Hereditary Lynch/Colorectal Cancer

Genetic testing with the Lynch/Colorectal High Risk Panel may be appropriate if your personal and/or family history is suggestive of Lynch syndrome or a hereditary predisposition to colorectal cancer and/or polyps. This includes:

- Colorectal or endometrial cancer* diagnosed under 50 years of age
- Multiple colon polyps (especially ≥ 20 adenomas) at any age
- Tumor testing which indicates an increased risk for a hereditary cancer syndrome known as Lynch syndrome (e.g. microsatellite instability and/or lack of immunohistochemistry staining for a mismatch repair protein)
- Multiple cancers in one person either of the same origin (such as two separate colorectal cancers) or of different origin (such as colon and endometrial cancer in the same individual)
- Multiple relatives diagnosed with the same or related cancers (such as colon, endometrial, ovarian, urinary tract, gastric) on the same side of the family and spanning multiple generations

Your healthcare provider will determine if genetic testing is medically necessary for you.

*if this is the primary indication for testing, a more comprehensive panel specifically geared at this diagnosis is available

Genes Included on the Lynch/Colorectal High Risk Panel are Listed in the Table Below

<table>
<thead>
<tr>
<th>Gene</th>
<th>Lifetime Cancer and/or Tumor Risks*</th>
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</thead>
<tbody>
<tr>
<td>APC</td>
<td>Colorectal (up to 93%), Duodenal or periampullary (4-12%), Gastric, Thyroid (up to 3%), Pancreatic, Brain-medulloblastoma, Liver-hepatoblastoma, Desmoid tumors, Gastrointestinal polyps</td>
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<tr>
<td>EPCAM**</td>
<td>Colorectal (69-75%), Endometrial (12-55%), Ovarian, Gastric, Pancreatic, Biliary tract, Urinary tract-transitional cell, Small bowel, Brain, Sebaceous neoplasms, Prostate</td>
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<tr>
<td>MLH1</td>
<td>Colorectal (34-46%), Endometrial (18-54%), Ovarian (10-20%), Gastric (6-20%), Urinary tract-transitional cell (1-4%), Pancreatic (1-4%), Biliary tract (2-3%), Small bowel (4-12%), Brain, Sebaceous neoplasms, Prostate</td>
</tr>
<tr>
<td>MSH2</td>
<td>Colorectal (37-48%), Endometrial (21-57%), Ovarian (10-24%), Urinary tract-transitional cell (8-20%), Gastric (&lt;1-9%), Pancreatic (1-4%), Biliary tract, Small bowel (1%), Brain, Sebaceous neoplasms, Prostate</td>
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<tr>
<td>MSH6**</td>
<td>Colorectal (20-44%), Endometrial (16-71%), Ovarian (1-13%), Gastric, Pancreatic, Biliary tract, Urinary tract-transitional cell, Small bowel, Brain, Sebaceous neoplasms, Prostate</td>
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<tr>
<td>MUTYH</td>
<td>Colorectal (up to 80%), Duodenal (up to 4%), Gastrointestinal polyps</td>
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<tr>
<td>PMS2**</td>
<td>Colorectal (11-20%), Endometrial (12-26%), Ovarian, Gastric, Pancreatic, Biliary tract, Urinary tract-transitional cell, Small bowel, Brain, Sebaceous neoplasms, Prostate</td>
</tr>
</tbody>
</table>

*Most commonly associated cancer/tumors listed; lifetime risks provided when available. Risks relate to carriers of a single pathogenic variant with the exception of the MUTYH.

**Tumor spectrum is representative of Lynch syndrome; data are limited with regard to the association of certain cancers with pathogenic variants in MSH6, PMS2 and EPCAM.
Medical Management Based on Genetic Test Results

Clinical guidelines may be available which provide options and recommendations for patients who have a positive (pathogenic or likely pathogenic variant) test result indicating an increased risk for cancer and/or tumors. Guidelines and recommendations for early detection and/or risk reduction are specific to the gene in which the pathogenic variant was found.

Recommendations may include:
• Clinical exams, such as skin or eye exams
• Urine analysis
• Imaging exams, such as a MRI, CT and/or ultrasound
• Screening procedures, such as a colonoscopy or endoscopy
• Risk-reducing surgery

In some cases, guidelines for screening and prevention are limited or not available for a positive result. Once your test results are available, a discussion with your healthcare provider is recommended to determine the most appropriate medical management options for you and your family.

Possible Outcomes of Genetic Testing

Positive
• Pathogenic or likely pathogenic variant identified
• Medical management recommendations may be available
• Family member testing may be recommended

Negative
• No significant genetic changes identified
• Medical management based on personal and/or family history

Variant of Uncertain Significance (VUS)
• A genetic change identified, but its association with disease is unclear
• Medical management based on personal and/or family history

Resources

General
American Cancer Society
www.cancer.org
GeneDx
www.genedx.com/ oncology
National Cancer Institute
www.cancer.gov

Colorectal Cancer
Colon Cancer Alliance
www.ccalliance.org
Fight Colorectal Cancer
www.fightcolorectalcancer.org
Hereditary Colon Cancer Takes Guts
www.hcctakesguts.org
Colon Cancer Alliance for Research and Education for Lynch Syndrome (CCARE)
www.fightlynch.org

Find a Genetic Counselor
Canadian Association of Genetic Counsellors
www.cagc-accg.ca
National Society of Genetic Counselors
www.nsgc.org