

Cystic Fibrosis Mechanisms

Cystic fibrosis is a hereditary disease leading to problems with Cl⁻ channels in the body. It is the most common lethal genetic disease in the Caucasian population. Patients develop recurrent pulmonary infections, bronchitis, infertility, pancreatic insufficiency, steatorrhea and malabsorption.



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Autosomal Recessive

Recessive-chocolate

Cystic fibrosis is inherited in an autosomal recessive manner.

CFTR Chromosome 7

CFTR-sifter Chrome 7

This disease occurs due to a defect in the CFTR gene on chromosome 7. Though there are many ways to effect the CFTR gene, commonly, a deletion of Phe508 occurs.

Cl⁻ channel Defect

Chlorine-dispenser Channel Broken

CFTR encodes an ATP-gated Cl⁻ channel. In the lungs and gut, this channel secretes Cl⁻, which leads to an H₂O gradient. In sweat glands, this channel reabsorbs Cl⁻. A defect in CFTR leads to defects in Cl⁻ secretion through these channels.

Decreased Chloride Secretion

Down-arrow Chlorine-dispenser Secreting into GI and Lungs

Defective Cl⁻ channels lead to increased chloride on the skin (not reabsorbed in sweat glands), as well as decreased chloride secretion (and subsequently water) in the gut and lungs.

Increased Na and Water Reabsorption

Up-arrow Salt-shaker and Water-bottle pulled out of Absorbing-sponge

In this disorder, Cl⁻ is not secreted into the lungs and GI tract. Thus, there is increased intracellular Cl⁻, which then causes a compensatory increase in Na⁺ reabsorption. Due to the high concentration of accumulated salt (NaCl) intracellularly, water is then reabsorbed.

Increased Na and Cl in Sweat

Up-arrow Salt-shaker and Chlorine-dispenser at Sweaty-sweatgland

In this disorder, Cl⁻ is not reabsorbed through sweat glands. Increased epithelial Cl⁻ causes a compensatory increase in Na⁺ excretion via epithelial channels.

Dehydration of Mucous Layers

Dried-up Mucous Layers of body

As Cl⁻ is not secreted into the lungs and GI tract and is "trapped" intracellularly, Na⁺ follows and H₂O is reabsorbed. This leads to abnormally thick mucus secreted into the lungs and GI tract.