

MUTYH-Associated Polyposis

MUTYH-associated polyposis (MAP) is one of the three main hereditary colorectal cancer syndromes. People with two variants or two copies of a variant in the MUTYH gene tend to develop colon and rectal polyps and have an increased risk of developing colorectal cancer. They may also have a slightly increased risk of developing certain other cancers. This test includes two genetic variants in the MUTYH gene that are most common and best studied in people of Northern European descent.

map2_zero_variants, you do not have the two genetic variants we tested.

However, this test does not include the majority of variants known to increase colorectal cancer risk. In addition, most cases of colorectal cancer are not caused by inherited variants, so people without a variant are still at risk of developing colorectal cancer. It's important to continue with any cancer screenings your healthcare provider recommends.



If you have a personal or family history of colorectal cancer or colorectal polyps, you should talk to a healthcare professional, who may recommend additional screening or genetic testing options for you.

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 in a clinical setting 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 MUTYH-Associated Polyposis 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 the **Y179C** and **G396D** variants in the MUTYH gene. These variants are linked to MAP, which increases a person's risk of developing certain cancers.
- Provides information on whether a person's genetic result is associated with an increased risk for colorectal cancer and may be associated with a slightly increased risk for certain other cancers.

Limitations

- Does **not** test for all possible variants in the MUTYH gene. More than 100 variants in the MUTYH gene are known to increase colorectal cancer risk. Only two of those variants are included in this test.
- Does **not** test for variants in other genes linked to hereditary colorectal cancer syndromes, such as Lynch syndrome and familial adenomatous polyposis (FAP).
- Does **not** account for non-genetic factors, such as environment and lifestyle, that influence overall cancer risk.

Important Ethnicities

- The variants included in this test are most common and best studied in people of **Northern European** descent. However, these two variants have also been found in other ethnicities.

You **do not have** the two variants we tested linked to MAP.

However, people without these variants are still at risk for colorectal cancer because most cases of colorectal cancer are caused by other factors.



You could still have a variant not included in this test.

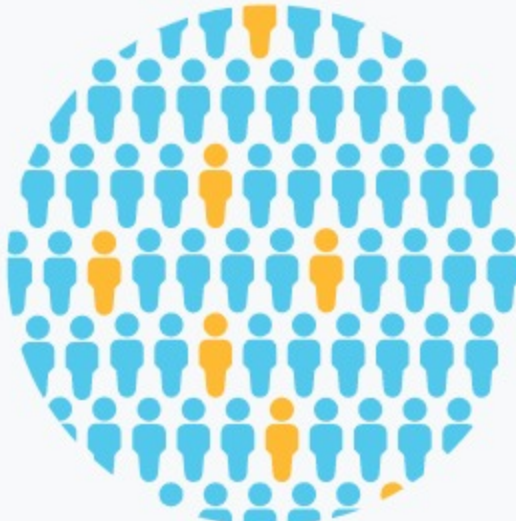
More than 100 variants in the MUTYH gene have been linked to MAP, which increases the risk for colorectal cancer. This test only includes two of those variants. In addition, this test does not include variants in other genes which can also increase colorectal cancer risk, including genes linked to Lynch syndrome and familial adenomatous polyposis (FAP).

[See Scientific Details](#)

In the general population, about **1 in 25** people will be diagnosed with colorectal cancer during their lifetime.

Less than 1% of colorectal cancer cases are caused by the two genetic variants in this report. Your risk is influenced by many other factors, including lifestyle, family history, and other genetic factors.

[See Scientific Details](#)



If you have a personal or family history of colorectal cancer or multiple colorectal polyps, talk to a healthcare professional about other testing options.

A genetic counselor can help you assess your overall cancer risk. [Learn more about genetic counseling.](#)

Lifestyle, family history, and other factors can also influence the chances of developing colorectal cancer.

Consult with a healthcare professional before making any major lifestyle changes.

Age

The risk of developing colorectal cancer generally increases with age. The majority of colorectal cancer cases are diagnosed in people over the age of 50.

[See Scientific Details for more information](#)

Age

Other genetic variants

Ethnicity

Family history

Inflammatory bowel disease

Lifestyle

About MUTYH-Associated Polyposis

MUTYH-associated polyposis (MAP) is a genetic condition where having two variants or two copies of a variant in the MUTYH gene increases a person's chance of developing colorectal cancer. This is because individuals with these variants are prone to developing colon and rectal polyps that, over time, can become cancerous. Variants in the MUTYH gene may also be associated with a slightly increased risk for certain other cancers.

When it develops

Most colorectal cancers start as abnormal growths on the inner lining of the colon or rectum, called polyps. People with MAP tend to develop between ten and a hundred polyps by age 50. These polyps can become cancerous. However, some people with MAP may develop colorectal cancer in the absence of colon or rectal polyps.

Lifetime cancer risks

- Studies suggest that people with MAP have a 43-100% chance of developing colorectal cancer in their lifetime without appropriate surveillance. These individuals may also have a slightly increased risk for certain other cancers.
- [See Scientific Details to learn more about these risks](#)

How common is MAP?

MAP accounts for about 1% of all colorectal cancer cases. Between 1 in 10,000 and 1 in 40,000 people of Northern European descent are expected to have MAP.

Screening and prevention

Professional guidelines recommend that individuals with two variants or two copies of a variant in the MUTYH gene should be screened for colon and rectal polyps earlier and more often, and undergo surveillance for small bowel polyps.

Current U.S. guidelines recommend that individuals with one MUTYH variant follow colorectal screening recommendations for the general population. However, for people who have had a first-degree relative with colorectal cancer and people who have a personal history of colorectal polyps (regardless of whether they have a MUTYH variant), these guidelines have different recommendations, which may include screening earlier and more often than the general population.

Read more at: [National Cancer Institute](#) [GeneReviews](#)

Learn more about MAP and colorectal cancer.



See our Frequently Asked Questions for more information.

FAQs



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

Print report



Learn more about colorectal cancer screening to help you and your doctor create a screening plan that's right for you.

Learn more

MUTYH-Associated Polyposis

Frequently Asked Questions

MUTYH-associated polyposis (MAP) is one of the three main hereditary colorectal cancer syndromes. People with two variants or two copies of a variant in the MUTYH gene tend to develop colon and rectal polyps and have an increased risk of developing colorectal cancer. They may also have a slightly increased risk of developing certain other cancers. This test includes two genetic variants in the MUTYH gene that are most common and best studied in people of Northern European descent.

MUTYH-Associated Polyposis

What does this test do?

This test looks for two specific genetic variants in the MUTYH gene, called Y179C and G396D. These variants are linked to MAP, which increases a person's risk of developing colorectal cancer.

This test provides information on whether a person's genetic result is associated with an increased risk for colorectal cancer and may also be associated with a slightly increased risk for certain other cancers.

This test does not include all possible variants in the MUTYH gene that may increase a person's risk of developing colorectal cancer.

This test does not include variants in other genes that are linked to other hereditary colorectal cancer syndromes, such as Lynch syndrome and familial adenomatous polyposis (FAP).

Is this answer helpful?

Yes

No

What does this test not do?

This test does not diagnose any type of cancer or any other health conditions. Only a healthcare professional can do that.

This test should not be used to make medical decisions. Results should be confirmed in a clinical setting before taking any medical action.

This test does not tell you if you have cancer or if you will definitely develop cancer in the future.

This test does not take into account other risk factors for colorectal cancer, such as personal and family health history. Thus, this test does not provide a complete assessment of your overall risk of developing colorectal cancer.

This test does not include all possible variants in the MUTYH gene that may increase a person's risk of developing colorectal cancer.

This test does not include variants in other genes that are linked to other hereditary colorectal cancer syndromes, such as Lynch syndrome and familial adenomatous polyposis (FAP).

Is this answer helpful?

Yes

No

The report says the variants included in this test are most common and best studied in people of Northern European descent. What if I'm not of Northern European descent?

Even though these two variants are most common in people of Northern European descent, they have also been observed in people of other ethnicities.

Similarly, even though the effect of these variants on a person's risk of developing colorectal cancer is best understood in people of Northern European descent, the effect is expected to be similar in people of other ethnicities. For example, if a person who is not of Northern European descent has both of the variants included in this report, he/she is still expected to have a similar elevated risk of developing colorectal cancer. [See Scientific Details for more information.](#)

Is this answer helpful?

Yes

No

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

You can learn more about **MAP** from the following resources:

- [Cancer.net \(American Society of Clinical Oncology\)*](#)

You can learn more about **colorectal cancer** from the following resources:

- [American Cancer Society*](#)
- [Colorectal Cancer Alliance*](#)
- [Fight Colorectal Cancer*](#)

If you have questions about your results or how they might affect you or your family, a genetic counselor may be able to help. [Learn more about genetic counseling.](#)

You can review the MUTYH-Associated Polyposis tutorial [here](#)*.

Is this answer helpful?

Yes

No

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

This means you do not have either of the two genetic variants we tested. However, it does **not** mean your colorectal cancer risk is reduced.

More than 100 variants in the MUTYH gene have been linked to MAP, which increases the risk for colorectal cancer. This report only includes two of those variants. In addition, this test does not include variants in other genes that can also increase colorectal cancer risk, including genes linked to Lynch syndrome and familial adenomatous polyposis (FAP). So you could still have a variant that is not included in this test.

About 1 in 25 people will be diagnosed with colorectal cancer during their lifetime, and the majority of these cases are not caused by inherited genetic variants. This means that other factors such as lifestyle, age, and family history are also important. For example, the risk for colorectal cancer is higher in people with a family history of colorectal cancer. [Learn more about other factors.](#)

Is this answer helpful?

Yes

No

My report says zero variants were detected. Does this mean I'm not at risk of developing colorectal cancer?

No. People with zero variants detected still have a risk of developing colorectal cancer.

More than 100 variants in the MUTYH gene have been linked to MAP, which increases the risk for colorectal cancer. This report only includes two of those variants. In addition, this test does not include variants in other genes that can also increase colorectal cancer risk, including genes linked to Lynch syndrome and familial adenomatous polyposis (FAP). So you could still have a variant that is not included in this test.

About 1 in 25 people will be diagnosed with colorectal cancer during their lifetime, and the majority of these cases are not caused by inherited genetic variants. This means that other factors such as lifestyle, age, and family history are also important. For example, the risk for colorectal cancer is higher in people with a family history of colorectal cancer. [Learn more about other factors.](#)

Is this answer helpful?

Yes

No

My report says zero variants were detected, but I have a personal or family history of colorectal cancer. What does this mean for me?

People with a family history of colorectal cancer have a higher risk of developing colorectal cancer themselves. Specifically, people with a [first-degree relative](#) who has been diagnosed with colorectal cancer are about twice as likely to develop colorectal cancer themselves, compared to the general population. The risk is even higher if that relative was diagnosed before the age of 45 or if there are two or more first-degree relatives who have developed colorectal cancer.

Even though you do not have the two genetic variants we tested, you could still have a variant that is not included in this test. More than 100 variants in the MUTYH gene have been linked to MAP, which increases the risk for colorectal cancer. This report only includes two of those variants. In addition, this test does not include variants in other genes that can also increase colorectal cancer risk, including genes linked to Lynch syndrome and familial adenomatous polyposis (FAP).

Other non-genetic factors may also influence your risk of developing colorectal cancer, even if you do not have any genetic variants. [Learn more about other factors.](#)

It is important to discuss your personal or family history of cancer with a healthcare professional, who can help you determine if additional genetic testing is appropriate. Genetic counseling can also help you understand your results and your options for additional testing. [Learn more about genetic counseling.](#)

Is this answer helpful?

Yes

No

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

Your genetic result means you do not have either of the two genetic variants we tested. However, because there are more than 100 genetic variants in the MUTYH gene linked to MAP and they only account for about 1% of colorectal cancer cases, your result doesn't give you much new information about your risk for colorectal cancer.

There are many other genetic and non-genetic factors that can affect your risk, which this test does not take into account. [Learn more about other factors.](#)

It is important to continue with any colorectal cancer screenings your healthcare provider recommends. Current U.S. guidelines recommend that people in the general population should start regular screening at age 50. Learn more from the [U.S. Preventive Services Task Force](#)*.

If you have a family history of colorectal cancer or a personal history of colorectal polyps, consider talking to a healthcare professional to learn more about options for screening and prevention. Your doctor may have specific screening recommendations for you, such as earlier or more frequent screening.

Talk to a healthcare professional if:

- You have a personal or family history of cancer.
- You think you might have colorectal polyps, colorectal cancer, or any other type of cancer.
- You have questions about other risk factors you may have.

Is this answer helpful?

Yes

No