Policy: MP098
Section: Medical Benefit Policy
Subject: Genetic Testing Related to Colorectal Cancer

I. Policy: Genetic Testing Related to Colorectal Cancer

II. Purpose/Objective:
   To provide a policy of coverage regarding Genetic Testing Related to Colorectal Cancer

III. Responsibility:
   A. Medical Directors
   B. Medical Management

IV. Required Definitions

1. Attachment – a supporting document that is developed and maintained by the policy writer or department requiring/authoring the policy.
2. Exhibit – a supporting document developed and maintained in a department other than the department requiring/authoring the policy.
3. Devised – the date the policy was implemented.
4. Revised – the date of every revision to the policy, including typographical and grammatical changes.
5. Reviewed – the date documenting the annual review if the policy has no revisions necessary.

V. Additional Definitions
Medical Necessity or Medically Necessary means Covered Services rendered by a Health Care Provider that the Plan determines are:

   a. appropriate for the symptoms and diagnosis or treatment of the Member's condition, illness, disease or injury;
   b. provided for the diagnosis, and the direct care and treatment of the Member's condition, illness disease or injury;
   c. in accordance with current standards of good medical treatment practiced by the general medical community.
   d. not primarily for the convenience of the Member, or the Member's Health Care Provider; and
   e. the most appropriate source or level of service that can safely be provided to the Member. When applied to hospitalization, this further means that the Member requires acute care as an inpatient due to the nature of the services rendered or the Member's condition, and the Member cannot receive safe or adequate care as an outpatient.

Medicaid Business Segment
Medical Necessity shall mean a service or benefit that is compensable under the Medical Assistance Program and if it meets any one of the following standards:

   (i) The service or benefit will, or is reasonably expected to, prevent the onset of an illness, condition or disability.
   (ii) The service or benefit will, or is reasonably expected to, reduce or ameliorate the physical, mental or development effects of an illness, condition, injury or disability.
   (iii) The service or benefit will assist the Member to achieve or maintain maximum functional
DESCRIPTION:
Genetic testing involves the analysis of DNA, RNA, chromosomes, proteins, and certain metabolites in order to detect heritable disease-related genotypes. There are currently two well-defined types of hereditary colorectal cancer, familial adenomatous polyposis (FAP) and hereditary nonpolyposis colorectal cancer (HNPCC).

INDICATIONS:
*REQUIRES PRIOR MEDICAL DIRECTOR or DESIGNEE AUTHORIZATION
- Genetic testing to determine the carrier status of the adenosis polyposis coli (APC) gene when criteria are met. (See applicable medical criteria below)
- Genetic testing to determine the carrier status of the MutY homolog [MYH] when the member meets criteria (*See applicable medical criteria below)
- Genetic testing to determine the carrier status of the HNPCC associated genes when either the Amsterdam or Bethesda criteria is met. (*See applicable medical criteria below)

MEDICAL CRITERIA:
For APC gene testing for familial adenomatous polyposis (FAP) and Attenuated FAP (AFAP):
To determine carrier status of the adenomatous polyposis coli gene (APC) for familial adenomatous polyposis (FAP) or attenuated familial adenomatous polyposis (AFAP) in the following subjects:
1. Individuals with greater than 10 adenomatous colonic polyps in their lifetime; OR
2. In first-degree relatives (e.g., siblings, parents, offspring) of an individual diagnosed with FAP or AFAP; OR
3. Personal history of desmoid tumor

For MutY human homolog [MYH]) gene testing for MYH-associated polyposis (MAP):
MYH-associated polyposis (MAP) genetic testing (gene MutY human homolog [MYH]) is covered in ANY of the following situations:
- Confirmatory testing for individuals with a history of adenomatous polyposis (>10 adenomas) and negative APC mutation testing: or
- For predictive testing when an individual has a sibling with known MYH polyposis; or
- For predictive testing when an individual has at least one affected sibling with findings consistent with recessive inheritance (i.e., MAP)

For HNPCC; Lynch Syndrome genetic testing (gene MLH1, MSH2, MSH6, PMS2, EPCAM):
NOTE: COLARIS Test® is a patented test for assessment of colorectal cancer risk. It detects mutations in MLH1, MSH2, PMS2 and EPCAM genes. COLARIS AP detects mutations in the APC and MYH genes.

Hereditary non-polyposis colorectal cancer (HNPCC)/Lynch syndrome (LS): (MLH1, MSH2, MSH6, PMS2, EPCAM sequence analysis) gene testing is considered medically necessary for members who meet any one of the following criteria:
The member:
- is diagnosed with colorectal cancer with the MSI histology at any age; or
- is diagnosed with endometrial cancer before age 50 years; or
- meets Amsterdam or Revised Bethesda Guidelines; or
- is diagnosed with a synchronous, or metachronous Lynch Syndrome (LS)-associated tumors*, regardless of age; or
- has a 1st- or 2nd-degree relative with a disease confirmed to be caused by a HNPCC mutation (genes MLH1, MSH2, MSH6, PMS2, EPCAM); or
- has ≥5% risk of LS on a validated mutation prediction model (eg, MMRpro, PREMM, MMRpredict)
  - [http://premm.dfci.harvard.edu/](http://premm.dfci.harvard.edu/)
  - [http://hnpccpredict.hgu.mrc.ac.uk/](http://hnpccpredict.hgu.mrc.ac.uk/)
  - [http://www4.utsouthwestern.edu/breasthealth/cagene/](http://www4.utsouthwestern.edu/breasthealth/cagene/).

* Lynch syndrome-related cancers include colorectal, endometrial, gastric, ovarian, pancreas, ureter and renal pelvis, biliary tract, brain (usually glioblastoma), and small intestinal cancers, as well as sebaceous gland adenomas and keratoacanthomas
<table>
<thead>
<tr>
<th>Amsterdam II Criteria</th>
<th>Revised Bethesda Guidelines</th>
</tr>
</thead>
<tbody>
<tr>
<td>Must Meet ALL of the following:</td>
<td>Must meet at least ONE of the following criteria:</td>
</tr>
<tr>
<td>• Three or more relatives, one of whom is a first-degree relative, with any of several histologically verified HNPCC-associated cancers (colorectal, endometrial, small bowel, ureter or renal pelvis); and</td>
<td>• Cancer in families that meet the Amsterdam criteria; or</td>
</tr>
<tr>
<td>• Colorectal cancer involving at least two generations; and</td>
<td>• Two HNPCC-related cancers, including synchronous and metachronous colorectal cancers or associated extracolonic cancers (biliary, endometrial, urinary or ovarian); or</td>
</tr>
<tr>
<td>• One or more colorectal cancers diagnosed before 50 years of age.</td>
<td>• Colorectal cancer and a first-degree relative with colorectal cancer and/or HNPCC-related extracolonic cancer and/or a colorectal adenoma; one of the cancers diagnosed before age 50 years, and the adenoma diagnosed before age 40 years; or</td>
</tr>
<tr>
<td>• Familial adenomatous polyposis (FAP) should be excluded in the colorectal cancer case(s), if any exists.</td>
<td>• Colorectal cancer or endometrial cancer diagnosed before age 50 years; or</td>
</tr>
<tr>
<td></td>
<td>• Right sided colorectal cancer with undifferentiated pattern (solid/cribiform), on histology, diagnosed before age 45 years; or</td>
</tr>
<tr>
<td></td>
<td>• Signet ring cell type colorectal cancer diagnosed before age 50 years; or</td>
</tr>
<tr>
<td></td>
<td>• Adenomas diagnosed before age 40 years; or</td>
</tr>
<tr>
<td></td>
<td>• Asymptomatic individuals with a first or second degree relative with a documented HNPCC mutation.</td>
</tr>
</tbody>
</table>

**Microsatellite instability (MSI) Testing or immunohistochemical (IHC) Analysis**

Microsatellite instability (MSI) testing or immunohistochemical (IHC) analysis of the tumor is considered medically necessary when any of the following criteria are met:

The member has:

- CRC diagnosed <70 yrs; or
- CRC diagnosed > 70 yrs and meets Bethesda guidelines; or
- endometrial cancer diagnosed before age 50 years

MSI testing or IHC analysis should be used as an initial test in members with colorectal or endometrial cancer to identify those who should proceed with HNPCC mutation analysis.

*NCCN Guidelines v1.2017 Lynch Syndrome*: the panel recommends tumor testing with IHC and/or MSI be used as the primary approach for pathology-lab-based universal screening. If tumor is available, LS-specific testing or multi-gene testing without IHC or MSI should be utilized in select cases under direction of a clinician with expertise in genetics, and should not be used as a universal screening strategy.

**COLOGUARD TESTING:**

**Fecal DNA Testing**: *(e.g., Cologuard.) DOES NOT REQUIRE PRIOR AUTHORIZATION* a noninvasive, multitarget fecal DNA test for the qualitative detection of colorectal neoplasia-associated DNA markers in addition to the presence of occult hemoglobin in stool is covered as a preventive screening methodology once every 3 years according to the following criteria:

- Age 45 to 85 years; and
- Asymptomatic (no signs/symptoms including but not limited to, lower gastrointestinal pain, blood in stool, positive guaiac fecal occult blood test, or fecal immunochemical test); and
- There has been no documentation of a normal colonoscopy in the previous 10 years; and
- At average risk of developing CRC defined as:
  - no personal history of adenomatous polyps, colorectal cancer, or inflammatory bowel disease including Crohn’s disease and ulcerative colitis; and
  - no family history of colorectal cancer or adenomatous polyps, familial adenomatous polyposis, or hereditary nonpolyposis CRC
Genetic testing is appropriate only when offered in a setting where a licensed or certified genetic counselor* or adequately trained health care professional is able to provide appropriate pre- and post-test genetic counseling, and medical necessity is supported by ALL of the following criteria:

1. The information is needed to adequately assess risk; and
2. The information will be used in the immediate care plan; and
3. Pedigree analysis establishes a high risk group for the disease; or
4. Clinical presentation of symptomology is evident and diagnosis cannot be established with conventional evaluation testing.

*A genetic counselor is considered by the Plan to be qualified if the following are met:

- M.S. or Ph.D. degree from a genetic counseling program approved/recognized by the American Board of Genetic Counseling or the American Board of Medical Genetics.
- Board certified or board qualified/eligible in the orderly process of obtaining board certification by the American Board of Genetic Counseling or American Board of Medical Genetics
- Proof of current competence and demonstrated ability (minimum of two years recent and continual experience within the past three years).

EXCLUSIONS:
There is no evidence to support the use of genetic testing for individuals from the general population with average risk. This is considered NOT MEDICALLY NECESSARY.

The ColonSentry testing panel is considered experimental, investigational, or unproven and is NOT COVERED. There are no current evidence-based guidelines from medical professional organizations or public health agencies that recommend ColonSentry for colorectal cancer screening and no evidence to support the use of this testing.

For Commercial and Medicaid Business Segments: The Epi proColon test is considered experimental, investigational, or unproven and is NOT COVERED. The Geisinger Technology Assessment Committee evaluated this technology and concluded that there is insufficient evidence in the peer-reviewed published medical literature to establish the effectiveness of this test on health outcomes when compared to established tests or technologies.

For Medicare Business Segment: Epi proColon testing is covered as an alternative colorectal cancer screening strategy.

Note: A complete description of the process by which a given technology or service is evaluated and determined to be experimental, investigational or unproven is outlined in MP 15 - Experimental Investigational or Unproven Services or Treatment.

PROCESS:
The Plan will utilize available published, peer reviewed medical literature, independent technology assessment program reports and/or review by the Geisinger Clinic Technology Assessment Committee to evaluate the following criteria when assessing the validity and efficacy of a specific genetic test:

- The analytical power of a test determined by its sensitivity and specificity is established.
- There is evidence of defined outcome measures that permits conclusions concerning the effect on health outcomes.
- The net health outcomes benefit of the test as compared to established alternatives is evaluated if applicable.
- The availability of test to membership is evaluated.
- The appropriate governmental and/or professional regulatory body approval is established.
- The test result will influence the treatment or alter the management of the member medical care plan.
- The test result has the capability to confirm a diagnosis when conventional medical evaluation is equivocal.

CODING ASSOCIATED WITH: Genetic Testing Related to Colorectal Cancer
The following codes are included below for informational purposes and may not be all inclusive. Inclusion of a procedure or device code(s) does not constitute or imply coverage nor does it imply or guarantee provider reimbursement. Coverage is determined by the member specific benefit plan document and any applicable laws regarding coverage of specific services. Please note that per Medicare coverage rules, only specific CPT/HCPCS Codes may be covered for the Medicare Business Segment. Please consult the CMS website at www.cms.gov or
the local Medicare Administrative Carrier (MAC) for more information on Medicare coverage and coding requirements.

81201 APC (ADENOMATOUS POLYPOSIS COLI) (EG, FAMILIAL ADENOMATOSIS POLYPOSIS [FAP], ATTENUATED FAP) GENE ANALYSIS; FULL GENE SEQUENCE
81202 APC (ADENOMATOUS POLYPOSIS COLI) (EG, FAMILIAL ADENOMATOSIS POLYPOSIS [FAP], ATTENUATED FAP) GENE ANALYSIS; KNOWN FAMILIAL VARIANTS
81203 APC (ADENOMATOUS POLYPOSIS COLI) (EG, FAMILIAL ADENOMATOSIS POLYPOSIS [FAP], ATTENUATED FAP) GENE ANALYSIS; DUPLICATION/DELETION VARIANTS
81298 MLH1 (mutL homolog 1, colon cancer, nonpolyposis type 2) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; full sequence analysis
81292 MLH1 (mutL homolog 1, colon cancer, nonpolyposis type 2) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; full sequence analysis
81293 MLH1 (mutL homolog 1, colon cancer, nonpolyposis type 2) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; known familial variants
81294 MLH1 (mutL homolog 1, colon cancer, nonpolyposis type 2) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; duplication/deletion variants
81295 MSH2 (mutS homolog 2, colon cancer, nonpolyposis type 1) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; full sequence analysis
81296 MSH2 (mutS homolog 2, colon cancer, nonpolyposis type 1) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; full sequence analysis
81297 MSH2 (mutS homolog 2, colon cancer, nonpolyposis type 1) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; known familial variants
81298 MSH6 (mutS homolog 6 [E. coli]) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; full sequence analysis
81299 MSH6 (mutS homolog 6 [E. coli]) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; known familial variants
81300 MSH6 (mutS homolog 6 [E. coli]) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; duplication/deletion variants
81301 Microsatellite instability analysis (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) of markers for mismatch repair deficiency (eg, BAT25, BAT26), includes comparison of neoplastic and normal tissue, if performed
81309 PIK3CA (phosphatidylinositol-4, 5-biphosphate 3-kinase, catalytic subunit alpha)(eg. colorectal and breast cancer) gene analysis, targeted sequence analysis (eg, exons 7,9, 20)
81317 PMS2 (postmeiotic segregation increased 2 [S. cerevisiae]) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; full sequence analysis
81318 PMS2 (postmeiotic segregation increased 2 [S. cerevisiae]) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; known familial variants
81319 PMS2 (postmeiotic segregation increased 2 [S. cerevisiae]) (eg, hereditary non-polyposis colorectal cancer, Lynch syndrome) gene analysis; duplication/deletion variants
81327 Sept9 (septin 9) (eg, colorectal cancer) methylation analysis
0101U Hereditary colon cancer disorders (eg, Lynch syndrome, PTEN hamartoma syndrome, Cowden syndrome, familial adenomatosis polyposis), genomic sequence analysis panel utilizing a combination of NGS, Sangar, MLPA, and array CGH, with MRNA analytics to resolve v
0155U (phosphatidylinositol-4, 5-biphosphate 3-kinase, catalytic subunit alpha)(eg. colorectal and breast cancer) gene analysis,


LINE OF BUSINESS:
Eligibility and contract specific benefits, limitations and/or exclusions will apply. Coverage statements found in the line of business specific benefit document will supersede this policy. For Medicare, applicable LCD’s and NCD’s will supersede this policy. For PA Medicaid Business segment, this policy applies as written.
REFERENCES:
This policy will be revised as necessary and reviewed no less than annually.
ECRI, HTAIS Windows on Medical Technology, Microsatellite Instability Testing for Hereditary Nonpolyposis Colorectal Cancer. Issue No. 64, Jan. 2002
ECRI, HTAIS Custom Hotline Response. Guidelines for Genetic Testing to Identify Persons at risk for Colorectal Cancer. Current as of 03/14/06.


Centers for Medicare & Medicaid Services. 2019 Clinical Lab Fee Schedule (CLFS) December 14, 2018

U.S. Food & Drug Administration. Premarket Approval (PMA). Cologuard Stool DNA-Based Colorectal Cancer Screening Test. Sept. 20, 2019

This policy will be revised as necessary and reviewed no less than annually.

Devised: 6/03

Revised: 7/23/04(criteria); 6/05 (update Bethesda criteria); 6/06; 6/07(revised criteria); 6/09 (coding); 6/10 (updated criteria); 8/12, 11/14 (Added medicare, Medicaid, healthy PA for Fecal DNA), 11/15 (added exclusion); 5/16 (expanded coverage for fecal DNA testing); 11/16 (remove PA for fecal DNA screening test); 4/17 (clarified criteria), 4/18 (criteria update); 7/19 (add Medicare Epi proColon coverage), 11/19 (update Cologuard age recommendation)

Reviewed: 7/08, 1/11, 1/12, 8/13, 8/14, 4/19