

Females do not have a prostate and are not at risk for prostate cancer. You can use this report to learn whether you have a genetic variant that may not affect your health, but could affect the health of your male relatives.

Health > Health Predisposition

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Hereditary Prostate Cancer (HOXB13-Related)

HOXB13-related hereditary prostate cancer is caused by a specific genetic variant in the HOXB13 gene. This variant is associated with an increased risk of developing prostate cancer and is most commonly found in people of European descent, especially people of Northern European descent.

Overview

Scientific Details

Frequently Asked Questions

Jamie, you do not have the G84E variant we tested.

Females do not have a prostate, so they are not at risk for prostate cancer.

However, your male relatives could still have this variant or another variant not tested that could increase their risk for prostate cancer. In addition, most cases of prostate cancer are not caused by inherited variants.

0 variants detected
in the HOXB13 gene

If you have a personal or family history of cancer, you should talk to a healthcare professional.

How To Use This Test

This test does not diagnose cancer or any other health conditions and should not be used to make medical decisions. Results should be confirmed by an independent genetic test prescribed by your own healthcare provider before taking any medical action.

Please talk to a healthcare professional if cancer runs in your family, you think you might have cancer, or you have any concerns about your results.

[Review the Hereditary Prostate Cancer \(HOXB13-Related\) tutorial](#)

[See Frequently Asked Questions](#)

[See Scientific Details for complete Indications for Use statement and full list of Warnings, Precautions, and Limitations](#)

+ Intended Uses

- Tests for the **G84E** variant in the HOXB13 gene. This variant is associated with an increased risk of developing prostate cancer.
- For males: Provides information on whether their genetic result is associated with an increased risk for prostate cancer.
- For females: Provides information on whether they have the tested variant, which means that their male relatives may be at increased risk for HOXB13-related prostate cancer.

- Limitations

- Does **not** test for all possible variants in the HOXB13 gene.
- Does **not** test for variants in other genes linked to hereditary prostate cancer, such as variants in BRCA1, BRCA2, and genes linked to Lynch syndrome.
- Does **not** account for non-genetic factors, like environment and lifestyle, that influence overall prostate cancer risk.
- The interpretation of your genetic result depends on the birth sex you reported in your account settings.

🌐 Ethnicity Considerations

- The variant included in this test is most commonly found in people of **European** descent, especially in people of **Northern European** descent.

You do not have the genetic variant we tested.

However, your male relatives could still have this variant or another variant not tested that could increase their risk for prostate cancer. In addition, most cases of prostate cancer are not caused by inherited variants.



You do not have the G84E variant we tested.

This variant is most commonly found in people of **European** descent, especially in people of **Northern European** descent.

[See Scientific Details](#)

Your male relatives are still at risk for prostate cancer.

This test does not include variants in other genes that can also increase prostate cancer risk in males, including variants in BRCA1, BRCA2, and genes linked to Lynch syndrome. In addition, most cases of prostate cancer are not caused by inherited variants. ⓘ

[See Scientific Details](#)



If you have a personal or family history of cancer, talk to a healthcare professional.

About Hereditary Prostate Cancer (HOXB13-Related)

The G84E variant in the HOXB13 gene is associated with an increased risk for prostate cancer. **Since females do not have a prostate, they are not at risk for prostate cancer.**



Lifetime prostate cancer risk

Males with the HOXB13 G84E variant have a 33–53% chance of developing prostate cancer by age 80, compared to about a 13% lifetime risk for males in the general population. [See Scientific Details to learn more about these risks.](#)



When prostate cancer develops

In general, the chances of developing prostate cancer increase with age. However, males with the HOXB13 G84E variant may develop prostate cancer at an earlier age.



How common is the HOXB13 G84E variant?

Up to 1 in 70 people of European descent has the HOXB13 G84E variant. This variant is expected to be less common in people of other ethnicities.



Prostate cancer screening

National guidelines for prostate cancer screening vary. Some guidelines recommend earlier prostate cancer screening in males with the HOXB13 G84E variant.

A healthcare professional can help your male relatives find an appropriate screening strategy.

Read more at: [MedlinePlus](#) [American Cancer Society](#)

Learn more about HOXB13-related hereditary prostate cancer.

FAQs

See our Frequently Asked Questions for more information.

FAQs



If you have a personal or family history of cancer, consult with a healthcare professional.

Print report



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Overview Scientific Details Frequently Asked Questions

The G84E variant in the HOXB13 gene is associated with an increased risk for prostate cancer.

This report does not include variants in other genes associated with increased prostate cancer risk.

HOXB13

The HOXB13 gene contains instructions for making a protein that helps regulate the expression of other genes. The HOXB13 protein acts as a tumor suppressor, preventing cells from growing and dividing too rapidly. Although the exact mechanism is not well understood, the G84E variant in the HOXB13 gene increases the risk for prostate cancer.

Read more at MedlinePlus*

Chromosome 17



You do not have the G84E variant we tested.

Variants Detected		View All Tested Markers	
Marker Tested	Your Genotype*	Additional Information	
G84E Gene: HOXB13 Marker: rs138213197	C Typical copy from one of your parents	C Typical copy from your other parent	<ul style="list-style-type: none"> Biological explanation Typical vs. variant DNA sequence(s) Percent of 23andMe customers with variant References [1, 7, 9, 13, 19] ClinVar*

* The percent of 23andMe customers with a variant may not be representative of the general population.

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 Interpretation

This report provides risk estimates for prostate cancer associated with the HOXB13 G84E variant. This test does not take into account non-genetic factors that influence overall prostate cancer risk.

Health Risk Estimates

Risk estimates are based on clinical studies that identify an association between a genotype and a health condition.

Numerical risk estimates are not available for people who have two copies of the HOXB13 G84E variant. An interpretation of "increased risk" is provided to males with this result. It is likely that their risk is at least as high as the risk for males with just one copy of the variant. More research is needed to understand the risk associated with this result.

Consider talking to a healthcare professional if you have any concerns about your results.

References [1, 9, 13, 18]

Lifetime risk

Risk by age

The risk estimates for prostate cancer shown below represent the proportion of males expected to develop prostate cancer during their lifetime (for the general population) or by age 80 (for those with the HOXB13 G84E variant). Estimates for the general population are based on observed prostate cancer among males in the United States. Estimates for males with the HOXB13 G84E variant are based primarily on studies of people of European descent.

	General population	HOXB13 G84E variant (one copy) ⓘ
Prostate cancer	12.6%	33-53%

See risk estimates by ethnicity for the general U.S. population

Test Details

Indications for Use

The 23andMe Personal Genome Service (PGS) uses qualitative genotyping to detect select clinically relevant variants in genomic DNA isolated from human saliva collected from individuals ≥18 years for the purpose of reporting and interpreting genetic health risks, including the 23andMe PGS Genetic Health Risk Report for Hereditary Prostate Cancer (HOXB13-Related). The 23andMe PGS Genetic Health Risk Report for Hereditary Prostate Cancer (HOXB13-Related) is indicated for reporting of the G84E variant in the HOXB13 gene. The report describes if a person has the G84E variant and if a male is at increased risk for prostate cancer. The variant included in this report is most common in people of European descent. The test report does not describe a person's overall risk of developing any type of cancer, and the absence of a variant tested does not rule out the presence of other variants that may be cancer-related. This test is not a substitute for visits to a healthcare provider for recommended screenings or appropriate follow-up and should not be used for diagnosis or to determine any treatments or medical interventions.

Special Considerations

- Genetic testing for the HOXB13 G84E variant in the general population is not currently recommended by any healthcare professional organizations.
- Prostate cancer risk associated with the HOXB13 G84E variant varies from person to person. Overall risk depends on family history and other factors.

Test Performance Summary

Clinical Performance

[9, 19]

The variant included in this report is associated with an increased risk of developing prostate cancer. However, not everyone with this variant develops prostate cancer. In addition, most cases of prostate cancer are not caused by inherited genetic variants.

- The HOXB13 G84E variant accounts for up to 5% of hereditary prostate cancer in families of European descent. In some Scandinavian countries this variant accounts for about 22% (Finland) and 8% (Sweden) of hereditary prostate cancer cases.
- Up to 1 in 70 people of European descent has the HOXB13 G84E variant. This variant is expected to be less common in people of other ethnicities.

Analytical Performance

Accuracy was determined by comparing results from this test with results from sequencing. Greater than 99% of test results were correct. The 95% confidence interval for the total number of samples tested was 83.2% to 100%. While unlikely, this test may provide false positive or false negative results. It is possible that the presence of certain mutations in your sample may interfere with the performance of this test. The effects of the interfering mutations on the performance of this test have not been studied. For more details on the analytical performance of this test, refer to the package insert.

Warnings, Precautions, and Limitations

- This test does not diagnose cancer or any other health conditions and cannot determine your overall risk of developing cancer in the future.
- This test should not be used to make medical decisions. Results should be confirmed by an independent genetic test prescribed by your own healthcare provider before taking any medical action.
- This test does not cover all variants that could increase risk for cancer.* The absence of a variant tested does not rule out the presence of other genetic variants that may impact cancer risk.
- Other factors, such as environmental and lifestyle risk factors, may affect your risk of developing cancer. This test does not account for those factors, and does not test for variants in other genes linked to hereditary cancers.
- Your ethnicity may affect how relevant this test is for you. The variant included in this test is most commonly found in people of European descent, especially in people of Northern European descent.
- This test is intended to provide you with genetic information to inform conversations with your doctor or other healthcare professional.
- This device is not intended for prenatal testing.
- This test should not be used to assess the presence of genetic variants that may impact response to medications.
- This test is not intended to detect the presence of deterministic variants in autosomal dominant diseases or conditions.
- This test is not a substitute for visits to a healthcare professional for recommended screenings. Consult with a healthcare professional if you have any questions or concerns about your results or your current state of health.
- Some people feel a little anxious after getting genetic health risk results. This is normal. If you feel very anxious, you should speak to your doctor or a genetic counselor.

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

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

References

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See all references ▼

Change Log

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

Date	Change
March 22, 2023	Hereditary Prostate Cancer (HOXB13-Related) report created.



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[Overview](#)[Scientific Details](#)[Frequently Asked Questions](#)

Hereditary Prostate Cancer (HOXB13-Related)

What does this test do?



What does this test **not** do?



Who is at risk for prostate cancer?



The report says the variant included in this test is most common in people of **European** descent, especially in people of **Northern European** descent. What if I'm not of European descent?



Where can I learn more about prostate cancer, support groups, and other resources?



My report says **zero variants** were detected. What does this mean?



My report says **zero variants** were detected. Does this mean my male family members are not at risk of developing prostate cancer?



My report says **zero variants** were detected, but I have a family history of prostate cancer. What does this mean for my male family members?



My report says **zero variants** were detected. What are some things I could do?



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