

OverviewScientific Details

Cystic Fibrosis

Scientific Details

Cystic fibrosis is a rare genetic disorder characterized by impaired lung and digestive function. A person must have two variants in the CFTR gene in order to have this condition.

Print

Cystic fibrosis is caused by variants in the CFTR gene.

CFTR

The CFTR gene contains instructions for making a protein called cystic fibrosis transmembrane conductance regulator. This protein helps control the salt and water balance of certain organs by allowing chloride ions to pass in and out of cells. Certain variants in the CFTR gene disrupt this function, causing the lungs, pancreas, and other organs to produce abnormally thick mucus. This mucus can clog the respiratory tract, leading to signs and symptoms of cystic fibrosis.

Read more at [Genetics Home Reference](#)

Chromosome 7

Gene: CFTR

You have two variants detected by this test.

Variants Detected		View All Tested Markers	
Marker Tested	Genotype*	Additional Information	
DeltaF508 Gene: CFTR Marker: i3000001	(–) Variant copy from one of your parents	CTT Typical copy from your other parent	Biological explanation Typical vs. variant DNA sequence(s) Percent of 23andMe customers with variant References [5] ClinVar
DeltaI507 Gene: CFTR Marker: i4000292	(–) Variant copy from one of your parents	ATC Typical copy from your other parent	Biological explanation Typical vs. variant DNA sequence(s) Percent of 23andMe customers with variant References [5] ClinVar

*This test cannot distinguish which copy you received from which parent. This test also cannot determine whether multiple variants, if detected, were inherited from only one parent or from both parents. This may impact how these variants are passed down.

23andMe always reports genotypes based on the 'positive' strand of the human genome reference sequence (build 37). Other sources sometimes report genotypes using the opposite strand.

Test Details

Indications for Use

The 23andMe PGS Carrier Status Test for Cystic Fibrosis is indicated for the detection of 28 variants in the CFTR gene. This test is intended to be used to determine carrier status for cystic fibrosis in adults, but cannot determine if a person has two copies of a tested variant. The test is most relevant for people of Ashkenazi Jewish, European, and Hispanic/Latino descent.

Special Considerations

- Symptoms of cystic fibrosis may vary depending on the variants involved.
- ACMG recommends carrier testing for cystic fibrosis for people of all ethnicities considering having children. This test includes 22 of 23 variants recommended for testing by ACMG.

Test Performance Summary

Carrier Detection Rate & Relevant Ethnicities

The "carrier detection rate" is an estimate of the percentage of carriers for this condition that would be identified by this test. Carrier detection rate differs by ethnicity and is provided only where sufficient data is available.

Ashkenazi Jewish	94%	[3, 5]
European	89%	[3, 5]
Hispanic/Latino	73%	[3, 5]
African American	65%	[3, 5]
Asian	55%	[3, 5]

Analytical Performance

Accuracy was determined by comparing results from this test with results from sequencing. Greater than 99% of test results were correct. While unlikely, this test may provide false positive or false negative results. For more details on the analytical performance of this test, refer to the package insert.

Warnings and Limitations

- This test does not cover all variants that could cause this condition.*
- This test does not diagnose any health conditions.
- Positive results in individuals whose ethnicities are not commonly associated with this condition may be incorrect. Individuals in this situation should consider genetic counseling and follow-up testing.
- Share results with your healthcare professional for any medical purposes.
- If you are concerned about your results, consult with a healthcare professional.

See the [Package Insert](#) for more details on use and performance of this test.

* Variants not included in this test may be very rare, may not be available on our genotyping platform, or may not pass our testing standards.

References

1. 510(k): Illumina MiSeqDx Cystic Fibrosis Clinical Sequencing Assay

2. American College of Obstetricians and Gynecologists Committee on Genetics. (2011). "ACOG Committee Opinion No. 486: Update on carrier screening for cystic fibrosis." Obstet Gynecol. 117(4):1028-31.

3. Bobadilla JL et al. (2002). "Cystic fibrosis: a worldwide analysis of CFTR mutations--correlation with incidence data and application to screening." Hum Mutat. 19(6):575-606.

4. Ong T et al. (2001). "Cystic Fibrosis and Congenital Absence of the Vas Deferens." [Updated 2017 Feb 2].

5. Watson MS et al. (2004). "Cystic fibrosis population carrier screening: 2004 revision of American College of Medical Genetics mutation panel." Genet Med. 6(5):387-91.

Change Log

Your report may occasionally be updated based on new information. This Change Log describes updates and revisions to this report.

Date	Change
Feb 14, 2019	The variant 2789+5G>A (i4000320) was added to the report. Customers who have this variant will see this variant detected in their result, and see updated content in their report.
Feb 14, 2019	The carrier detection rate was updated for customers who self-report having Ashkenazi Jewish ancestry. The chances of still being a carrier were also updated for customers with no variants detected who self-report having Ashkenazi Jewish ancestry.
Feb 14, 2019	The chances of still being a carrier were updated for customers with no variants detected who self-report having European or African American ancestry.
Feb 14, 2019	Due to improvements in data analysis, some customers who previously received a "Not Determined" result for one or more of the following genetic markers may see a genotype at these markers: i4000291, i4000294, i4000296, i4000300, i4000301, i4000305, i4000307, i4000308, i4000311, i4000314, i4000315, i4000317, i4000318, i4000321, i4000323, i4000324, i4000325. This may also update the overall report result for these customers.
Feb 14, 2019	Cystic Fibrosis report created.

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