Consent for Hereditary Cancer Genetic Testing for Lynch Syndrome

Introduction. This form describes the benefits, risks, and limitations of genetic testing for inherited susceptibility to cancer. Please read this form carefully and discuss any questions that you have with your team of providers.

Purpose. This test analyzes a specific gene or genes for genetic changes called mutations. The gene(s) analyzed are associated with a specific hereditary cancer syndrome called Lynch Syndrome, or Hereditary Nonpolyposis Colorectal Cancer Syndrome (HNPCC). This test will help determine if a person has a significantly increased risk of developing certain tumors due to a mutation(s) in a cancer-predisposing gene.

Lynch Syndrome is a hereditary cancer syndrome caused by alteration in any one of a class of genes called mismatch repair (MMR) genes. There are five known genes: The mismatch repair genes: MLH1, MSH2, MSH6, and PMS2, and a related gene known as EPCAM. The majority of cases of Lynch Syndrome are due to mutations in these genes. Individuals who carry a Lynch Syndrome gene mutation have a greater risk to develop certain types of cancer than those who do not have a mutation. However, not everyone who has an altered gene will develop cancer.

The cancer risks associated with a Lynch Syndrome gene mutation are listed below. The percentages cited are estimates of lifetime risk and may change as more research is done.


Both men and women with a Lynch Syndrome gene mutation have a 50% (1 in 2) chance of passing it on to each of their children. There is no evidence that being born with a Lynch Syndrome gene mutation increases the risk for childhood cancers.

Test Procedure. Hartford Hospital personnel will draw approximately 2 tubes of blood and send them to the appropriate genetic testing laboratory. The laboratory will analyze your Lynch Syndrome genes to look for mutations. No other tests will be run on your sample without your written consent.

You are consenting to the following tests:

- Comprehensive Lynch Syndrome testing: sequencing and large rearrangement analysis
- Gene specific analysis: MLH1, MSH2, MSH6, PMS2 or EPCAM
- Single site analysis

___________Patient initials

Test Results and Interpretation. Listed below are the possible results from testing:

- Positive – A mutation that is associated with an increased risk for hereditary cancer was identified. Knowing this information may help you and your doctor make more informed choices about your health care, such as screening, risk-reducing surgeries and preventive medication strategies.
- Negative – A mutation was not identified.
  - If you are the first person tested in your family, you still have at least the same risk of cancer as does a person in the general population. You may still be at greater than average risk for hereditary cancer due to a genetic predisposition that cannot be detected by this test, either in the gene(s) you were tested for or in another gene linked to hereditary cancer.
  - If you test negative for a mutation known to be in your family, you are considered to have the same risks as others in the general population.
- Uncertain – A genetic change (a Variant of Uncertain Significance, or VUS) was detected but it is not known if this change is linked to cancer risk. You still have at least the same risk of cancer as the general population, and may still be at greater than average risk due to a genetic predisposition that cannot be detected by this test.

The genetic counselor will discuss the results with you and what they mean. Genetic test results have implications for blood relatives. If you choose, the genetic counselor will help you inform family members if a mutation is found.

Benefits. Your genetic test results may help you and your doctor make more informed choices about your health care, such as screening, risk-reducing surgeries and preventive medication strategies. The identification of gene mutation(s) in a family enables other blood relatives to determine whether or not they share the same hereditary cancer risks. If you test positive, you should discuss with your healthcare provider how hereditary cancer is inherited and learn about the chance your children and blood relatives may have inherited the same mutation(s) in the gene(s) tested. If you test negative for a known mutation in your family, you cannot pass on that mutation to your children. If you have had cancer or a pre-cancerous condition, you would still have some increased risk based on your own history. If you have not had cancer, you would still have the general population risk of cancer.
Risks. Learning that you have a genetic mutation may be upsetting and may increase feelings of depression, anxiety, and vulnerability. You could feel that you have to make difficult decisions about your medical care. Learning of a positive result may affect relationships with family members. News of an inconclusive result may increase feelings of distress. If more than one family member is tested, there is a possibility of learning sensitive information about your family, such as non-paternity or undisclosed adoption.

To address concerns regarding possible health insurance discrimination, most states and the federal government have enacted laws to prohibit genetic discrimination. Furthermore, broad federal legislation (HIPAA) prohibits unauthorized disclosure of confidential personal health information. However, for the most part, these laws do not prevent life and disability insurers from using genetic testing information in determining coverage. **It is our policy to include genetic information and test results into your medical record.**

Limitations. This test analyzes only the Lynch Syndrome genes. If no mutation is found, you may still be at risk for hereditary cancer due to a genetic predisposition that cannot be detected by this test, either in the gene(s) you were tested for or in another gene linked to hereditary cancer. Genetic testing clarifies cancer risks for only those cancers related to the genes analyzed.

Financial Responsibility. Genetic testing of appropriate individuals is typically reimbursed by health insurance. You are responsible for any cost of the genetic test not reimbursed by insurance, including all applicable co-pays and deductibles.

Future Correspondence. Due to the dynamics of this field, there continues to be new information and data. It is recommended that you keep in contact with your healthcare provider, annually, to learn of any new developments in cancer genetics and to provide any updates to your personal or family history which may affect your cancer risks.

I confirm that the possible benefits, risks, limitations and costs of genetic testing for Lynch Syndrome have been explained to me. All my questions have been answered. I have read this consent form and will be given a copy for my records. My signature below indicates my willingness to have genetic testing for Lynch Syndrome gene mutation(s). I understand the confidentiality policy outlined in the consent form. This means that I recognize that genetic information and a copy of the test result will be included in my medical record.

Signature of Patient ___________________________ Date ___________ Time ___________

Print name

I have explained the possible risks, limitations and costs as well as potential benefits of genetic testing for Lynch Syndrome gene mutations, and have answered any questions regarding the test to the best of my ability.

Genetic Counselor/Clinician ___________________________ Date ___________ Time ___________

Patient initials ___________