairs the intrinsic coagulation pathway. Researchers suggest that this lack (deficiency) of functional plasma kallikrein protein does not generally cause any symptoms because another process called the extrinsic coagulation pathway (also known as the tissue factor pathway) can compensate for the impaired intrinsic coagulation pathway.

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

- congenital prekallikrein deficiency
- Fletcher factor deficiency
- Fletcher trait
- PKK deficiency

Diagnosis & Management

Genetic Testing Information

- What is genetic testing? https://primer/testing/genetictesting

Other Diagnosis and Management Resources

- Massachusetts General Hospital Laboratory Handbook: Prekallikrein http://mghlabtest.partners.org/coagbook/CO002600.htm

Additional Information & Resources

Health Information from MedlinePlus

- Health Topic: Bleeding Disorders https://medlineplus.gov/bleedingdisorders.html

Genetic and Rare Diseases Information Center

- Prekallikrein deficiency, congenital https://rarediseases.info.nih.gov/diseases/4477/prekallikrein-deficiency-congenital

Educational Resources

- MalaCards: prekallikrein deficiency, congenital https://www.malacards.org/card/prekallikrein_deficiency_congenital
- Orphanet: Congenital prekallikrein deficiency https://www.orpha.net/consor/cgi-bin/OC_Exp.php?Lng=EN&Expert=749

Patient Support and Advocacy Resources

- National Blood Clot Alliance https://www.stoptheclot.org/
**Scientific Articles on PubMed**

- PubMed
  https://www.ncbi.nlm.nih.gov/pubmed?term=%28prekallikrein+deficiency%5BTIAB%5D%29+AND+english%5Bla%5D+AND+human%5Bmh%5D

**Catalog of Genes and Diseases from OMIM**

- PREKALLIKREIN DEFICIENCY
  http://omim.org/entry/612423

**Sources for This Summary**

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