A TEST FOR PATIENTS WITH COLORECTAL OR UTERINE CANCER

TumorNext-Lynch®
PATIENT GUIDE

Ambry Genetics®
A Konica Minolta Company
Germline vs. Tumor Testing

TumorNext-Lynch® looks for mutations and other changes in your tumor (also referred to as somatic testing) and inherited (germline) mutations in your blood.

It is important to understand the key differences between germline and tumor genetic tests, since they can give you very different information about your health and your family.

**WHAT’S THE DIFFERENCE?**

<table>
<thead>
<tr>
<th>WHAT IS TESTED?</th>
<th>INHERITANCE</th>
<th>RISKS</th>
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<tbody>
<tr>
<td><strong>INHERITED (GERMLINE)</strong></td>
<td>Blood or saliva</td>
<td>Can be inherited and passed on to family members</td>
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<tr>
<td></td>
<td>Genes that are identical in all cells of your body</td>
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<tr>
<td><strong>TUMOR (SOMATIC)</strong></td>
<td>Your tumor tissue for cancer-specific changes</td>
<td>Not inherited and only present in your tumor cells. Cannot be passed to family members</td>
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What is TumorNext-
hrd?

TumorNext-Lynch® tests for both germline (inherited) mutations that cause Lynch syndrome, as well as mutations and other changes in your tumor that may be able to rule out Lynch syndrome. The results of this testing can help your healthcare provider know more about your cancer risks, so they can make a better plan for how to care for you and your family members.

**HOW CAN TUMORNEXT-LYNCH® HELP?**

**GERMLINE (INHERITED)**

- Learning if you have Lynch syndrome can help you understand your future cancer risks and if you need to undergo more cancer screening or consider preventive options.
- Your test results could also impact your family members, as they may also have an increased cancer risk.

**TUMOR (SOMATIC)**

- Learning if you have certain genetic changes or mutations in your tumor can help to rule out a diagnosis of Lynch syndrome and can help your healthcare provider understand if you may or may not be a good candidate for certain treatments.

**WHO SHOULD HAVE TESTING WITH TUMORNEXT-LYNCH?**

Patients with a personal history of colorectal or uterine cancer who:

- Have tumor screening results that suggest a possible diagnosis of Lynch syndrome, like abnormal MSI or IHC* results results
- Have a personal and family history of cancer suspicious for Lynch syndrome

*WHAT IS MSI/IHC?*

Immunohistochemistry (IHC) and microsatellite instability analysis (MSI) are two tests performed on tumor tissue to screen for Lynch syndrome and they may be done automatically at some hospitals for patients with certain types of cancer, like colorectal or uterine. The results of these tests can tell us if a tumor has certain traits that may make it more likely for the patient to have Lynch syndrome, but it can’t diagnose or confirm that someone has the condition. When people have abnormal MSI or IHC results, additional testing, such as TumorNext-Lynch, is needed to confirm or rule out Lynch syndrome.
Many people have a family history of cancer, but only 5-10% of cancer is hereditary.

People who have these gene mutations are born with them – they do not develop over time.

Learning if you have an inherited mutation can help you know more about your cancer risks.

People with a higher chance of developing cancer may need screening, like colonoscopies, that start at younger ages, and occur more often.

Understanding The Basics

CANCER FALLS INTO 1 OF 3 CATEGORIES

**FAMILIAL CANCER**
A pattern of the same or related types of cancer within a family that may be due to genes and/or other shared factors, such as environment and lifestyle.

**HEREDITARY CANCER**
A pattern of the same or related types of cancer within a family due to inherited gene changes (called mutations), that can be passed from parent to child.

**SPORADIC CANCER**
Cancers that occur by chance in one or two family members that are not due to an inherited gene mutation. Approximately 80% of all cancers are sporadic.

ABOUT HEREDITARY CANCER
YOUR GENES CARRY A STORY THAT IS UNIQUE TO YOU AND MAKES YOU WHO YOU ARE. GENETIC TESTING CAN HELP YOU BETTER UNDERSTAND YOUR RISKS FOR CANCER.

Lynch syndrome is caused by inherited mutations in either MLH1, MSH2, MSH6, PMS2, or EPCAM. It is the most common cause of hereditary colorectal and uterine cancer and occurs in an estimated 1/279-1/440 Americans. People with Lynch syndrome have an increased risk for multiple types of cancer including colorectal, uterine, ovarian, and others. The graph below highlights some of the cancer risks for a person with Lynch syndrome compared to someone in the general population who does not have this condition.

**Lynch Syndrome Lifetime Cancer Risks (%)**

*Recent publications suggest lower risks for MSH6 and PMS2 mutation carriers*
Benefits of Tumor Next-Lynch®:

Provides more information that can help clarify whether or not you have Lynch syndrome, so that your healthcare provider knows how to best care for you and your family.

Benefits of Both Tumor and Germline Testing:

Your healthcare provider may be able to discuss additional personalized treatment options based on your genetic test results.

Benefits of Germline Genetic Testing:

Your healthcare provider can adjust your cancer screening plan (such as age of initial screening, type, and frequency) based on your genetic test results.

- An example of cancer screening is a colonoscopy.

Your healthcare provider may discuss possible cancer prevention options, such as preventive surgery to reduce the risk for certain cancers.

- An example is prophylactic hysterectomy and oophorectomy (removing the uterus, ovaries and Fallopian tubes before a cancer occurs).
Men and women have the same chance of inheriting a mutation, but their chance of developing cancer may be different.
Possible Genetic Test Results

**CONSISTENT WITH A DIAGNOSIS OF LYNCH SYNDROME**

- You are at an increased risk for cancers such as colorectal and uterine. Your healthcare provider will discuss the best management recommendations for you.

- Genetic testing for certain family members may be recommended.

**RULES OUT A DIAGNOSIS OF LYNCH SYNDROME**

- You were not found to have an inherited gene mutation that causes Lynch syndrome.

- If you had previously abnormal Lynch syndrome screening results (MSI/IHC), the testing explained these results and found that you most likely do not have Lynch syndrome.

**PERSONALIZED TREATMENT OPTIONS**

- Based on certain traits of your tumor, you may be a candidate for personalized treatment options, such as immunotherapy. Talk to your doctor about which treatment options may be right for you.

**INCONCLUSIVE RESULTS THAT CANNOT CONFIRM OR RULE OUT A DIAGNOSIS OF LYNCH SYNDROME**

- Cancer risk(s) and treatment recommendations are based on personal and family history.

It is possible to have a combination of positive and VUS results, since multiple genes are tested.
Resources For You

Ambry’s Patient Education Website
ambrygen.com/patient

Cancercare
cancercare.org

American Cancer Society
cancer.org

Genetic Information Nondiscrimination Act
ginahelp.org

American Society of Clinical Oncology
cancer.net

National Cancer Institute
cancer.gov

FIND A GENETIC COUNSELOR

National Society of Genetic Counselors
nsgc.org

Canadian Association of Genetic Counsellors
cagc-accg.ca
1 HOW IS GENETIC TESTING PERFORMED AND HOW LONG DOES IT TAKE?

Genetic testing requires a blood or saliva sample, which is collected using a special kit that is shipped overnight to Ambry Genetics by your healthcare provider. The testing, which looks for mutations that cause an increased risk for cancer, takes less than three weeks to complete, and results are sent to your healthcare provider.

2 WHAT WILL HAPPEN WHEN MY RESULTS ARE READY?

Your healthcare provider will receive your results; they will not be sent directly to you. Every healthcare provider may have a different method and time frame for reviewing your results with you, so it is important to discuss this process with them when your test is performed. Your healthcare provider will discuss recommended next steps based on your test results.

3 WILL MY GENETIC TEST RESULTS AFFECT MY INSURANCE COVERAGE?

In the U.S., the Genetic Information Nondiscrimination Act (2008) prohibits discrimination by health insurance companies and employers, based on genetic information. Depending on where you live in the world, you may have different (or fewer) laws in this area. Visit ginahelp.org to learn more.

4 HOW WILL MY TEST RESULTS BE PROTECTED?

We are required by law to maintain the confidentiality of your protected health information in accordance with the Health Insurance Portability and Accountability Act (HIPAA). Visit HHS.gov to learn more.
5 SHOULD I TELL MY FAMILY MEMBERS ABOUT MY GENETIC TEST RESULTS?

It is important to share your results with your family members, because they may provide additional information about their own cancer risks and management options. If you feel unsure about how to approach the subject, your healthcare provider may be able to offer some advice.

6 WILL GENETIC TESTING BE COVERED BY MY INSURANCE?

Many insurance plans cover genetic testing, and Ambry Genetics is contracted with the majority of U.S. health plans. Your out-of-pocket cost may vary based on your individual plan. A team of dedicated specialists is available to help you get access to the genetic testing you need, and provide further details about our payment options. Please call or email our Billing department at +1.949.900.5795 or billing@ambrygen.com with any questions. Visit ambrygen.com/patientbilling for more information.

7 WHAT IS AN EXPLANATION OF BENEFITS (EOB)?

Your insurance company sends you an EOB to explain any services paid on your behalf. You can contact us directly to speak with a Billing specialist with any questions or concerns about Ambry Genetics genetic testing that appears on your EOB. It is important to remember that insurance companies can take several weeks or even a couple of months to process claims.

STILL HAVE QUESTIONS?

Talk to your healthcare provider or visit our website: ambrygen.com