

[Revision log](#)

CONCERT GENETICS GENETIC TESTING: HEREDITARY CANCER SUSCEPTIBILITY

See [Important Reminder](#) at the end of this policy for important regulatory and legal information.

OVERVIEW

Genetic testing for hereditary cancer susceptibility is performed when an individual has risk factors that increase suspicion that they could develop an inherited form of cancer. These risk factors may include an individual's personal and/or medical histories, as well as their family medical history. When a genetic test is positive for hereditary cancer susceptibility, the individual is at an increased risk for cancer and this information may impact medical management, including screening, prevention, and treatment decisions.

Genetic testing for hereditary cancer susceptibility is a germline test and can be performed on individual genes (e.g., *BRCA1*) or on many genes simultaneously (i.e., multi-gene panels). Panels can range from a more limited number of genes associated with hereditary susceptibility to one specific type of cancer (e.g., breast cancer panel), or a pan-cancer hereditary cancer susceptibility panel (i.e., a panel that tests for many genes associated with hereditary cancer susceptibility at the same time).

Of note, the National Society of Genetic Counselors (NSGC) endorses the use of multi-gene panel tests when clinically warranted and appropriately applied. Specifically, the NSGC recommends thorough evaluation of the analytic and clinical validity of the test, as well as its clinical utility³. For this reason, several of the criteria in this policy require that panel tests do not include genes without known association with the disease in question.

Targeted mutation testing is the process of analyzing one single pathogenic or likely pathogenic (P/LP) variant in one gene. Generally, this type of testing is recommended when there is a known P/LP variant in an individual's close relative. Importantly, an individual meeting criteria for broader testing (i.e. full gene or multi-gene panel testing) based on clinical history should have

broader testing performed. Of note, if a variant of unknown significance (VUS) is detected in an individual, it is not recommended that family members also be tested for the VUS, unless the VUS is reclassified to a pathogenic or likely pathogenic variant.

Targeted germline genetic testing may also be recommended when there is a P/LP variant found on somatic tumor profiling. It should be noted that there is language in several National Comprehensive Cancer Network (NCCN) guidelines stating that somatic P/LP variants are common in some genes and may not indicate the need for germline testing unless the clinical/family history is consistent with a P/LP variant in the germline. However, given these tests are targeted and have significant implications for a patient's medical management, it is clinically appropriate to allow for a path to coverage for this type of testing.

POLICY REFERENCE TABLE

Coding Implications

This clinical policy references Current Procedural Terminology (CPT®). CPT® is a registered trademark of the American Medical Association. All CPT codes and descriptions are copyrighted 2022, American Medical Association. All rights reserved. CPT codes and CPT descriptions are from the current manuals and those included herein are not intended to be all-inclusive and are included for informational purposes only. Codes referenced in this clinical policy are for informational purposes only. Inclusion or exclusion of any codes does not guarantee coverage. Providers should reference the most up-to-date sources of professional coding guidance prior to the submission of claims for reimbursement of covered services.

. Please see the [Concert Genetics Platform](#) for a comprehensive list of registered tests.

Criteria Sections	Example Tests (Labs)	Common CPT Codes	Common ICD10 Codes	Ref
Pan-Cancer Hereditary Cancer Susceptibility Panels	MyRisk (Myriad Genetics)	81432, 81433	C15-26, C50-58, Z17, Z80, Z85.0-Z85.9	1, 2, 3, 11
	Common Hereditary Cancers Panel (Invitae)			
	CancerNext (Ambry Genetics)			
	Tempus xG Hereditary Cancer Panel			

	+RNAinsight with CancerNext (Ambry Genetics)	0134U		
Hereditary Breast Cancer Susceptibility Panels	VistaSeq Breast Cancer Panel (Labcorp) Breast Cancer Panel (Invitae) Breast Cancer STAT NGS Panel (Sequencing & Deletion/Duplication) (Fulgent Genetics) Breast Cancer - High Risk Panel (PreventionGenetics, part of Exact Sciences) Breast Cancer High-Risk Panel plus PALB2 (GeneDx)	81162, 81163, 81164, 81165, 81166, 81167, 81216, 81307, 81321, 81351, 81432, 81433	C50, Z80.3, Z83, Z84, Z85, Z86	1
	BRCAPlus (Ambry Genetics)	0129U		
Hereditary GI/Colon Cancer Panel Tests	Colorectal Cancer Panel (Invitae)	81435, 81436	C15-26, Z80, Z83, Z84, Z85, Z86	2
	ColoNext (Ambry Genetics)	0101U		
	+RNAinsight for ColoNext (Ambry Genetics)	0130U, 0162U		
Hereditary Gastric Cancer Panels	Invitae Gastric Cancer Panel (Invitae)	81201, 81203, 81292, 81294, 81295, 81297, 81298, 81300, 81317, 81319, 81403, 81404, 81405, 81406, 81408, 81479	C16, Z80, Z85, Z86	7
	Gastric Cancer Panel (PreventionGenetics, part of Exact Sciences)			
Hereditary Pancreatic Cancer Susceptibility Panels	Pancreatic Cancer Panel (Invitae)	81162, 81163, 81201, 81292, 81295, 81298, 81351, 81433, 81479	C25, Z80, Z84, Z85, Z86	1
	PancNext (Ambry Genetics)			
Hereditary Polyposis Panels	Hereditary Polyposis Panel (PreventionGenetics, part of Exact Sciences)	81201, 81203, 81406, 81479	D12, K63.5, Z80, Z84, Z85, Z86	2
	COLARIS AP (Myriad Genetics)			

Hereditary Prostate Cancer Susceptibility Panels	Prostate Cancer Panel-Primary Panel (Invitae)	81162, 81292, 81295, 81351, 81479	C61, Z80, Z84, Z85, Z86	1, 20
	ProstateNext (Ambry Genetics)			
	+RNAinsight for ProstateNext (Ambry Genetics)	0133U		
Hereditary Neuroendocrine Cancer Susceptibility Panels	Hereditary Paraganglioma-Pheochromocytoma Panel (Invitae)	81437, 81438	C74, C75, C7A Z80, Z84, Z85, Z86	6
	PGLNext (Ambry Genetics)			
BRCA1 and BRCA2 Gene Testing				
BRCA1 or BRCA2 Targeted Variant or Known Familial Variant Analysis	BRCA1 or BRCA2 Targeted Variant-Single Test (GeneDx)	81215, 81217	C50, C56, D05, Z17, Z80, Z83, Z84, Z85, Z86	1
BRCA1 and/or BRCA2 Targeted Variant Analysis - Ashkenazi Jewish Founder Variants	BRCA1/2 Ashkenazi Jewish 3-Site Mutation Panel (Ambry Genetics)	81212		
	MultiSite 3 BRCAanalysis (Myriad Genetics)			
BRCA1 and BRCA2 Sequencing and/or Deletion/Duplication Analysis	Hereditary BRCA1/2 Panel (Invitae)	81162, 81163, 81164, 81165,		
	BRCA1/2 Seq and Del/Dup (Ambry Genetics)	81166, 81167, 81216		
	+RNAinsight for BRCA1/2 (Ambry Genetics)	0138U		
PALB2 Gene Testing				
PALB2 Targeted Variant Analysis	PALB2 Targeted Variant (GeneDx)	81308	C15-26, Z80, Z84, Z85, Z86	1
PALB2 Sequencing and/or Deletion/Duplication Analysis	PALB2 Sequencing PALB2 Deletion/Duplication (Quest)	81307, 81479		
	PALB2 with +RNA insight (Ambry Genetics)	0137U		
ATM and/or CHEK2 Gene Testing				
ATM or CHEK2 Targeted Variant	ATM Targeted Variant - Single Test (GeneDx)	81479	C50, D05, Z80, Z84, Z85, Z86	1

Analysis	CHEK2 Targeted Variant - Single Test (GeneDx)			
ATM or CHEK2 Sequencing and/or Deletion/Duplication Analysis	Ataxia-Telangiectasia Test (Invitae)	81408, 81479		
	Hereditary Breast Cancer via the CHEK2 Gene (PreventionGenetics, part of Exact Sciences)	81479		
	+RNAinsight for ATM (Ambry Genetics)	0136U		
Lynch Syndrome / Hereditary Nonpolyposis Colorectal Cancer (HNPCC)				
MLH1, MSH2, MSH6, PMS2, or EPCAM Targeted Mutation Analysis	MSH6 Targeted Variant; PMS2 Targeted Variant; EPCAM Targeted Variant (GeneDx)	81299, 81318, 81479	C15-22, C24-6, C26 C53-57 Z80, Z84, Z85, Z86	2
	Hereditary Nonpolyposis Colorectal Cancer (HNPCC): MLH1 (Known Mutation) (Labcorp)	81293		
	Hereditary Nonpolyposis Colorectal Cancer (HNPCC): MSH2 (Known Mutation) (Labcorp)	81296		
MLH1, MSH2, MSH6, PMS2, or EPCAM Sequencing and/or Deletion/Duplication Analysis	HNPCC Concurrent (Ambry Genetics)	81292, 81294, 81295, 81297, 81298, 81300, 81317, 81319, 81403		
	Lynch Syndrome Panel (Invitae)			
	CustomNext + RNA: MLH1, MSH2, MSH6, and/or PMS2 (Ambry Genetics)	0158U, 0159U, 0160U, 0161U, 0162U		
BAP1-Tumor Predisposition Syndrome				
BAP1 Targeted Variant Analysis	BAP1: Site Specific Analysis (familial) (Univ of Pennsylvania School of Medicine-Genetic Diagnostic Laboratory)	81403	C22, C45, C64 C69, D22, D32, Z80, Z84, Z85, Z86	8
BAP1 Sequencing and/or Deletion/Duplication Analysis	BAP1 Full Gene Sequencing and Deletion/Duplication (Invitae)	81479		5, 8, 12 13, 14
Birt-Hogg-Dube syndrome (BHDS)				
FLCN Targeted Variant	FLCN Targeted Variant - Single	81479	C65, D14.3,	8

Analysis	Test (GeneDx)		D23.9, Z84, Z85, Z86	
FLCN Sequencing and/or Deletion/Duplication Analysis	Birt-Hogg-Dube Syndrome Test (Invitae)	81479		8, 10
<u>Cowden Syndrome (CS)/PTEN Hamartoma Tumor Syndrome (PHTS)</u>				
PTEN Targeted Variant Analysis	PTEN Targeted Variant - Single Test (GeneDx)	81322	C15-21, C26, C50, C54, C55, C64, C73, D12, D13, D17, D23, D24, F78, F84.0, Q75.3, Q87.89, Z80, Z84, Z85, Z86	1
PTEN Sequencing and/or Deletion/Duplication Analysis	PTEN Gene Sequencing and Del/Dup (GeneDx)	81321, 81323		
<u>Adenomatous Polyposis Conditions (Familial Adenomatous Polyposis syndrome (FAP)/Attenuated FAP (AFAP) and MUTYH-Associated Polyposis Syndrome (MAP))</u>				
APC or MUTYH Targeted Variant Analysis	APC Targeted Variant - Single Test (GeneDx)	81202	C15-21, D12, Z80, Z84, Z85, Z86	2
	MUTYH Targeted Variant - Single Test (GeneDx)	81403, 81401		
APC and/or MUTYH Sequencing and/or Deletion/Duplication Analysis	APC Seq and Del/Dup (Ambry Genetics)	81201, 81203		
	Familial Adenomatous Polyposis Test (Invitae)			
	+RNAInsight for APC (Ambry Genetics)	0157U		
	MUTYH Full Gene Sequencing and Deletion/Duplication (Invitae)	81406, 81479		
<u>Familial Atypical Multiple Mole Melanoma Syndrome (FAMMM)</u>				
CDKN2A Targeted Variant Analysis	CDKN2A Targeted Variant - Single Test (GeneDx)	81479	C43, Z12.83, Z80, Z84, Z85, Z86	19
CDKN2A Sequencing and/or Deletion/Duplication Analysis	CDKN2A Full Gene Sequencing and Deletion/Duplication (Invitae)	81404, 81479		5
<u>Hereditary Diffuse Gastric Cancer</u> (aka, Signet Ring Cell Gastric Cancer)				

CDHI Targeted Variant Analysis	CDH1 Targeted Variant - Single Test (GeneDx)	81479	C16, C50, Q35, Q36, Z80, Z84, Z85, Z86	1, 7
CDHI Sequencing and/or Deletion/Duplication Analysis	Hereditary Diffuse Gastric Cancer Syndrome Test (Invitae)	81406, 81479		7
Juvenile Polyposis Syndrome (JPS)				
SMAD4 and/or BMPR1A Targeted Variant Analysis	Targeted Variant: SMAD4 (PreventionGenetics, part of Exact Sciences)	81403	C15-C26, D12, Z80, Z84, Z85, Z86	2
	Targeted Variant: BMPR1A (PreventionGenetics, part of Exact Sciences)	81403		
SMAD4 and/or BMPR1A Sequencing and/or Deletion/Duplication Analysis	Juvenile Polyposis Syndrome Panel (Invitae)	81405, 81406, 81479		
	BMPR1A, SMAD4 Gene Sequencing and Del/Dup (GeneDx)			
Hereditary Leiomyomatosis and Renal Cell Cancer (HLRCC)				
FH Targeted Variant Analysis	FH Sequence Analysis (Familial Mutation/Variant Analysis) (Baylor Genetics)	81403	C44, C55, C64, D23, D25, Z84, Z85, Z86	8
FH Sequencing and/or Deletion/Duplication Analysis	Hereditary Leiomyomatosis and Renal Cell Carcinoma (Ambry Genetics)	81405, 81479		8, 18
Li-Fraumeni Syndrome (LFS)				
TP53 Targeted Variant Analysis	TP53 Targeted Variant - Single Test (GeneDx)	81352	C30-41, C15-26, C45, C47-49, C50, C71, C95.9, Z80, Z84, Z85, Z86	1
TP53 Sequencing and/or Deletion/Duplication Analysis	TP53 Full Gene Sequencing and Deletion/Duplication (Invitae)	81351, 81479		
	Li-Fraumeni Syndrome, TP53 Sequencing and Deletion/Duplication (Quest Diagnostics)			
Multiple Endocrine Neoplasia - Type 1 (MEN1)				
MEN1 Targeted Variant Analysis	MEN1 Targeted Variant - Single Test (GeneDx)	81479	C25, C75.0, D35.2, E31.2,	6

MEN1 Sequencing and/or Deletion/Duplication Analysis	MEN1 Gene Sequencing and Del/Dup (GeneDx)	81404, 81405	Z80, Z84, Z85, Z86	
	Multiple Endocrine Neoplasia Type 1 Test (Invitae)			
Multiple Endocrine Neoplasia Type 2 (MEN2)				
RET Targeted Variant Analysis	RET Targeted Variant - Single Test (GeneDx)	81404	C73-75, C7A, D3A, Z80, Z84, Z85, Z86	6
RET Sequencing and/or Deletion/Duplication Analysis	RET Full Gene Sequencing and Deletion/Duplication (Invitae)	81406, 81479, S3840		6, 17
Nevoid Basal Cell Carcinoma Syndrome (NBCCS) (aka Gorlin syndrome)				
PTCH1 and/or SUFU Targeted Variant Analysis	Targeted Variant: PTCH1 or SUFU (GeneDx)	81479	C44, C71.6, G93, M27.4, Z84, Z85, Z86	15
PTCH1 and SUFU Sequencing and/or Deletion/Duplication Analysis	Basal Cell Nevus Syndrome Panel (Invitae)	81479		
Hereditary Paraganglioma/Pheochromocytoma Syndrome (PGL/PCC)				
MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, or TMEM127 Targeted Variant Analysis	SDHB, SDHD, SDHC, MAX, SDHAF2, or TMEM127 Targeted Variant - Single Test (GeneDx) Targeted Variants: MAX, SDHAF2, TMEM127 (PreventionGenetics, part of Exact Sciences)	81479	C7A, C74.1, D35.00, D44.7, Z84, Z85, Z86	8
MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, and/or TMEM127 Sequencing and/or Deletion/Duplication Analysis	SHDB Full Gene Sequencing and Deletion/Duplication (Invitae)	81405, 81479		16
	SDHA Full Gene Sequencing and Deletion/Duplication (Invitae)	81406, 81479		
	SDHC Full Gene Sequencing and Deletion/Duplication (Invitae)	81404, 81405		
	SDHD Full Gene Sequencing and Deletion/Duplication (Invitae)	81404, 81479		
	MAX Full Gene Sequencing and Deletion/Duplication (Invitae)	81479		
	SDHAF2 Full Gene Sequencing and Deletion/Duplication (Invitae)			

	TMEM127 Full Gene Sequencing and Deletion/Duplication (Invitae)			
<u>Peutz-Jeghers Syndrome (PJS)</u>				
<u>STK11 Targeted Variant Analysis</u>	STK11 Targeted Variant - Single Test (GeneDx)	81479	C50, Q85.8, Z80, Z84, Z85, Z86	2
<u>STK11 Sequencing and/or Deletion/Duplication Analysis</u>	STK11 Gene Sequencing & Del/Dup (GeneDx)	81404, 81405		
<u>Retinoblastoma</u>				
<u>RB1 Targeted Variant Analysis</u>	Retinoblastoma: Site Specific Analysis (Familial) (Univ of Pennsylvania School of Medicine-Genetic Diagnostic Laboratory)	81403	C69, C75.3, Z80, Z84, Z85, Z86	9
<u>RB1 Sequencing and/or Deletion/Duplication Analysis</u>	RB1 Full Gene Sequencing and Deletion/Duplication (Invitae)	81479, S3841		
<u>Von Hippel-Lindau Syndrome (VHL)</u>				
<u>VHL Targeted Variant Analysis</u>	VHL Sequence Analysis (Familial Mutation/Variant Analysis) (Baylor Genetics, LLC)	81403	C64, C7A, D3A, D35.00, K86.2, N28, N50.3, Q85.8, Z80, Z84, Z85, Z86	8
<u>VHL Sequencing and/or Deletion/Duplication Analysis</u>	VHL Full Gene Sequencing and Deletion/Duplication (Invitae)	81403, 81404, S3842		
	VHL Gene Sequencing and Del/Dup (GeneDx)			

OTHER RELATED POLICIES

This policy document provides criteria for genetic testing for hereditary cancer susceptibility. Please refer to:

- ***Genetic Testing: Multisystem Inherited Disorders, Intellectual Disability, and Developmental Delay*** for criteria related to diagnostic testing for Fanconi anemia.

- ***Oncology: Algorithmic Testing*** for criteria related to tests that give prognostic information for an individual with cancer, or any oncology related test that involved an algorithmic portion.
- ***Oncology: Molecular Analysis of Solid Tumors and Hematologic Malignancies*** for criteria related to somatic tumor testing, including Microsatellite Instability for colon cancer, and blood cancer testing
- ***Oncology: Cancer Screening*** for criteria related to tests that screen for the presence of cancer.
- ***Oncology: Circulating Tumor DNA and Circulating Tumor Cells (Liquid Biopsy)*** for criteria related to the testing of tumor DNA circulating in an individual's blood stream.
- ***Genetic Testing: General Approach to Genetic and Molecular Testing*** for criteria related to hereditary cancer susceptibility that is not specifically discussed in this or other non-general policies.

CRITERIA

It is the policy of health plans affiliated with Centene Corporation® that the specific genetic testing noted below is **medically necessary** when meeting the related criteria:

PAN-CANCER HEREDITARY CANCER SUSCEPTIBILITY PANELS

A pan-cancer hereditary cancer susceptibility panel includes genes that are associated with inherited susceptibility to several different types of cancer (e.g., breast cancer, colon cancer, stomach cancer, etc.).

- I. Genetic testing using a pan-cancer hereditary cancer susceptibility panel (81432, 81433) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee meets at least one of the following:
 1. The member/enrollee meets clinical criteria for [BRCA1 and BRCA2 sequencing and/or deletion/duplication analysis](#), **OR**

2. The member/enrollee meets clinical criteria for [Lynch syndrome/HNPCC](#) [MLH1, MSH2, MSH6, PMS2, or EPCAM sequencing and/or deletion/duplication analysis](#), **AND**
 - C. The panel includes, at a minimum, sequencing of the following genes: *BRCA1*, *BRCA2*, *EPCAM*, *MLH1*, *MSH2*, *MSH6*, *PMS2*, **AND**
 - D. The panel does not include genes without a known association with cancer by [ClinGen](#).
- II. Genetic testing using a pan-cancer hereditary cancer susceptibility panel (81432, 81433) is considered **investigational** for all other indications.
- III. Hereditary cancer susceptibility panel targeted mRNA sequencing analysis for the interpretation of variants of unknown significance (0134U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance or follow up testing without proven utility.

Note: If a multigene cancer panel is performed, the appropriate panel code should be used.

[back to top](#)

HEREDITARY BREAST CANCER SUSCEPTIBILITY PANELS

A hereditary breast cancer susceptibility panel includes genes that are associated with inherited susceptibility to breast cancer.

- I. Genetic testing using a hereditary breast cancer susceptibility panel (81162, 81163, 81164, 81165, 81166, 81167, 81216, 81307, 81321, 81351, 81432, 81433, 0129U) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee meets sequencing and/or deletion/duplication clinical criteria for at least one of the following:
 1. [BRCA1 and BRCA2 Sequencing and Deletion/Duplication analysis](#), **OR**
 2. [PALB2 Sequencing or Deletion/Duplication analysis](#), **AND**
 - C. The panel includes, at a minimum, sequencing of the following genes: *BRCA1*, *BRCA2*, *PALB2*, **AND**

- D. The panel does not include genes without known association with breast cancer by [ClinGen](#).
- II. Genetic testing using a STAT hereditary breast cancer panel (81162, 81163, 81164, 81165, 81166, 81167, 81216) is considered medically necessary when:
 - A. The member/enrollee meets any of the above criteria, **AND**
 - B. The member/enrollee requires a rapid turn-around-time for decision making related to surgical interventions and treatment.
- III. Genetic testing using a hereditary breast cancer susceptibility panel (81162, 81163, 81164, 81165, 81166, 81167, 81216, 81307, 81321, 81351, 81432, 81433, 0129U) is considered **investigational** for all other indications.

[back to top](#)

HEREDITARY GI/COLON CANCER PANEL TESTS

A hereditary colorectal cancer susceptibility panel includes genes that are associated with inherited susceptibility to colorectal cancer.

- I. Genetic testing using a hereditary colorectal cancer susceptibility panel (81435, 81436, 0101U) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee meets at least one of the following:
 - 1. The member/enrollee has a personal history of, or at least one blood relative with any of the following:
 - a) At least 10 adenomatous polyps, **OR**
 - b) At least 2 hamartomatous polyps, **OR**
 - c) At least 5 serrated polyps/lesions proximal to the rectum, **OR**
 - 2. The member/enrollee has a personal history of colorectal cancer under 50 years of age, **OR**

3. The member/enrollee meets clinical criteria for Lynch syndrome/HNPCC [MLH1, MSH2, MSH6, PMS2, or EPCAM Sequencing and/or Deletion/Duplication Analysis](#), **AND**
 - C. The panel includes, at a minimum, sequencing of the following genes: *APC*, *MUTYH*, *MLH1*, *MSH2*, *MSH6*, *PMS2*, *EPCAM*, *BMPR1A*, *SMAD4*, *PTEN*, *STK11*, and *TP53*, **AND**
 - D. The panel does not include genes without a known association with colorectal or gastrointestinal cancer by [ClinGen](#).
- II. Genetic testing using a hereditary colorectal cancer susceptibility panel (81435, 81436, 0101U) is considered **investigational** for all other indications.
- III. Hereditary colorectal cancer susceptibility panel targeted mRNA sequencing analysis for the interpretation of variants of unknown significance (0130U, 0162U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance or follow up testing without proven utility.

Note: If a multigene cancer panel is performed, the appropriate panel code should be used.

[back to top](#)

HEREDITARY GASTRIC CANCER PANELS

A hereditary gastric cancer panel includes genes that are associated with inherited susceptibility to gastric (stomach) cancer.

- I. Genetic testing using a hereditary gastric susceptibility panel (81201, 81203, 81292, 81294, 81295, 81297, 81298, 81300, 81317, 81319, 81403, 81404, 81405, 81406, 81408, 81479) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee meets sequencing and/or deletion/duplication clinical criteria for at least one of the following:
 1. [Lynch syndrome/Hereditary Nonpolyposis Colorectal Cancer](#), **OR**
 2. [Hereditary Diffuse Gastric Cancer](#), **OR**
 3. [Juvenile Polyposis Syndrome](#), **OR**

- 4. [Peutz-Jeghers Syndrome](#), **OR**
 - 5. [Familial Adenomatous Polyposis](#), **AND**
 - C. The panel includes, at a minimum, sequencing of the following genes: *APC*, *BMPRIA*, *CDH1*, *EPCAM*, *MLH1*, *MSH2*, *MSH6*, *PMS2*, *SMAD4*, *STK11*, **AND**
 - D. The panel does not include genes without a known association with gastric (stomach) cancer by [ClinGen](#).
- II. Genetic testing using a hereditary gastric cancer susceptibility panel (81201, 81203, 81292, 81294, 81295, 81297, 81298, 81300, 81317, 81319, 81403, 81404, 81405, 81406, 81408, 81479) is considered **investigational** for all other indications.

[back to top](#)

HEREDITARY PANCREATIC CANCER SUSCEPTIBILITY PANELS

A hereditary pancreatic cancer susceptibility panel includes genes that are associated with inherited susceptibility to pancreatic cancer.

- I. Genetic testing using a hereditary pancreatic cancer susceptibility panel (81162, 81163, 81201, 81292, 81295, 81298, 81351, 81433, 81479) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee meets criteria for [BRCA1 and BRCA2 sequencing and/or deletion/duplication analysis](#), **AND**
 - C. The panel includes, at a minimum, sequencing of the following genes: *ATM*, *BRCA1*, *BRCA2*, *CDKN2A*, *EPCAM*, *MLH1*, *MSH2*, *MSH6*, *PALB2*, *STK11*, *TP53*, **AND**
 - D. The panel does not include genes without a known association with pancreatic cancer by [ClinGen](#).
- II. Genetic testing using a hereditary pancreatic cancer susceptibility panel (81162, 81163, 81201, 81292, 81295, 81298, 81351, 81433, 81479) is considered **investigational** for all other indications.

[back to top](#)

HEREDITARY POLYPOSIS PANELS

A hereditary polyposis panel is one that includes genes that are associated with inherited susceptibility to colon polyposis.

- I. Genetic testing using a hereditary polyposis panel (81201, 81203, 81406, 81479) is considered **medically necessary** when:
 - A. The member/enrollee meets criteria for sequencing and/or deletion/duplication analysis for [Adenomatous Polyposis conditions \(Familial Adenomatous Polyposis Syndrome \(FAP\)/Attenuated FAP \(AFAP\) and MUTYH-Associated Polyposis Syndrome \(MAP\)](#), **AND**
 - B. The panel includes, at a minimum, sequencing of the following genes: *APC* and *MUTYH*, **AND**
 - C. The panel does not include genes without a known association with colon polyposis by [ClinGen](#).
- II. Genetic testing using a hereditary polyposis panel (81201, 81203, 81406, 81479) is considered **investigational** for all other indications.

[back to top](#)

HEREDITARY PROSTATE CANCER SUSCEPTIBILITY PANELS

A hereditary prostate cancer susceptibility panel is one that includes genes that are associated with inherited susceptibility to prostate cancer.

- I. Genetic testing using a hereditary prostate cancer susceptibility panel (81162, 81292, 81295, 81351, 81479,) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The patient has a personal history of prostate cancer diagnosed at any age with any of the following characteristics:

1. Metastatic prostate cancer, **OR**
 2. [High- or very-high risk localized prostate cancer](#), **OR**
 3. Regional (node positive) prostate cancer, **OR**
- C. A personal history of prostate cancer and any of the following:
1. One or more [close relatives](#) with any of the following:
 - a) Breast cancer at or under age 50, **OR**
 - b) Colorectal or endometrial cancer at or under age 50, **OR**
 - c) Male (sex assigned at birth) breast cancer at any age, **OR**
 - d) Ovarian cancer at any age, **OR**
 - e) Exocrine pancreatic cancer at any age, **OR**
 - f) Metastatic, regional, very-high-risk, or high-risk prostate cancer at any age, **OR**
 2. One or more first degree relatives with prostate cancer at or under age 60, **OR**
 3. Two or more [close relatives](#) with either of the following:
 - a) Breast cancer at any age, **OR**
 - b) Prostate cancer at any age, **OR**
 4. Three or more first or second degree relatives with a [Lynch syndrome related cancer](#), especially if diagnosed under age 50, **OR**
 5. Ashkenazi Jewish ancestry, **OR**
 6. A personal history of breast cancer, **OR**
- D. The member/enrollee meets criteria for [BRCA1 and BRCA2 sequencing and/or deletion/duplication analysis](#), **AND**
- E. The panel includes, at a minimum, sequencing of the following genes: *BRCA1*, *BRCA2*, **AND**

- F. The panel does not include genes without a known association with prostate cancer by [ClinGen](#).
- II. Genetic testing using a hereditary prostate cancer susceptibility panel (81162, 81292, 81295, 81351, 81479, is considered **investigational** for all other indications.
- III. Hereditary prostate cancer susceptibility panel targeted mRNA sequencing analysis for the interpretation of variants of unknown significance (0133U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance, or follow up testing without proven utility.

[back to top](#)

HEREDITARY NEUROENDOCRINE CANCER SUSCEPTIBILITY PANELS

A hereditary neuroendocrine cancer susceptibility panel is one that includes genes that are associated with inherited susceptibility to a neuroendocrine cancer.

- I. Genetic testing using a hereditary neuroendocrine cancer susceptibility panel (81437, 81438) is considered **medically necessary** when:
 - A. The member/enrollee meets criteria for sequencing and/or deletion/duplication analysis for at least one of the following (see specific criteria sections below):
 - 1. [Von-Hippel Lindau syndrome \(VHL\)](#), **OR**
 - 2. [Hereditary Paraganglioma-Pheochromocytoma syndrome \(PGL/PCC\)](#), **OR**
 - 3. [Multiple Endocrine Neoplasia Type 1 \(MEN1\)](#), **OR**
 - 4. [Multiple Endocrine Neoplasia Type 2 \(MEN2\)](#), **AND**
 - B. The panel includes, at a minimum, sequencing of the following genes: *MAX*, *SDHB*, *SDHC*, *SDHD*, *TMEM127*, *SDHAF2*, *SDHA*, *VHL*, *MEN1*, *RET*, **AND**
 - C. The panel does not include genes without a known association with a neuroendocrine cancer by [ClinGen](#).

- II. Genetic testing using a hereditary neuroendocrine cancer susceptibility panel (81437, 81438) is considered **investigational** for all other indications.

Note: If a multigene cancer panel is performed, the appropriate panel code should be used.

[back to top](#)

***BRCA1* AND *BRCA2* GENE TESTING**

***BRCA1* or *BRCA2* Targeted Variant or Known Familial Variant Analysis**

- I. *BRCA1* (81215) or *BRCA2* (81217) targeted variant or known familial variant analysis for hereditary cancer susceptibility is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. One of the following:
 1. The member/enrollee has a family history of a known *BRCA1* or *BRCA2* pathogenic or likely pathogenic variant, **OR**
 2. A *BRCA1* or *BRCA2* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *BRCA1* (81215) or *BRCA2* (81217) targeted variant analysis for hereditary cancer susceptibility is considered **investigational** for all other indications.

***BRCA1* and/or *BRCA2* Targeted Variant Analysis - Ashkenazi Jewish Founder Variants**

- I. *BRCA1* and *BRCA2* (81212) targeted variant analysis for the 185delAG, 5385insC, 6174delT variants is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee is of Ashkenazi Jewish ancestry (at least one grandparent of Ashkenazi Jewish ancestry).
- II. *BRCA1* and *BRCA2* (81212) targeted variant analysis for the 185delAG, 5385insC, 6174delT variants is considered **investigational** for all other indications.

BRCA1 and BRCA2 Sequencing and/or Deletion/Duplication Analysis

- I. *BRCA1* and *BRCA2* (81162, 81163, 81164, 81165, 81166, 81167, 81216) sequencing and/or deletion/duplication analysis for hereditary breast and/or ovarian cancer susceptibility is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 1. The member/enrollee has a personal history of any of the following:
 - a) Male breast cancer, **OR**
 - b) Triple-negative breast cancer, **OR**
 - c) Epithelial ovarian cancer (including fallopian tube cancer or peritoneal cancer), **OR**
 - d) Pancreatic cancer, **OR**
 - e) Metastatic prostate cancer, **OR**
 - f) [High- or very-high-risk group prostate cancer](#), **OR**
 - g) Multiple primary breast cancers (diagnosed synchronously or metachronously), **OR**
 2. The member/enrollee has a personal history of breast cancer **AND** any of the following:
 - a) Female breast cancer diagnosed at age 50 years or younger, **OR**
 - b) Ashkenazi Jewish ancestry, **OR**
 - c) One or more [close relatives](#) with any of the following:
 - (1) Female breast cancer diagnosed at age 50 years or younger, **OR**
 - (2) Male breast cancer, **OR**
 - (3) Ovarian cancer, **OR**
 - (4) Pancreatic cancer, **OR**
 - (5) Metastatic, or [high- or very-high-risk group prostate cancer](#), **OR**

- d) Three or more total diagnoses of breast cancer in the member/enrollee and/or [close relatives](#), **OR**
 - e) Two or more [close relatives](#) with either breast or prostate cancer (of any grade), **OR**
- 3. The member/enrollee does not have a personal history of a BRCA1/2-related cancer, but has a [first- or second-degree relative](#) meeting any of the above criteria, **OR**
- 4. The member/enrollee has metastatic breast cancer and is being considered for systemic treatment using PARP inhibitors, **OR**
- 5. The member/enrollee has [high-risk](#), HER2-negative breast cancer and is being considered for adjuvant treatment with olaparib, **OR**
- 6. The member/enrollee's probability of having a *BRCA1* or *BRCA2* pathogenic variant is greater than 5% based on prior probability models (examples: Tyrer-Cuzick, BRCAPro, CanRisk).
- II. *BRCA1* and *BRCA2* (81162, 81163, 81164, 81165, 81166, 81167, 81216) sequencing and/or deletion/duplication analysis for hereditary breast and/or ovarian cancer susceptibility is considered **investigational** for all other indications.
- III. *BRCA1/BRCA2* mRNA sequencing analysis for the interpretation of variants of unknown significance (0138U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance or follow up testing without proven utility.

[back to top](#)

***PALB2* GENE TESTING**

***PALB2* Targeted Variant Analysis**

- I. *PALB2* targeted variant analysis (81308) for hereditary breast and/or ovarian cancer susceptibility is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. One of the following:

1. The member/enrollee has a family history of a known pathogenic or likely pathogenic variant in *PALB2*, **OR**
 2. A pathogenic or likely pathogenic variant was detected by tumor profiling in *PALB2*, and germline analysis has not yet been performed.
- II. *PALB2* targeted variant analysis (81308) for hereditary breast and/or ovarian cancer susceptibility is considered **investigational** for all other indications.

***PALB2* Sequencing and/or Deletion/Duplication Analysis**

- I. *PALB2* (81307, 81479) sequencing and/or deletion/duplication analysis for hereditary breast and/or ovarian cancer susceptibility is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 1. The member/enrollee has a personal history of any of the following:
 - a) Male breast cancer, **OR**
 - b) Triple-negative breast cancer, **OR**
 - c) Epithelial ovarian cancer (including fallopian tube cancer or peritoneal cancer), **OR**
 - d) Pancreatic cancer, **OR**
 - e) Multiple primary breast cancers (diagnosed synchronously or metachronously), **OR**
 2. The member/enrollee has a personal history of breast cancer **AND** any of the following:
 - a) Female breast cancer diagnosed at age 50 years or younger, **OR**
 - b) Ashkenazi Jewish ancestry, **OR**
 - c) One or more [close relatives](#) with any of the following:
 - (1) Female breast cancer diagnosed at age 50 years or younger, **OR**
 - (2) Male breast cancer, **OR**
 - (3) Ovarian cancer, **OR**

(4) Pancreatic cancer, **OR**

d) Three or more total diagnoses of breast cancer in the member/enrollee and/or close relatives, **OR**

3. The member/enrollee does not have a personal history of a PALB2-related cancer, but has a [first- or second-degree relative](#) meeting any of the above criteria, **OR**
 4. The member/enrollee has breast cancer and is being considered for systemic treatment decisions using PARP inhibitors for metastatic breast cancer, **OR**
 5. The member/enrollee has [high-risk](#), HER2-negative breast cancer and is being considered for adjuvant treatment with olaparib, **OR**
 6. The member/enrollee's probability of having a *BRCA1* or *BRCA2* pathogenic variant is greater than 5% based on prior probability models (examples: Tyrer-Curzick, BRCAPro, CanRisk).
- II. *PALB2* (81307) sequencing and/or deletion/duplication analysis for hereditary breast and/or ovarian cancer susceptibility is considered **investigational** for all other indications.
 - III. *PALB2* mRNA sequencing analysis for the interpretation of variants of unknown significance (0137U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance or follow up testing without proven utility.

[back to top](#)

***ATM* AND/OR *CHEK2* GENE TESTING**

***ATM* or *CHEK2* Targeted Variant Analysis**

- I. *ATM* (81479) or *CHEK2* (81479) targeted variant analysis for hereditary breast and/or ovarian cancer susceptibility is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. One of the following:
 1. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *ATM* or *CHEK2*, **OR**

2. A pathogenic or likely pathogenic variant was detected by tumor profiling in *ATM* or *CHEK2* and germline analysis has not yet been performed.
- II. *ATM* (81479) or *CHEK2* (81479) targeted variant analysis for hereditary breast and/or ovarian cancer susceptibility is considered **investigational** for all other indications.

***ATM* and/or *CHEK2* Sequencing and/or Deletion/Duplication Analysis**

- I. *ATM* (81408, 81479) and/or *CHEK2* (81479) sequencing and/or deletion/duplication analysis for hereditary breast and/or ovarian cancer susceptibility, as a stand alone test, is considered **investigational**.
- II. *ATM* mRNA sequencing analysis for the interpretation of variants of unknown significance (0136U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance or follow up testing without proven utility.

[back to top](#)

LYNCH SYNDROME / HEREDITARY NONPOLYPOSIS COLORECTAL CANCER (HNPCC) TESTING

***MLH1*, *MSH2*, *MSH6*, *PMS2*, or *EPCAM* Targeted Variant Analysis**

- I. *MLH1* (81293), *MSH2* (81296), *MSH6* (81299), *PMS2* (81318), or *EPCAM* (81479) targeted variant analysis for Lynch syndrome/HNPCC is considered **medically necessary** when:
 - A. The member/enrollee has a blood relative with a known pathogenic or likely pathogenic variant in *MLH1*, *MSH2*, *MSH6*, *PMS2*, or *EPCAM*, **OR**
 - B. A pathogenic or likely pathogenic variant was detected by tumor profiling in *MLH1*, *MSH2*, *MSH6*, *PMS2*, or *EPCAM* and germline analysis has not yet been performed.
- II. *MLH1* (81293), *MSH2* (81296), *MSH6* (81299), *PMS2* (81318), or *EPCAM* (81479) targeted variant analysis for Lynch syndrome/HNPCC is considered **investigational** for all other indications.

***MLH1, MSH2, MSH6, PMS2, or EPCAM* Sequencing and/or Deletion/Duplication Analysis**

- I. *MLH1* (81292, 81294), *MSH2* (81295, 81297), *MSH6* (81298, 81300), *PMS2* (81317, 81319), and/or *EPCAM* (81403) sequencing and/or duplication analysis for Lynch syndrome/HNPCC is considered **medically necessary** when:
 - A. The member/enrollee has a [Lynch syndrome-related cancer](#) **and** the tumor shows evidence of mismatch repair (MMR) deficiency (either by microsatellite instability (MSI) or loss of MMR protein expression), **OR**
 - B. The member/enrollee has a diagnosis of a [Lynch syndrome-related cancer](#), **AND** any of the following:
 1. Diagnosed before age 50, **OR**
 2. Diagnosed at any age with an additional [Lynch syndrome-related cancer](#) **OR**
 3. Diagnosed at any age with one or more [first- or second-degree relatives](#) diagnosed before age 50 with a [Lynch syndrome-related cancer](#), **OR**
 4. Diagnosed at any age with two or more [first- or second-degree relatives](#) diagnosed at any age with a [Lynch syndrome-related cancer](#), **OR**
 - C. The member/enrollee has a family history of **any** of the following:
 1. One or more [first-degree relatives](#) diagnosed with colorectal or endometrial cancer before age 50, **OR**
 2. One or more [first-degree relatives](#) diagnosed with colorectal or endometrial cancer and an additional [Lynch syndrome-related cancer](#), **OR**
 3. Two or more [first- or second-degree relatives](#) diagnosed with a [Lynch syndrome-related cancer](#), one of whom was diagnosed before age 50, **OR**
 4. Three or more [first- or second-degree relatives](#) diagnosed with a [Lynch syndrome-related cancer](#), **OR**
 - D. The member/enrollee has a 5% or greater risk of Lynch syndrome on one of the following variant prediction models: MMRpro, PREMM5, MMRpredict, **OR**
 - E. The member/enrollee has a personal history of colorectal and/or endometrial cancer with a PREMM5 score of 2.5% or greater.

- II. *MLH1* (81292, 81294), *MSH2* (81295, 81297), *MSH6* (81298, 81300), *PMS2* (81317, 81319), and/or *EPCAM* (81403) sequencing and/or duplication analysis for Lynch syndrome/HNPCC is considered **investigational** for all other indications.
- III. *MLH1*, *MSH2*, *MSH6*, *PMS2* and *EPCAM* mRNA sequencing analysis for the interpretation of variants of unknown significance (0158U, 0159U, 0160U, 0161U, 0162U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance, or follow up testing without proven utility.

[back to top](#)

***BAP1*-TUMOR PREDISPOSITION SYNDROME**

***BAP1* Targeted Variant Analysis**

- I. *BAP1* targeted variant analysis (81403) for *BAP1*-tumor predisposition syndrome is considered **medically necessary** when:
 - A. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *BAP1*, **OR**
 - B. A pathogenic or likely pathogenic variant in *BAP1* was identified on tumor profiling and germline analysis has not yet been performed.
- II. *BAP1* targeted variant analysis (81403) for *BAP1*-tumor predisposition syndrome is considered **investigational** for all other indications.

***BAP1* Sequencing and/or Deletion/Duplication Analysis**

- I. *BAP1* sequencing and/or deletion/duplication analysis (81479) for *BAP1*-tumor predisposition syndrome is considered **medically necessary** when:
 - A. The member/enrollee has a personal history of:
 - 1. Two or more of the following:
 - a) *BAP1*-inactivated melanocytic tumors (aka atypical spitz tumor), **OR**
 - b) Uveal melanoma, **OR**

- c) Malignant mesothelioma, **OR**
- d) Renal cell carcinoma, **OR**
- e) Hepatocellular carcinoma, **OR**
- f) Cholangiocarcinoma, **OR**
- g) Meningioma, **OR**
- 2. One or more of the above listed tumors/cancer, **AND**
 - a) A [first- or second-degree relative](#) with any of the above listed tumors/cancers.
- II. *BAP1* sequencing and/or deletion/duplication analysis (81479) for *BAP1*-tumor predisposition syndrome is considered **investigational** for all other indications.

[back to top](#)

BIRT-HOGG-DUBE SYNDROME (BHDS)

***FLCN* Targeted Variant Analysis**

- I. *FLCN* targeted variant analysis (81479) for Birt-Hogg-Dube syndrome (BHDS) is considered **medically necessary** when:
 - A. The member/enrollee has a [first- or second-degree relative](#) with a known pathogenic or likely pathogenic variant in *FLCN*, **OR**
 - B. A pathogenic or likely pathogenic variant in *FLCN* was identified on tumor profiling and germline analysis has not yet been performed.
- II. *FLCN* targeted variant analysis (81479) for Birt-Hogg-Dube syndrome (BHDS) is considered **investigational** for all other indications.

***FLCN* Sequencing and/or Deletion/Duplication Analysis**

- I. *FLCN* sequencing and/or deletion/duplication analysis (81479) for Birt-Hogg-Dube syndrome (BHDS) is considered **medically necessary** when:
 - A. The member/enrollee has a personal history of any of the following:

1. 5 or more fibrofolliculomas/trichodiscomas with at least one confirmed histologically, **OR**
 2. Multiple lung cysts with no apparent cause, **OR**
 3. Renal cancer diagnosed before 50 years of age, **OR**
 4. Multifocal or bilateral renal cancer, **OR**
 5. Renal cancer of mixed chromophobe and oncocytic, clear cell, or papillary histology, **OR**
 6. Oncocytoma, **OR**
 7. Angiomyolipoma, **OR**
 8. A [first-degree relative](#) with BHDS.
- II. *FLCN* sequencing and/or deletion/duplication analysis (81479) for Birt-Hogg-Dube syndrome (BHDS) is considered **investigational** for all other indications.

[back to top](#)

COWDEN SYNDROME (CS)/PTEN HAMARTOMA TUMOR SYNDROME (PHTS)

***PTEN* Targeted Variant Analysis**

- I. *PTEN* targeted variant analysis (81322) for Cowden syndrome (CS)/*PTEN* hamartoma tumor syndrome (PHTS) is considered **medically necessary** when:
 - A. The member/enrollee has a blood relative with a known pathogenic or likely pathogenic variant in *PTEN*, **OR**
 - B. A pathogenic or likely pathogenic variant in *PTEN* was identified on tumor profiling and germline analysis has not yet been performed.
- II. *PTEN* targeted variant analysis (81322) for Cowden syndrome (CS)/*PTEN* hamartoma tumor syndrome (PHTS) is considered **investigational** for all other indications.

***PTEN* Sequencing and/or Deletion/Duplication Analysis**

- I. *PTEN* sequencing and/or deletion/duplication analysis (81321, 81323) for Cowden syndrome (CS)/*PTEN* hamartoma tumor syndrome (PHTS) is considered **medically necessary** when:
 - A. The member/enrollee has a personal history of any of the following:
 1. Bannayan Riley-Ruvalcaba syndrome (BRRS), **OR**
 2. Adult Lhermitte-Duclos disease (LDD) (defined as the presence of a cerebellar dysplastic gangliocytoma), **OR**
 3. Autism-spectrum disorder and macrocephaly, **OR**
 4. At least 2 biopsy-proven trichilemmomas, **OR**
 - B. The member/enrollee meets clinical criteria for CS/PHTS:
 1. Macrocephaly (greater than or equal to 97 percentile), **OR**
 2. Lhermitte-Duclos disease, **OR**
 3. Gastrointestinal hamartomas or ganglioneuromas, **AND**
 4. At least two of the following:
 - a) Breast Cancer, **OR**
 - b) Endometrial Cancer, **OR**
 - c) Thyroid Cancer (follicular), **OR**
 - d) Macular pigmentation of the glans penis, **OR**
 - e) Mucocutaneous lesions (One biopsy-proven trichilemmoma; multiple palmoplantar keratoses; multifocal or extensive oral mucosal papillomatosis; multiple cutaneous facial papules), **OR**
 - C. The member/enrollee has at least two of the following:
 1. Breast Cancer, **OR**
 2. Endometrial Cancer, **OR**
 3. Thyroid Cancer (follicular), **OR**

4. Multiple gastrointestinal hamartomas or ganglioneuromas, **OR**
5. Macrocephaly (greater than or equal to 97 percentile), **OR**
6. Macular pigmentation of the glans penis, **OR**
7. Mucocutaneous lesions (One biopsy-proven trichilemmoma; multiple palmoplantar keratoses; multifocal or extensive oral mucosal papillomatosis; multiple cutaneous facial papules), **AND**
8. At least three of the following:
 - a) Autism Spectrum Disorder, **OR**
 - b) Colon Cancer, **OR**
 - c) Esophageal glycogenic acanthosis (3 or more), **OR**
 - d) Lipomas, **OR**
 - e) Intellectual disability (i.e., IQ less than or equal to 75), **OR**
 - f) Thyroid cancer (papillary or follicular variant of papillary thyroid cancer), **OR**
 - g) Thyroid structural lesions (such as adenoma, multinodular goiter), **OR**
 - h) Renal cell carcinoma, **OR**
 - i) Single GI hamartoma or ganglioneuroma, **OR**
 - j) Testicular lipomatosis, **OR**
 - k) Vascular anomalies (including multiple intracranial developmental venous anomalies), **OR**

D. The member/enrollee has macrocephaly, AND

1. Breast Cancer, **OR**
2. Endometrial Cancer, **OR**
3. Thyroid Cancer (follicular), **OR**
4. Multiple gastrointestinal hamartomas or ganglioneuromas, **OR**

5. Macrocephaly (greater than or equal to 97 percentile), **OR**
6. Macular pigmentation of the glans penis, **OR**
7. Mucocutaneous lesions (One biopsy-proven trichilemmoma; multiple palmoplantar keratoses; multifocal or extensive oral mucosal papillomatosis; multiple cutaneous facial papules), **OR**

E. The member/enrollee has at least three of the following:

1. Breast Cancer, **OR**
2. Endometrial Cancer, **OR**
3. Thyroid Cancer (follicular), **OR**
4. Multiple gastrointestinal hamartomas or ganglioneuromas, **OR**
5. Macular pigmentation of the glans penis, **OR**
6. Mucocutaneous lesions (One biopsy-proven trichilemmoma; multiple palmoplantar keratoses; multifocal or extensive oral mucosal papillomatosis; multiple cutaneous facial papules), **OR**
7. The member/enrollee has a [close relative](#) with a clinical diagnosis of CS/PHTS or BRRS for whom testing has not been performed, **OR**

F. The member/enrollee has any of the following:

1. Breast Cancer, **OR**
2. Endometrial Cancer, **OR**
3. Thyroid Cancer (follicular), **OR**
4. Multiple gastrointestinal hamartomas or ganglioneuromas, **OR**
5. Macrocephaly (greater than or equal to 97 percentile), **OR**
6. Macular pigmentation of the glans penis, **OR**
7. Mucocutaneous lesions (One biopsy-proven trichilemmoma; multiple palmoplantar keratoses; multifocal or extensive oral mucosal papillomatosis; multiple cutaneous facial papules), **AND**
8. At least three of the following:

- a) Autism Spectrum Disorder, **OR**
- b) Colon Cancer, **OR**
- c) Esophageal glycogenic acanthosis (3 or more), **OR**
- d) Lipomas, **OR**
- e) Intellectual disability (ie, IQ less than or equal to 75), **OR**
- f) Thyroid cancer (papillary or follicular variant of papillary thyroid cancer), **OR**
- g) Thyroid structural lesions (such as adenoma, multinodular goiter), **OR**
- h) Renal cell carcinoma, **OR**
- i) Single GI hamartoma or ganglioneuroma, **OR**
- j) Testicular lipomatosis, **OR**
- k) Vascular anomalies (including multiple intracranial developmental venous anomalies), **OR**

G. The member/enrollee has at least four of the following:

- 1. Autism Spectrum Disorder, **OR**
- 2. Colon Cancer, **OR**
- 3. Esophageal glycogenic acanthosis (3 or more), **OR**
- 4. Lipomas, **OR**
- 5. Intellectual disability (i.e., IQ less than or equal to 75), **OR**
- 6. Thyroid cancer (papillary or follicular variant of papillary thyroid cancer), **OR**
- 7. Thyroid structural lesions (such as adenoma, multinodular goiter), **OR**
- 8. Renal cell carcinoma, **OR**
- 9. Single GI hamartoma or ganglioneuroma, **OR**

- 10. Testicular lipomatosis, **OR**
- 11. Vascular anomalies (including multiple intracranial developmental venous anomalies), **OR**
- H. The member/enrollee has a close relative with a clinical diagnosis of CS/PHTS or BRRS for whom testing has not been performed, **AND**
 - 1. The member/enrollee has at least one of the following:
 - a) Breast Cancer, **OR**
 - b) Endometrial Cancer, **OR**
 - c) Thyroid Cancer (follicular), **OR**
 - d) Multiple gastrointestinal hamartomas or ganglioneuromas, **OR**
 - e) Macrocephaly (greater than or equal to 97 percentile), **OR**
 - f) Macular pigmentation of the glans penis, **OR**
 - g) Mucocutaneous lesions (One biopsy-proven trichilemmoma; multiple palmoplantar keratoses; multifocal or extensive oral mucosal papillomatosis; multiple cutaneous facial papules), **OR**
 - 2. At least two of the following:
 - a) Autism Spectrum Disorder, **OR**
 - b) Colon Cancer, **OR**
 - c) Esophageal glycogenic acanthosis (3 or more), **OR**
 - d) Lipomas, **OR**
 - e) Intellectual disability (i.e., IQ less than or equal to 75), **OR**
 - f) Thyroid cancer (papillary or follicular variant of papillary thyroid cancer), **OR**
 - g) Thyroid structural lesions (such as adenoma, multinodular goiter), **OR**
 - h) Renal cell carcinoma, **OR**

- i) Single GI hamartoma or ganglioneuroma, **OR**
 - j) Testicular lipomatosis, **OR**
 - k) Vascular anomalies (including multiple intracranial developmental venous anomalies).
- II. *PTEN* sequencing and/or deletion/duplication analysis (81321, 81323,) for Cowden syndrome (CS)/*PTEN* hamartoma tumor syndrome (PHTS) is considered **investigational** for all other indications.

[back to top](#)

ADENOMATOUS POLYPOSIS CONDITIONS (Familial Adenomatous Polyposis Syndrome (FAP)/Attenuated FAP (AFAP) AND *MUTYH*-Associated Polyposis Syndrome (MAP)

***APC* OR *MUTYH* Targeted Variant Analysis**

- I. *APC* (81202) or *MUTYH* targeted variant analysis (81401, 81403) for [adenomatous polyposis](#) testing is considered **medically necessary** when:
 - A. The member/enrollee has a family history of a known pathogenic or likely pathogenic variant in *APC* or *MUTYH*, **OR**
 - B. An *APC* or *MUTYH* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *APC* (81202) or *MUTYH* (81401, 81403) targeted variant analysis for adenomatous polyposis conditions is considered **investigational** for all other indications.

***APC* and/or *MUTYH* Sequencing and/or Deletion/Duplication Analysis**

- I. *APC* sequencing and/or deletion/duplication analysis (81201, 81203) and/or *MUTYH* sequencing and/or deletion/duplication analysis (81406, 81479) for adenomatous polyposis conditions is considered **medically necessary** when:
 - A. The member/enrollee has a history of any of the following:
 - 1. 10 or more cumulative adenomas, **OR**
 - 2. Congenital hypertrophy of the retinal pigment epithelium (CHRPE), **OR**

3. Desmoid tumor, **OR**
 4. Hepatoblastoma, **OR**
 5. Cribriform-morular variant of papillary thyroid cancer, **OR**
 6. A clinical diagnosis of serrated-polyposis syndrome, with at least some adenomas, based on one of the following:
 - a) 5 or more serrated polyps proximal to the rectum, all being 5mm or greater in size and at least 2 being 10mm or greater in size, **OR**
 - b) More than 20 serrated polyps of any size distributed throughout the large bowel, with at 5 or more being proximal to the rectum, **OR**
 7. Duodenal cancer, **OR**
 8. Duodenal adenomas.
- II. *APC* sequencing and/or deletion/duplication analysis (81201, 81203) and/or *MUTYH* sequencing and/or deletion/duplication analysis (81406, 81479) for adenomatous polyposis conditions is considered **investigational** for all other indications.
 - III. *APC* mRNA sequencing analysis for the interpretation of variants of unknown significance (0157U), when billed in addition, is considered **investigational** because it is typically either considered an existing component of the genetic testing process for quality assurance or follow up testing without proven utility.

[back to top](#)

FAMILIAL ATYPICAL MULTIPLE MOLE MELANOMA (FAMMM) SYNDROME

CDKN2A Targeted Variant Analysis

- I. *CDKN2A* targeted variant analysis (81479) for familial atypical multiple mole melanoma (FAMMM) syndrome, also known as melanoma-pancreatic cancer syndrome, is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. One of the following:

1. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *CDKN2A*, **OR**
 2. A *CDKN2A* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *CDKN2A* targeted variant analysis (81479) for familial cutaneous malignant melanoma is considered **investigational** for all other indications.

***CDKN2A* Sequencing and/or Deletion/Duplication Analysis**

- I. *CDKN2A* sequencing and/or deletion/duplication analysis (81404, 81479) for familial atypical multiple mole melanoma (FAMMM) syndrome, also known as melanoma-pancreatic cancer syndrome, as a standalone test, is considered **investigational**.

[back to top](#)

HEREDITARY DIFFUSE GASTRIC CANCER (AKA, SIGNET RING CELL GASTRIC CANCER):

***CDH1* Targeted Variant Analysis**

- I. *CDH1* targeted variant analysis (81479) for Hereditary Diffuse Gastric Cancer (aka, Signet Ring Cell Gastric Cancer) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. One of the following:
 1. The member/enrollee has a blood relative with a known pathogenic or likely pathogenic variant in *CDH1*, **OR**
 2. A *CDH1* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *CDH1* targeted variant analysis (81479) for Hereditary Diffuse Gastric Cancer (aka, Signet Ring Cell Gastric Cancer) is considered **investigational** for all other indications.

***CDH1* Sequencing and/or Deletion/Duplication Analysis**

- I. *CDH1* sequencing and/or deletion/duplication analysis for Hereditary Diffuse Gastric Cancer (aka, Signet Ring Cell Gastric Cancer) (81406, 81479) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee meets at least one of the following criteria:
 1. Diffuse gastric cancer diagnosed before age 50 years, **OR**
 2. Diffuse gastric cancer diagnosed at any age in a member/enrollee with [Maori ancestry](#), **OR**
 3. Diffuse gastric cancer diagnosed at any age in a member/enrollee with a personal or family history of cleft lip/cleft palate, **OR**
 4. Bilateral lobular breast cancer diagnosed before age 70 years, **OR**
 5. Personal or family history of diffuse gastric cancer and lobular breast cancer, one diagnosed before age 70 years, **OR**
 6. Two cases of gastric cancer in the family, one of which is a confirmed case of diffuse gastric cancer, diagnosed at any age, **OR**
 7. Two cases of lobular breast cancer in family member before 50 years of age.
- II. *CDH1* sequencing and/or deletion/duplication analysis for Hereditary Diffuse Gastric Cancer (aka, Signet Ring Cell Gastric Cancer) (81406, 81479) is considered **investigational** for all other indications.

[back to top](#)

JUVENILE POLYPOSIS SYNDROME (JPS)

***SMAD4* or *BMPRI1A* Targeted Variant Analysis**

- I. *SMAD4* and/or *BMPRI1A* targeted variant analysis (81403) for juvenile polyposis syndrome (JPS) is considered **medically necessary** when:

- A. The member/enrollee has a blood relative with a known pathogenic or likely pathogenic variant in *SMAD4* and/or *BMPRIA*, **OR**
 - B. A *SMAD4* and/or *BMPRIA* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *SMAD4* and/or *BMPRIA* targeted variant analysis (81403) for juvenile polyposis syndrome (JPS) is considered **investigational** for all other indications.

***SMAD4* and/or *BMPRIA* Sequencing and/or Deletion/Duplication Analysis**

- I. *SMAD4* and/or *BMPRIA* sequencing and/or deletion/duplication analysis (81405, 81406, 81479) for juvenile polyposis syndrome (JPS) is considered **medically necessary** when:
- A. The member/enrollee has 5 or more [juvenile polyps](#) in the colon, **OR**
 - B. The member/enrollee has multiple [juvenile polyps](#) throughout the gastrointestinal tract, **OR**
 - C. The member/enrollee has [juvenile polyps](#) (any number) and a family history of JPS, **OR**
 - D. The member/enrollee has a personal history of cancer and a *SMAD4* or *BMPRIA* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *SMAD4* and/or *BMPRIA* sequencing and/or deletion/duplication analysis (81405, 81406, 81479) for juvenile polyposis syndrome (JPS) is considered **investigational** for all other indications.

[back to top](#)

HEREDITARY LEIOMYOMATOSIS AND RENAL CELL CANCER (HLRCC)

***FH* Targeted Variant Analysis**

- I. *FH* targeted variant analysis (81403) for hereditary leiomyomatosis and renal cell cancer (HLRCC) is considered **medically necessary** when:
- A. The member/enrollee is 18 years or older, **AND**

B. One of the following:

1. The member/enrollee has a [first- or second-degree relative](#) with a known pathogenic or likely pathogenic variant in *FH*, **OR**
 2. A *FH* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *FH* targeted variant analysis (81403) for hereditary leiomyomatosis and renal cell cancer (HLRCC) is considered **investigational** for all other indications.

***FH* Sequencing and/or Deletion/Duplication Analysis**

- I. *FH* sequencing and/or deletion/duplication analysis (81405, 81479) for hereditary leiomyomatosis and renal cell cancer (HLRCC) is considered **medically necessary** when:
 - A. The member/enrollee is 18 years or older, **AND**
 - B. The member/enrollee has at least one of the following:
 1. Cutaneous leiomyomata, **OR**
 2. Uterine leiomyomata (uterine fibroids), **OR**
 3. Renal cell carcinoma.
- II. *FH* sequencing and/or deletion/duplication analysis (81405, 81479) for hereditary leiomyomatosis and renal cell cancer (HLRCC) is considered **investigational** for all other indications.

[back to top](#)

LI-FRAUMENI SYNDROME (LFS)

***TP53* Targeted Variant Analysis**

- I. *TP53* targeted variant analysis (81352) for Li-Fraumeni syndrome (LFS) is considered **medically necessary** when:
 - A. The member/enrollee has a blood relative with a known pathogenic or likely pathogenic variant in *TP53*, **OR**

- B. A *TP53* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *TP53* targeted variant analysis (81352) for Li-Fraumeni syndrome (LFS) is considered **investigational** for all other indications.

***TP53* Sequencing and/or Deletion/Duplication Analysis**

- I. *TP53* sequencing and/or deletion/duplication analysis (81351, 81479) for Li-Fraumeni syndrome (LFS) is considered **medically necessary** when:
 - A. The member/enrollee was diagnosed with breast cancer before 31 years of age, **OR**
 - B. The member/enrollee was diagnosed with pediatric hypodiploid acute lymphoblastic leukemia, **OR**
 - C. The member/enrollee meets **all** of the following Classic LFS criteria:
 - 1. The member/enrollee was diagnosed with a sarcoma before 45 years of age, **AND**
 - 2. The member/enrollee has a [first-degree relative](#) diagnosed with any cancer before 45 years of age, **AND**
 - 3. At least one of the following:
 - a) The member/enrollee has an additional [first- or second-degree relative](#) diagnosed with any cancer before 45 years of age, **OR**
 - b) The member/enrollee has an additional [first- or second-degree relative](#) diagnosed with sarcoma at any age, **OR**
 - D. The member/enrollee meets **any** of the following Chompret clinical diagnostic criteria:
 - 1. The member/enrollee has been diagnosed at any age with an adrenocortical carcinoma, choroid plexus carcinoma, or rhabdomyosarcoma of embryonal anaplastic subtype, **OR**
 - 2. The member/enrollee has multiple tumors (except multiple breast tumors), two of which belong to the LFS tumor spectrum (soft tissue sarcoma, osteosarcoma, CNS tumor, breast cancer) with the initial cancer occurring before 46 years of age, **OR**

3. The member/enrollee has a diagnosis of soft tissue sarcoma, osteosarcoma, CNS tumor, breast cancer diagnosed before 46 years of age, **AND**
 - a) A [first- or second-degree relative](#) diagnosed with any of the aforementioned cancers (other than breast cancer if the proband has breast cancer) before 56 years of age, **OR**
 - b) Multiple primaries at any age.
- II. *TP53* sequencing and/or deletion/duplication analysis (81351, 81479) for Li-Fraumeni syndrome (LFS) is considered **investigational** for all other indications.

[back to top](#)

MULTIPLE ENDOCRINE NEOPLASIA TYPE 1 (MEN1)

***MEN1* Targeted Variant Analysis**

- I. *MEN1* targeted variant analysis (81479) for multiple endocrine neoplasia type 1 (MEN1) is considered **medically necessary** when:
 - A. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *MEN1*, **OR**
 - B. An *MEN1* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *MEN1* targeted variant analysis (81479) for multiple endocrine neoplasia type 1 (MEN1) is considered **investigational** for all other indications.

***MEN1* Sequencing and/or Deletion/Duplication Analysis**

- I. *MEN1* sequencing and/or deletion/duplication analysis (81404, 81405) for multiple endocrine neoplasia type 1 (MEN1) is considered **medically necessary** when:
 - A. The member/enrollee has a personal history of at least two of the following:
 1. Duodenal/pancreatic neuroendocrine tumor, **OR**
 2. Primary hyperparathyroidism, **OR**
 3. Pituitary adenoma, **OR**

- 4. Foregut (bronchial, thymic, or gastric) carcinoid, **OR**
- B. The member/enrollee has a personal history of one of the above, **AND**
 - 1. The member/enrollee has a family history of at least one of the above.
- II. *MEN1* sequencing and/or deletion/duplication analysis (81404, 81405) for multiple endocrine neoplasia type 1 (MEN1) is considered **investigational** for all other indications.

[back to top](#)

MULTIPLE ENDOCRINE NEOPLASIA TYPE 2 (MEN2)

***RET* Targeted Variant Analysis**

- I. *RET* targeted variant analysis (81404) for multiple endocrine neoplasia type 2 (MEN2) is considered **medically necessary** when:
 - A. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *RET*, **OR**
 - B. A *RET* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *RET* targeted variant analysis (81404) for multiple endocrine neoplasia type 2 (MEN2) is considered **investigational** for all other indications.

***RET* Sequencing and/or Deletion/Duplication Analysis**

- I. *RET* sequencing and/or deletion/duplication analysis (81406, 81479, S3840) for multiple endocrine neoplasia type 2 (MEN2) is considered **medically necessary** when:
 - A. The member/enrollee has a diagnosis of medullary thyroid cancer, **OR**
 - B. The member/enrollee has a diagnosis of primary C-cell hyperplasia, **OR**
 - C. The member/enrollee has a personal history of an adrenal pheochromocytoma and parathyroid hyperplasia, **OR**
 - D. The member/enrollee has a [first-degree relative](#) that meets at least one of the above criteria and has not previously undergone *RET* sequencing and/or deletion duplication analysis.

- II. *RET* sequencing and/or deletion/duplication analysis (81406, 81479, S3840) for multiple endocrine neoplasia type 2 (MEN2) is considered **investigational** for all other indications.

[back to top](#)

NEVOID BASAL CELL CARCINOMA SYNDROME (NBCCS) (aka Gorlin syndrome)

***PTCH1* or *SUFU* Targeted Variant Analysis**

- I. *PTCH1* or *SUFU* targeted variant analysis (81479) for nevoid basal cell carcinoma syndrome (NBCCS), also known as Gorlin syndrome, is considered **medically necessary** when:
 - A. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *PTCH1* or *SUFU*, **OR**
 - B. A *PTCH1* or *SUFU* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *PTCH1* or *SUFU* targeted variant analysis (81479) for nevoid basal cell carcinoma syndrome (NBCC), also known as Gorlin syndrome, is considered **investigational** for all other indications.

***PTCH1* and *SUFU* Sequencing and/or Deletion/Duplication Analysis**

- I. *PTCH1* and *SUFU* sequencing and/or deletion duplication analysis (81479) for nevoid basal cell carcinoma syndrome (NBCC), also known as Gorlin syndrome, is considered **medically necessary** when:
 - A. The member/enrollee has a personal history of either of the following:
 - 1. At least two of the following:
 - a) Lamellar calcification of the falx, **OR**
 - b) Jaw keratocyst, **OR**
 - c) Palmar/plantar pits (2 or more), **OR**

- d) Multiple basal cell carcinomas (more than 5 in lifetime) or a basal cell carcinoma diagnosed before 30 years of age, **OR**
 - e) A first-degree relative with NBCCS, **AND**
2. At least one of the following:
- a) Childhood medulloblastoma, **OR**
 - b) Lympho-mesenteric or pleural cysts, **OR**
 - c) Macrocephaly (OFC greater than 97th centile), **OR**
 - d) Cleft lip/palate, **OR**
 - e) Vertebral/rib anomalies (bifid/splayed/extra ribs; bifid vertebrae), **OR**
 - f) Pre- or post-axial polydactyly, **OR**
 - g) Ovarian fibromas, **OR**
 - h) Cardiac fibromas, **OR**
 - i) Ocular anomalies (examples: cataract, pigmentary changes of the retinal epithelium, developmental defects), **OR**
3. At least one of the following:
- a) Lamellar calcification of the falx, **OR**
 - b) Jaw keratocyst, **OR**
 - c) Palmar/plantar pits (2 or more), **OR**
 - d) Multiple basal cell carcinomas (more than 5 in lifetime) or a basal cell carcinoma diagnosed before 30 years of age, **OR**
 - e) A first-degree relative with NBCCS, **AND**
4. At least three of the following:
- a) Childhood medulloblastoma, **OR**
 - b) Lympho-mesenteric or pleural cysts, **OR**

- c) Macrocephaly (OFC greater than 97th centile), **OR**
 - d) Cleft lip/palate, **OR**
 - e) Vertebral/rib anomalies (bifid/splayed/extra ribs; bifid vertebrae), **OR**
 - f) Pre- or post-axial polydactyly, **OR**
 - g) Ovarian fibromas, **OR**
 - h) Cardiac fibromas, **OR**
 - i) Ocular anomalies (examples: cataract, pigmentary changes of the retinal epithelium, developmental defects).
- II. *PTCH1* and *SUFU* sequencing and/or deletion/duplication analysis (81479) is considered **investigational** for all other indications.

[back to top](#)

HEREDITARY PARAGANGLIOMA/PHEOCHROMOCYTOMA SYNDROME (PGL/PCC)

MAX, *SDHA*, *SDHAF2*, *SDHB*, *SDHC*, *SDHD*, or *TMEM127* Targeted Variant Analysis

- I. *MAX*, *SDHA*, *SDHAF2*, *SDHB*, *SDHC*, *SDHD*, or *TMEM127* targeted variant analysis (81403) for hereditary paraganglioma/pheochromocytoma syndrome (PGL/PCC) is considered **medically necessary** when:
 - A. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *MAX*, *SDHA*, *SDHAF2*, *SDHB*, *SDHC*, *SDHD*, or *TMEM127*, **OR**
 - B. A *MAX*, *SDHA*, *SDHAF2*, *SDHB*, *SDHC*, *SDHD*, or *TMEM127* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *MAX*, *SDHA*, *SDHAF2*, *SDHB*, *SDHC*, *SDHD*, or *TMEM127* targeted variant analysis (81403) for hereditary paraganglioma/pheochromocytoma syndrome (PGL/PCC) is considered **investigational** for all other indications.

***MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, and TMEM127* Sequencing and Deletion Duplication Analysis**

- I. *MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, and TMEM127* sequencing and/or deletion/duplication analysis (81404, 81405, 81406, 81479) for hereditary paraganglioma/pheochromocytoma syndrome (PGL/PCC) is considered **medically necessary** when:
 - A. The member/enrollee has a diagnosis of one or more of the following:
 1. Pheochromocytoma, **OR**
 2. Paraganglioma, **OR**
 3. Clear cell renal cell cancer, **OR**
 4. Gastrointestinal stromal tumor (GIST), **OR**
 5. Pulmonary chondromas, **OR**
 - B. The member/enrollee has a family history of paraganglioma or pheochromocytoma.
- II. *MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, and TMEM127* sequencing and/or deletion/duplication (81404, 81405, 81406, 81479) for hereditary paraganglioma/pheochromocytoma syndrome (PGL/PCC) is considered **investigational** for all other indications.

[back to top](#)

PEUTZ-JEGHERS SYNDROME (PJS)

***STK11* Targeted Variant Analysis**

- I. *STK11* targeted variant analysis (81479) for Peutz-Jeghers syndrome is considered **medically necessary** when:
 - A. The member/enrollee has a blood relative with a known pathogenic or likely pathogenic variant in *STK11*, **OR**
 - B. An *STK11* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.

- II. *STK11* targeted variant analysis (81479) for Peutz-Jeghers syndrome is considered **investigational** for all other indications.

***STK11* Sequencing and/or Deletion/Duplication Analysis**

- I. *STK11* sequencing and/or deletion/duplication analysis (81404, 81405) for Peutz-Jeghers syndrome (PJS) is considered **medically necessary** when:
 - A. The member/enrollee has a clinical diagnosis of Peutz-Jeghers syndrome based on the presence of any two of the following:
 - 1. At least two histologically confirmed Peutz-Jeghers-type hamartomatous polyps of the GI tract, **OR**
 - 2. Mucocutaneous pigmentation of the mouth, lips, nose, eyes, genitalia, or fingers, **OR**
 - 3. A family history of PJS.
- II. *STK11* sequencing and/or deletion/duplication analysis (81404, 81405) for Peutz-Jeghers syndrome is considered **investigational** for all other indications.

[back to top](#)

RETINOBLASTOMA

***RB1* Targeted Variant Analysis**

- I. *RB1* targeted variant analysis (81403) for retinoblastoma is considered **medically necessary** when:
 - A. The member/enrollee has a [close relative](#) with a known pathogenic or likely pathogenic variant in *RB1*, **OR**
 - B. An *RB1* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *RB1* targeted variant analysis (81403) for retinoblastoma is considered **investigational** for all other indications.

***RB1* Sequencing and/or Deletion/Duplication Analysis**

- I. *RB1* sequencing and/or deletion/duplication analysis (81479, S3841) for retinoblastoma is considered **medically necessary** when:
 - A. The member/enrollee has a diagnosis of retinoblastoma in one or both eyes, **OR**
 - B. The member/enrollee has a family history of retinoblastoma in one or both eyes
- II. *RB1* sequencing and/or deletion/duplication analysis (81479, S3841) for retinoblastoma is considered **investigational** for all other indications.

[back to top](#)

VON HIPPEL-LINDAU SYNDROME (VHL)

***VHL* Targeted Variant Analysis**

- I. *VHL* targeted variant analysis (81403) for Von Hippel-Lindau syndrome is considered **medically necessary** when:
 - A. The member/enrollee has a [first- or second-degree relative](#) with a known pathogenic or likely pathogenic variant in *VHL*, **OR**
 - B. A *VHL* pathogenic or likely pathogenic variant was detected by tumor profiling and germline analysis has not yet been performed.
- II. *VHL* targeted variant analysis (81403) for Von Hippel-Lindau syndrome is considered **investigational** for all other indications.

***VHL* Sequencing and/or Deletion/Duplication Analysis**

- I. *VHL* sequencing and/or deletion/duplication analysis (81403, 81404, S3842) for Von Hippel-Lindau syndrome is considered **medically necessary** when:
 - A. The member/enrollee has a diagnosis of one or more of the following:
 1. Hemangioblastoma of the retina, spine, or brain, **OR**
 2. Clear cell renal cell carcinoma diagnosed before age 40 years, **OR**
 3. Multiple and/or bilateral clear cell renal cell carcinoma diagnosed at any age, **OR**

4. Pheochromocytoma or paraganglioma (in abdomen, thorax, or neck), **OR**
 5. Retinal angiomas, **OR**
 6. Endolymphatic sac tumor, **OR**
 7. Epididymal or adnexal papillary cystadenoma, **OR**
 8. Pancreatic serous cystadenoma, **OR**
 9. Pancreatic neuroendocrine tumors, **OR**
 10. Multiple renal, pancreatic or hepatic cysts.
- II. *VHL* sequencing and/or deletion/duplication analysis (81403, 81404, S3842) for Von Hippel-Lindau syndrome is considered **investigational** for all other indications.

[back to top](#)

NOTES AND DEFINITIONS

1. **Close relatives** include first, second, and third degree blood relatives on the same side of the family:
 - a. **First-degree relatives** are parents, siblings, and children
 - b. **Second-degree relatives** are grandparents, aunts, uncles, nieces, nephews, grandchildren, and half siblings
 - c. **Third-degree relatives** are great grandparents, great aunts, great uncles, great grandchildren, and first cousins
2. A **limited history** is defined as a member/enrollee who has fewer than two 1st- or 2nd-degree female relatives in the same lineage that lived to age 45. The "limited family history" can occur on either side of the family. A 3-generation pedigree is needed to assess whether family history is limited
3. "**Breast cancer**" applies to patients with invasive cancer or ductal carcinoma in situ (DCIS).
4. **High-risk** breast cancer for olaparib therapy is defined as
 - a. Triple negative breast cancer treated with either:

- i. Adjuvant chemotherapy with axillary node-positive disease or an invasive primary tumor greater than or equal to 2 cm on pathology analysis, **OR**
 - ii. Neoadjuvant chemotherapy with residual invasive breast cancer in the breast or resected lymph nodes, **OR**
 - b. Hormone receptor positive disease treated with either:
 - i. Adjuvant chemotherapy with four or more positive pathologically confirmed lymph nodes, OR
 - ii. Neoadjuvant chemotherapy which did not have a complete pathologic response, with a CPS+CG score [pre-treatment clinical (CS) and post-treatment pathological stage (PS), estrogen-receptor status (E) and grade (G)] of 3 or higher
- 5. **Juvenile polyps** are polyps associated with Juvenile Polyposis Syndrome. These polyps are exophytic and eroded. They typically contain the following: marked edema and inflammation within the lamina propria, cystic glands filled with thick mucin, and some degree of smooth muscle proliferation.
- 6. **ClinGen** is a National Institutes of Health (NIH)-funded resource dedicated to building a central resource that defines the clinical relevance of genes and variants for use in precision medicine and research.
- 7. **Maori ancestry** describes individuals who are of indigenous New Zealand ethnic background
- 8. **High-risk-prostate cancer** is defined by NCCN as an individual who has no very-high-risk features but has exactly one of the following high-risk features:
 - a. cT3a, OR
 - b. Grade Group 4 or Grade Group 5, OR
 - c. PSA > 20ng/ml
- 9. **Very-high-risk prostate cancer** is defined by NCCN as an individual who has at least one of the following:
 - a. CT3b-cT4
 - b. Primary Gleason pattern 5
 - c. 2 or 3 high-risk features
 - d. >4 cores with Grade Group 4 or 5
- 10. **Targeted mutation testing** is the process of analyzing one specific pathogenic or likely pathogenic (P/LP) variant in one gene. This testing is typically performed when there is a

known familial mutation, or in cases where a P/LP variant is identified on somatic tumor profiling.

11. **Adenomatous polyposis** describes conditions that cause multiple adenomas (i.e., benign polyps) in the gastrointestinal tract.
12. **Lynch Syndrome related cancer** is defined as any of the following cancer types: colorectal, endometrial, gastric, ovarian, pancreatic, ureter and renal pelvic, brain (usually glioblastoma), biliary tract, small intestinal, sebaceous adenoma, sebaceous carcinoma, or keratoacanthoma.

BACKGROUND AND RATIONALE

Pan-Cancer Hereditary Susceptibility Panels

National Comprehensive Cancer Network (NCCN)

NCCN Breast, Ovarian, and/or Pancreatic Cancer Genetic Assessment guidelines (3.2023) recognize that next-generation sequencing technology has rapidly altered the clinical approach to testing at-risk patients and their families for hereditary forms of cancer and that when more than one gene can explain an inherited cancer syndrome, tailored multi-gene testing is often more efficient and/or cost effective than single-gene testing. However, there is the chance of finding a variant of uncertain significance in a well established gene, or finding a pathogenic variant in a gene with uncertain clinical management. These types of findings increase as additional genes are included in the multi-gene panel. It is recommended that multi-gene panel testing be offered by a professional genetic expert that provides detailed pre- and post-test counseling. (p. EVAL-A 3 of 10)

NCCN Guidelines for Genetic/Familial High-Risk Assessment Colorectal (1.2023) state that when more than one gene can explain an inherited cancer syndrome, multigene testing is more efficient than single-gene testing, or sequential single syndrome testing. There is also a role for multigene testing in individuals who have tested negative (indeterminate) for a single syndrome, but whose personal or family history remains strongly suggestive of an inherited susceptibility. (p. GENE-1)

National Society of Genetic Counselors (NSGC)

The National Society of Genetic Counselors released a position statement (2017) endorsing the use of multi-gene panels when clinically warranted and appropriately applied, stating the following:

“These tests can provide a comprehensive and efficient route to identifying the genetic causes of disease. Before ordering a multi-gene panel test, providers should thoroughly evaluate the analytic and clinical validity of the test, as well as its clinical utility. Additional factors to consider include, but are not limited to: clinical and family history information, gene content of the panel, limitations of the sequencing and informatics technologies, and variant interpretation and reporting practices.

Panels magnify the complexities of genetic testing and underscore the value of experts, such as genetic counselors, who can educate stakeholders about appropriate utilization of the technology to mitigate risks of patient harm and unnecessary costs to the healthcare system. NSGC supports straightforward and transparent pricing so that patients, providers, laboratories, and health plans can easily weigh the value of genetic testing in light of its cost.”

American College of Obstetricians and Gynecologists

ACOG published Committee Opinion Number 793 (2019) regarding hereditary cancer syndromes and risk assessment that included the following recommendations:

- A hereditary cancer risk assessment is the key to identifying patients and families who may be at increased risk of developing certain types of cancer. Assessments should be performed by obstetrician–gynecologists or other obstetric–gynecologic care providers and should be updated regularly.
- If a hereditary cancer risk assessment suggests an increased risk of a hereditary cancer syndrome, referral to a specialist in cancer genetics or a health care provider with expertise in genetics is recommended for expanded gathering of family history information, risk assessment, education, and counseling, which may lead to genetic testing and tailored cancer screening or risk reduction measures, or both.
- Genetic testing may be performed using a panel of multiple genes through next-generation sequencing technology. This multigene testing process increases the likelihood of finding variants of unknown significance, and it also allows for testing for pathogenic and likely pathogenic variants in multiple genes that may be associated with a specific cancer syndrome or family cancer phenotype (or multiple phenotypes). (p. e143)

Hereditary Breast Cancer Susceptibility Panels

National Comprehensive Cancer Network (NCCN)

The NCCN guidelines for Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic Cancers (3.2023, p. CRIT-2) outline clinical criteria for germline genetic testing of

high-penetrance breast cancer genes. These guidelines include:

1.) Personal history of breast cancer at 50 years of age or younger.

2.) Personal history of breast cancer at any age with specific features:

- Treatment indications
 - To aid in systemic treatment decisions using PARP inhibitors for breast cancer in the metastatic setting
 - To aid in adjuvant treatment decisions with olaparib for high-risk, HER2-negative breast cancer, triple-negative breast cancer
- Pathology/histology
 - Triple-negative breast cancer
 - Multiple primary breast cancers (synchronous or metachronous)
 - Male breast cancer
- Ashkenazi Jewish ancestry
- Family history of at least 1 close blood relative with:
 - Breast cancer at age 50 years or younger
 - Male breast cancer
 - Ovarian cancer
 - Pancreatic cancer
 - Prostate cancer with metastatic, or high- or very-high-risk group
 - 3 or more total diagnoses of breast cancer in patient and/or close blood relatives
 - 2 or more close blood relatives with either breast or prostate cancer (any grade)

3.) Personal history of epithelial ovarian cancer (including fallopian tube cancer or peritoneal cancer) at any age.

4.) Personal history of metastatic prostate cancer OR high- or very-high-risk group prostate cancer).

5.) Family history-based criteria: An affected individual (not meeting testing criteria listed above) or unaffected individual with a first- or second-degree blood relative meeting any of the criteria listed above (except unaffected individuals whose relatives meet criteria only for systemic therapy decision-making). If the affected relative has pancreatic cancer or prostate cancer only first-degree relatives should be offered testing unless indicated based on additional family history

6.) An affected or unaffected individual who otherwise does not meet the criteria above but has a probability of greater than 5% of a *BRCA1/2* pathogenic variant based on prior probability models (eg, Tyrer-Cuzick, BRCAPro, CanRisk).

Hereditary GI/Colon Cancer Panel Tests

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Genetic/Familial High-Risk Assessment: Colorectal (1.2023) outline criteria for assessment for hereditary colorectal syndromes as follows:

- Polyposis: Patient with a personal history of, or a single family member with, at least 10 adenomatous polyps, at least 2 hamartomatous polyps, or at least 5 serrated polyps/lesions proximal to the rectum (p. HRS-1)
- Personal history of colorectal cancer: Patient is under 50 years at age of diagnosis, or meets Lynch syndrome criteria (p. HRS-1, HRS-3, LS-1) (see [MLH1, MSH2, MSH6, PMS2, EPCAM Sequencing and/or Deletion/Duplication Analysis](#))
- Personal of family history of Lynch syndrome-related cancer that meets Lynch syndrome criteria (p. HRS-3, LS-1) (see [MLH1, MSH2, MSH6, PMS2, EPCAM Sequencing and/or Deletion/Duplication Analysis](#)).

NCCN also states that the minimum number of CRC-risk associated genes to include in germline multi-gene panel testing are as follows: *APC, MUTYH, MLH1, MSH2, MSH6, PMS2, EPCAM, BMPR1A, SMAD4, PTEN, STK11*, and *TP53*. (p. HRS-A 2 of 2)

Hereditary Gastric Cancer Panels

National Comprehensive Cancer Network (NCCN)

NCCN Gastric Cancer guidelines (1.2023) outline criteria for further genetic risk assessment for high-risk syndromes associated with gastric cancer, including: hereditary diffuse gastric cancer, Lynch syndrome, Juvenile Polyposis Syndrome, Peutz-Jeghers syndrome, and Familial Adenomatous Polyposis. (p. GAST-D 3 of 7 and p. GAST-D 4 of 7)

Hereditary Pancreatic Cancer Susceptibility Panels

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) recommend genetic counseling and germline testing for all individuals diagnosed with exocrine pancreatic cancer, as well as individuals with a first-degree relative diagnosed with exocrine pancreatic cancer. These guidelines list the following genes as those that are typically tested for pancreatic cancer risks: *ATM, BRCA1, BRCA2, CDKN2A, MLH1, MSH2, MSH6*,

EPCAM, PALB2, STK11, TP53. (p. CRIT-5)

Hereditary Polyposis Panels

National Comprehensive Cancer Network (NCCN)

The NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline recommendations for evaluating individuals with adenomatous polyposis (defined as 10 or more adenomas) for germline mutations in *APC* and *MUTYH*. (p. HRS-2)

Hereditary Prostate Cancer Susceptibility Panels

National Comprehensive Cancer Network (NCCN)

NCCN Prostate Cancer guidelines (3.2023) state that germline testing is recommended in patients with a personal history of prostate cancer in the following scenarios:

- By prostate cancer stage or risk group (diagnosed at any age)
- Metastatic, regional (node positive), very-high-risk localized, or high-risk localized prostate cancer
- By family history and/or ancestry:
- 1 or more first-, second-, or third-degree relative with:
 - breast cancer at age <50 y
 - colorectal or endometrial cancer at age <50 y
 - male (sex assigned at birth) breast cancer at any age
 - ovarian cancer at any age
 - exocrine pancreatic cancer at any age
 - metastatic, regional, very-high-risk, or high-risk prostate cancer at any age
 - 1 or more first-degree relative (parent or sibling) with:
 - prostate cancer at age <60 y
 - 2 or more first-, second-, or third-degree relatives with:
 - breast cancer at any age
 - prostate cancer at any age
 - 3 or more first- or second-degree relatives with:
 - Lynch syndrome-related cancers, especially if diagnosed <50 y: colorectal, endometrial, gastric, ovarian, exocrine pancreas, upper tract urothelial, glioblastoma, biliary tract, and small intestinal cancer
- A known family history of familial cancer risk mutation (pathogenic/likely pathogenic variants), especially in: *BRCA1, BRCA2, ATM, PALB2, CHEK2*,

- MLH1, MSH2, MSH6, PMS2, and EPCAM*
- Ashkenazi Jewish ancestry

NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) recommend the following testing criteria for prostate cancer susceptibility genes:

Personal history of prostate cancer with specific clinical features: metastatic disease, specific histology (intraductal/ cribriform, high- or very-high risk group), or with specific family history/ancestry features: 1 or more close blood relative with breast cancer at age 50 years or younger; triple-negative breast cancer at any age; male breast cancer at any age; ovarian cancer any age; pancreatic cancer any age; metastatic, intraductal/ cribriform histology, or high- or very-high risk group at any age; 2 or more close blood relatives with either breast or prostate cancer (any grade) at any age; Ashkenazi Jewish ancestry. Another fulfilling criterion is an individual with or without prostate cancer affected (not meeting testing criteria listed above) with a first-degree blood relative meeting any of the criteria listed above (except unaffected individuals whose relatives meet criteria only for systemic therapy decision-making). (p. CRIT-6)

Hereditary Neuroendocrine Cancer Susceptibility Panels

National Comprehensive Cancer Network (NCCN)

The NCCN Neuroendocrine and Adrenal Tumors Guideline (1.2023) states that multigene panel testing may be a more efficient and cost-effective solution for evaluating a patient for a hereditary endocrine cancer syndrome, as there is clinical overlap between several genetic conditions that predispose to endocrine neoplasms. (p. NE-E 2 of 8)

These guidelines also outline common clinical and tumor manifestations of different hereditary endocrine neoplasia syndromes, including hereditary paraganglioma/pheochromocytoma syndrome, multiple endocrine neoplasia type 1 (MEN1), multiple endocrine neoplasia type 2 (MEN2), and von Hippel-Lindau syndrome. (p. NE-E 4 of 8 and p. NE-E 5 of 8)

BRCA1 AND BRCA2 GENE TESTING

BRCA1/BRCA2 Targeted Variant or Known Familial Variant Analysis

National Comprehensive Cancer Network (NCCN)

The NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) states that testing should be performed in the following situations:

- 1) Individuals with any blood relative with a known pathogenic/likely pathogenic variant in a cancer susceptibility gene
- 2) Individuals with a pathogenic/likely pathogenic (P/LP) variant identified on tumor genomic testing that has clinical implications if also identified in the germline. (p.CRIT-1)

BRCA1/BRCA2 Targeted Variant Analysis - Ashkenazi Jewish Founder Variants

National Comprehensive Cancer Network (NCCN)

The NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) states that testing for the three known Ashkenazi Jewish founder *BRCA1/2* mutations is appropriate for individuals who are age 18 years or older and have at least one grandparent who is of Ashkenazi Jewish ancestry. (p. CRIT-6 and p. CRIT-6A)

BRCA1 and BRCA2 Sequencing and/or Deletion/Duplication Analysis

National Comprehensive Cancer Network (NCCN)

The NCCN guidelines for Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) outline clinical criteria for germline genetic testing of high-penetrance breast cancer genes, including *BRCA1* and *BRCA2*. These guidelines include:

Personal history of breast cancer with specific features:

- Diagnosed 50 years of age or younger
- Diagnosed at any age: To aid in systemic treatment decisions using PARP inhibitors for breast cancer in the metastatic setting; to aid in adjuvant treatment decisions with olaparib for high-risk, HER2-negative breast cancer; triple-negative breast cancer; multiple primary breast cancers (synchronous or metachronous); Male breast cancer; Ashkenazi Jewish ancestry; at least 1 close blood relative with: breast cancer at age 50 years or younger, male breast cancer, ovarian cancer, pancreatic cancer, prostate cancer with metastatic, or high- or very-high-risk group, 3 or more total diagnoses of breast cancer in patient and/or close blood relatives, 2 or more close blood relatives with either breast or prostate cancer (any grade).
- Family history-based criteria: An affected individual (not meeting testing criteria listed

above) or unaffected individual with a first- or second degree blood relative meeting any of the criteria listed above (except unaffected individuals whose relatives meet criteria only for systemic therapy decision-making). If the affected relative has pancreatic cancer or prostate cancer only first-degree relatives should be offered testing unless indicated based on additional family history.

- An affected or unaffected individual who otherwise does not meet the criteria above but has a probability of greater than 5% of a *BRCA1/2* pathogenic variant based on prior probability models (eg, Tyrer-Cuzick, BRCAPro, CanRisk) (p. CRIT-2)

US Preventive Services Task Force (USPSTF)

The USPSTF published a recommendation statement (2019) on risk assessment, genetic counseling, and genetic testing for *BRCA*-related cancer that included the following conclusion and recommendation:

“The USPSTF recommends that primary care clinicians assess women with a personal or family history of breast, ovarian, tubal, or peritoneal cancer or who have an ancestry associated with *BRCA1/2* gene mutations with an appropriate brief familial risk assessment tool. Women with a positive result on the risk assessment tool should receive genetic counseling and, if indicated after counseling, genetic testing. (B recommendation) The USPSTF recommends against routine risk assessment, genetic counseling, or genetic testing for women whose personal or family history or ancestry is not associated with potentially harmful *BRCA1/2* gene mutations. (D recommendation)”. (p. 652)

***PALB2* GENE TESTING**

***PALB2* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

The NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) states that testing should be performed in the following situations:

- Individuals with any blood relative with a known pathogenic/likely pathogenic variant in a cancer susceptibility gene
- Individuals with a pathogenic/likely pathogenic (P/LP) variant identified on tumor genomic testing that has clinical implications if also identified in the germline. (p.CRIT-1)

***PALB2* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

The NCCN guidelines for Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) outline clinical criteria for germline genetic testing of high-penetrance breast cancer genes, including *PALB2*. These guidelines include:

Personal history of breast cancer with specific features:

- Diagnosed 50 years of age or younger
- Diagnosed at any age: To aid in systemic treatment decisions using PARP inhibitors for breast cancer in the metastatic setting; to aid in adjuvant treatment decisions with olaparib for high-risk, HER2-negative breast cancer, triple-negative breast cancer; multiple primary breast cancers (synchronous or metachronous); Male breast cancer; Ashkenazi Jewish ancestry; at least 1 close blood relative with: breast cancer at age 50 years or younger, male breast cancer, ovarian cancer, pancreatic cancer, prostate cancer with metastatic, or high- or very-high-risk group, 3 or more total diagnoses of breast cancer in patient and/or close blood relatives, 2 or more close blood relatives with either breast or prostate cancer (any grade),
- Family history-based criteria: An affected individual (not meeting testing criteria listed above) or unaffected individual with a first- or second degree blood relative meeting any of the criteria listed above (except unaffected individuals whose relatives meet criteria only for systemic therapy decision-making). If the affected relative has pancreatic cancer or prostate cancer only first-degree relatives should be offered testing unless indicated based on additional family history.
- An affected or unaffected individual who otherwise does not meet the criteria above but has a probability of greater than 5% of a *BRCA1/2* pathogenic variant based on prior probability models (eg, Tyrer-Cuzick, BRCAPro, CanRisk) (p. CRIT-2)

ATM AND CHEK2 GENE TESTING

ATM or CHEK2 Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

The NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) state that testing should be performed in the following situations:

- Individuals with any blood relative with a known pathogenic/likely pathogenic variant in a cancer susceptibility gene

- Individuals with a pathogenic/likely pathogenic (P/LP) variant identified on tumor genomic testing that has clinical implications if also identified in the germline. (p.CRIT-1)

ATM or CHEK2 Sequencing and/or Deletion/Duplication Analysis

National Comprehensive Cancer Network (NCCN)

While the NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) do provide surveillance recommendations for individuals with germline *ATM* and *CHEK2* mutations (p. GENE-A 1 of 10 and p. GENE-A 4 of 10), these genes are not considered high-penetrance breast cancer susceptibility genes, and the guidelines do not include gene-specific clinical criteria for *ATM* and *CHEK2* as they do for the high-penetrance breast cancer susceptibility genes.

LYNCH SYNDROME/HEREDITARY NONPOLYPOSIS COLORECTAL CANCER (HNPCC) TESTING

MLH1, MSH2, MSH6, PMS2, or EPCAM Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline testing criteria for the evaluation of Lynch syndrome. NCCN recommends analysis of *MLH1*, *MSH2*, *MSH6*, *PMS2* and/or *EPCAM* in individuals with a known pathogenic variant in the family. (p. HRS-5)

Additionally, NCCN states that tumor testing can be complementary to germline testing and can assist in interpretation of results. Although germline origin can sometimes be inferred with a high degree of confidence, confirmatory germline testing is indicated for pathogenic/likely pathogenic variants with a reasonable clinical suspicion of being a germline origin (based on patient/family history or clinical characteristics, presence of founder mutation, and in some cases variant allele frequency). (p. HRS-A 4 of 7)

MLH1, MSH2, MSH6, PMS2, or EPCAM Sequencing and/or Deletion/Duplication Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline testing criteria for the evaluation of Lynch syndrome. NCCN recommends analysis of *MLH1*, *MSH2*,

MSH6, *PMS2* and/or *EPCAM* in individuals with a personal and/or family history of Lynch syndrome-related cancers, such as colorectal, endometrial, gastric, ovarian, pancreatic, ureter and renal pelvic, brain (usually glioblastoma), biliary tract, small intestinal, sebaceous adenoma, sebaceous carcinoma, or keratoacanthoma. These criteria include:

- An individual with a Lynch-syndrome (LS)-related cancer or cancer and any of the following: Diagnosed younger than 50 years; a synchronous or metachronous LS -related cancer regardless of age; 1 first-degree or second-degree relative with an LS-related cancer diagnosed younger than 50 years; or 2 or more first-degree or second-degree relatives with an LS-related cancer regardless of age
- Family history of any of the following: at least 1 first-degree relative with a colorectal or endometrial cancer diagnosed younger than 50 years; at least 1 first-degree relative with a colorectal or endometrial cancer and a synchronous or metachronous LS-related cancer regardless of age; 2 or more first-degree or second-degree relatives with LS-related cancers, one of whom was diagnosed before age 50; 3 or more first-degree or second-degree relatives with LS-related cancers regardless of age
- An individual with a 5% risk or greater of having an MMR gene pathogenic variant based on predictive models (i.e., PREMM5, MMRpro, MMRpredict)
- An individual with a personal history of CRC and/or endometrial cancer with a PREMM5 score of 2.5% or greater should be considered for multi-gene panel testing.

For individuals without a personal history of CRC and/or endometrial cancer, some data have suggested using a PREMM5 score threshold of 2.5% or greater rather than 5% or greater to select individuals for MMR genetic testing. Based on these data, it is reasonable for testing to be done based on the 2.5% or greater score result and clinical judgment. (p. HRS-5)

NCCN states that the minimum number of CRC-risk associated genes to include in germline multi-gene panel testing are as follows: *APC*, *MUTYH*, *MLH1*, *MSH2*, *MSH6*, *PMS2*, *EPCAM*, *BMPRI1A*, *SMAD4*, *PTEN*, *STK11*, and *TP53*. (p. HRS-A 2 of 2)

***BAP1* TUMOR PREDISPOSITION SYNDROME**

***BAP1* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Kidney Cancer (1.2024) include *BAP1* tumor predisposition syndrome in their overview of hereditary renal cell carcinoma syndromes, and state that this testing is indicated for an individual with a close blood relative with a known pathogenic/likely pathogenic variant. (p. HERED-RCC-1 and HERED-RCC-2)

***BAP1* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Cutaneous Melanoma (2.2023) state that individuals with the presence of germline mutations in *CDKN2a*, *CDK4*, *MC1R*, *BRCA2*, *BAP1* and potentially other genes, are predisposed to develop single or multiple primary melanomas. (p. ME-A 1 of 2)

NCCN guidelines for Uveal Melanoma (1.2023) include germline *BAP1* mutations as a risk factor for developing uveal melanoma. (p. UM-A 1 of 2)

NCCN guidelines for Malignant Pleural Mesothelioma (1.2023) state that approximately 12-16% of patients with pleural or peritoneal mesothelioma have a germline mutation, including in *BAP1*. (p. MPM-A 5 of 8)

NCCN guidelines for Kidney cancer (1.2024) include *BAP1* tumor predisposition syndrome in their overview of hereditary renal cell carcinoma syndromes (p. HERED-RCC-2)

GeneReviews: BAP1 Tumor Predisposition Syndrome (BAP1-TPDS)

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online. The clinical description and testing indications for *BAP1* Tumor Predisposition syndrome are as follows:

BAP1-TPDS should be suspected in an individual who has EITHER of the following:

- Two or more confirmed *BAP1*-TPDS tumors*
- One *BAP1*-TPDS tumor and a first- or second-degree relative with a confirmed *BAP1*-TPDS tumor*

*Excluding two basal cell cancers and/or cutaneous melanomas, given their high frequency in the general population

In addition to *BAP1*-inactivated melanocytic tumors, uveal melanoma, malignant mesothelioma, cutaneous melanoma, renal cell carcinoma, and basal cell carcinoma, individuals with germline mutations in *BAP1* may have an increased risk for hepatocellular carcinoma, cholangiocarcinoma, and meningioma.

BIRT-HOGG DUBE SYNDROME (BHDS)

FLCN Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Kidney Cancer (1.2024) includes Birt-Hogg-Dube syndrome in their overview of hereditary renal cell carcinoma syndromes, and state that this testing is indicated for an individual with a close blood relative with a known pathogenic/likely pathogenic variant. (p. HERED-RCC-1 and HERED-RCC-2)

FLCN Sequencing and/or Deletion/Duplication Analysis

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Kidney Cancer (1.2024 include Birt-Hogg-Dube syndrome in their overview of hereditary renal cell carcinoma syndromes. (p. HERED-RCC-2). Commonly seen histologies include chromophobe, hybrid oncocytic tumors, clear cell, oncocytomas, angiomyolipomas, and papillary RCC. (p. HERED-RCC-2)

GeneReviews: Birt-Hogg-Dube Syndrome (BHDS)

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online. The clinical description and testing indications for Birt-Hogg-Dube syndrome (BHDS) are as follows:

BHDS should be suspected in individuals with any of the following major or minor criteria.

Major criteria

- Five or more fibrofolliculomas/trichodiscomas with at least one confirmed histologically
- Identification of a heterozygous pathogenic variant in *FLCN*

Minor criteria

- Multiple lung cysts. Bilateral basally located lung cysts with no other apparent cause, with or without spontaneous primary pneumothorax
- Early-onset renal cancer (age <50 years)
- Multifocal or bilateral renal cancer
- Renal cancer of mixed chromophobe and oncocytic histology
- First-degree relative with BHDS

The diagnosis of BHDS is established in a proband with:

- One major criteria (Note: Identification of a heterozygous pathogenic variant in FLCN is one of the major criteria); **OR**
- Two minor criteria

COWDEN SYNDROME (CS)/PTEN HAMARTOMA TUMOR SYNDROME (PHTS)

PTEN Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

The NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) states that testing should be performed in the following situations:

- 1) Individuals with any blood relative with a known pathogenic/likely pathogenic variant in a cancer susceptibility gene
- 2) Individuals with a pathogenic/likely pathogenic (P/LP) variant identified on tumor genomic testing that has clinical implications if also identified in the germline. (p.CRIT-1)

PTEN Sequencing and/or Deletion/Duplication Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) outline clinical criteria for the genetic testing for Cowden syndrome (CS)/PTEN hamartoma tumor syndrome (PHTS) in individuals with a personal or family history of PHTS/CS. These include:

- Individual from a family with a known *PTEN* pathogenic or likely pathogenic variant
- Individual with a personal history of Bannayan-Riley-Ruvalcaba syndrome (BRRS)
- Individual meeting clinical diagnostic criteria* for CS/PHTS [Cowden syndrome/PTEN hamartoma tumor syndrome]
- Individual not meeting clinical diagnostic criteria for CS/PHTS with a personal history of: Adult Lhermitte-Duclos disease (cerebellar tumors); Autism spectrum disorder and macrocephaly; Two or more biopsy-proven trichilemmomas; Two or more major criteria (one must be macrocephaly); Three major criteria, without macrocephaly; One major and 3 or more minor criteria; 4 or more minor criteria
- At-risk individual with a relative with a clinical diagnosis of CS/PHTS or BRRS for whom testing has not been performed. The at-risk individual must have the following: Any one major criterion or two minor criteria

- *PTEN* pathogenic or likely pathogenic variant detected by tumor genomic testing on any tumor type in the absence of germline analysis (p. CRIT-8 and CRIT-8A)

*Revised Clinical Diagnostic Criteria for PTEN Hamartoma Tumor Syndrome:

Operational diagnosis in an individual (either of the following:

1. Three or more major criteria, but one must include macrocephaly, Lhermitte-Duclos disease, or GI hamartomas; or
2. Two major and three minor criteria (CRIT-8A)

ADENOMATOUS POLYPOSIS CONDITIONS (FAMILIAL ADENOMATOUS POLYPOSIS SYNDROME (FAP)/ATTENUATED FAP (AFAP) AND MUTYH-ASSOCIATED POLYPOSIS SYNDROME (MAP))

***APC* or *MUTYH* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline clinical criteria for the genetic testing, which includes a known pathogenic variant in an adenomatous polyposis gene in the family. (p. POLYP-1)

Of note, NCCN recommends analysis of *MUTYH* in individuals where the family pathogenic variant is known. Specifically, siblings of a patient with MAP are recommended to have site-specific testing for the familial pathogenic variants. (p. MAP-1 and MAP-3)

Additionally, NCCN states that tumor testing can be complementary to germline testing and can assist in interpretation of results. Although germline origin can sometimes be inferred with a high degree of confidence, confirmatory germline testing is indicated for pathogenic/likely pathogenic variants with a reasonable clinical suspicion of being a germline origin (based on patient/family history or clinical characteristics, presence of founder mutation, and in some cases variant allele frequency). (p. HRS-A 4 of 7)

***APC* and/or *MUTYH* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline Adenomatous Polyposis testing criteria. These include: Personal history of greater than or equal

to 20 cumulative adenomas, known pathogenic variant in adenomatous polyposis gene in family, or multifocal/bilateral congenital hypertrophy of retinal pigment epithelium (CHRPE) (p. POLYP-1)

The guidelines also note that biallelic *MUTYH* mutations have also been implicated in rare cases of serrated polyposis syndrome (defined as 5 or more serrated polyps proximal to the rectum all being 5mm or larger with 2 or more being 10 or more mm in size, or more than 20 serrated polyps of any size distributed throughout the colon, with 5 or more being proximal to the rectum). (p. SPS-1)

FAMILIAL ATYPICAL MULTIPLE MOLE MELANOMA (FAMMM) SYNDROME

***CDKN2A* Targeted Variant Analysis**

Genetic Support Foundation

The Genetic Support Foundation's Genetics 101 information on inheritance patterns says the following about testing for familial pathogenic variants:

Genetic testing for someone who may be at risk for an inherited disease is always easier if we know the specific genetic cause. Oftentimes, the best way to find the genetic cause is to start by testing someone in the family who is known or strongly suspected to have the disease. If their testing is positive, then we can say that we have found the familial pathogenic (harmful) variant. We can use this as a marker to test other members of the family to see who is also at risk.

***CDKN2A* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Cutaneous Melanoma guidelines (2.2023) recommend considering genetic counseling referral for *p16/CDKN2A* mutation testing (and possibly other genes) when a patient has 3 or more invasive cutaneous melanomas, or a personal or family history of a mix of invasive melanoma, pancreatic cancer, and/or astrocytoma diagnoses. (p. ME-11)

HEREDITARY DIFFUSE GASTRIC CANCER (aka, Signet Ring Cell Gastric Cancer):

***CDH1* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Gastric Cancer guidelines (1.2023) outline testing criteria for germline *CDH1* testing, which states that a known mutation in a gastric cancer susceptibility gene in a close relative is criteria for further risk evaluation. (p.GAST-D 1 of 8)

NCCN guidelines for Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic Cancers (3.2023) address tumor genomic testing and state that somatic pathogenic and likely pathogenic variants may be of germline or somatic origin, and germline testing should be considered when clinically indicated. (p. EVAL-A 5 of 10)

***CDH1* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Gastric Cancer guidelines (1.2023) outline testing criteria for germline *CDH1* testing which incorporates both personal and family history of gastric cancer and lobular breast cancer. These include:

- Two gastric cancer cases in a family, one confirmed diffuse gastric cancer (DGC) regardless of age
- DGC diagnosed before age 50 years without a family history
- Personal or family history of DGC and lobular breast cancer, one diagnosed before age 70 years
- Two cases of lobular breast cancer in family members before 50 years of age
- DGC at any age in individuals of Māori ethnicity, or with a personal or family history of cleft lip/cleft palate
- Bilateral lobular breast cancer before age 70 years. (p. GAST-D 3 of 7)

JUVENILE POLYPOSIS SYNDROME (JPS)

***SMAD4* and *BMPRIA* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline clinical criteria for the genetic testing, which states that genetic testing should be performed for individuals with a family history of JPS. (p. JPS-1)

Additionally, NCCN states that tumor testing can be complementary to germline testing and can

assist in interpretation of results. Although germline origin can sometimes be inferred with a high degree of confidence, confirmatory germline testing is indicated for pathogenic/likely pathogenic variants with a reasonable clinical suspicion of being a germline origin (based on patient/family history or clinical characteristics, presence of founder mutation, and in some cases variant allele frequency). (p. HRS-A 4 of 7)

***SMAD4* and *BMPRI1A* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline clinical criteria for the genetic testing for juvenile polyposis syndrome (JPS) in individuals with a personal and/or family history suggestive of JPS, noting that clinical genetic testing is recommended as approximately 50% of JPS cases occurring due to pathogenic variants in *BMPRI1A* and *SMAD4*. These criteria include 5 or more colonic juvenile polyps, multiple juvenile polyps throughout the gastrointestinal tract, and any number of juvenile polyps in someone with a family history of JPS. (p. JPS-1) These guidelines also acknowledge that pathogenic or likely pathogenic variants report in tumor genetic analyses may be of germline or somatic origin, and confirmatory germline testing is indicated if applicable. (p. HRS-A 4 of 7).

HEREDITARY LEIOMYOMATOSIS AND RENAL CELL CANCER (HLRCC)

***FH* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Kidney Cancer (1.2024) include Hereditary leiomyomatosis and renal cell carcinoma (HLRCC) in their overview of hereditary renal cell carcinoma syndromes, and state that this testing is indicated for an individual with a close blood relative with a known pathogenic/likely pathogenic variant. (p. HERED-RCC-1 and HERED-RCC-2)

***FH* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Kidney Cancer (1.2024) outline criteria for further genetic risk evaluation for hereditary renal cell carcinoma syndromes, including HLRCC-associated renal cell carcinoma. (p. HERED-RCC-2)

GeneReviews: FH Tumor Predisposition Syndrome

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online. The recommended testing for FH tumor predisposition syndrome (HLRCC) is as follows:

FH tumor predisposition syndrome should be suspected in individuals with the following features:

Cutaneous leiomyomata (~50%):

- Skin-colored to light brown/reddish papules or nodules distributed over the trunk, extremities, and occasionally on the face and neck
- May be single, grouped/clustered, segmental, or disseminated
- Histopathology shows bundles of smooth muscle fibers with central, long blunt-edged nuclei

Uterine leiomyomata (uterine fibroids) (~90% of females):

- Fibroids tend to be numerous and large.
- Fibroids often demonstrate loss of FH staining and positive cytoplasmic staining for S-(2-succino) cysteine

Renal tumors (~15%) are usually solitary, highly aggressive renal cell carcinoma (RCC) that metastasizes early.

The spectrum of renal tumors includes type 2 papillary, undefined papillary, unclassified, tubulocystic, and collecting-duct carcinoma.

LI-FRAUMENI SYNDROME (LFS)

***TP53* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

The NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) states that testing should be performed in the following situations:

- 1) Individuals with any blood relative with a known pathogenic/likely pathogenic variant in a cancer susceptibility gene
- 2) Individuals with a pathogenic/likely pathogenic (P/LP) variant identified on tumor genomic testing that has clinical implications if also identified in the germline. (p.CRIT-1)

TP53 Sequencing and/or Deletion/Duplication Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic guidelines (3.2023) outline clinical testing criteria for the genetic testing for Li-Fraumeni syndrome. This includes classic Li-Fraumeni syndrome criteria and Chompret criteria and considerations for family history:

Classic Li-Fraumeni syndrome (LFS) criteria:

- Combination of an individual diagnosed at age younger than 45 years with a sarcoma **AND**
- A first-degree relative diagnosed at age younger than 45 years with cancer **AND**
- An additional first- or second-degree relative in the same lineage with cancer diagnosed at age younger than 45 years, or a sarcoma at any age

Chompret criteria:

- Individual with a tumor from LFS tumor spectrum (e.g., soft tissue sarcoma, osteosarcoma, CNS tumor, breast cancer, adrenocortical carcinoma), before 46 years of age, **AND**
 - At least one first- or second-degree relative with any of the aforementioned cancers (other than breast cancer if the proband has breast cancer) before the age of 56 years or with multiple primaries at any age, **OR**
- Individual with multiple tumors (except multiple breast tumors), two of which belong to LFS tumor spectrum with the initial cancer occurring before the age of 46 years, **OR**
- Individual with adrenocortical carcinoma, or choroid plexus carcinoma or rhabdomyosarcoma of embryonal anaplastic subtype, at any age of onset, regardless of family history, **OR**
- Breast cancer before 31 years of age, **OR**
- Pediatric hypodiploid acute lymphoblastic leukemia, **OR**
- Affected individual with pathogenic/likely pathogenic variant identified on tumor genomic testing that may have implications if also identified on germline testing (p. CRIT-7)

MULTIPLE ENDOCRINE NEOPLASIA TYPE 1 (MEN1)

MEN1 Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Neuroendocrine and Adrenal Tumors guidelines (1.2023) recommend that targeted genetic testing for *MEN1* be performed for individuals with a close blood relative with a known pathogenic variant/likely pathogenic variant in a cancer susceptibility gene. (p. NE-E 3 of 8)

Additionally, NCCN recommends genetic risk evaluation and genetic testing for Hereditary Endocrine Neoplasia Syndromes when a mutation is identified on tumor genomic testing that has clinical implications if also identified in the germline. (p NE-E 3 of 8)

***MEN1* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Neuroendocrine and Adrenal Tumors guidelines (1.2023) recommend that patients with two or more of the following, or 1 AND a family history of 1 or more of the following, be evaluated for *MEN1* germline mutations:

- Foregut carcinoid (bronchial, thymic, or gastric)
- Pituitary adenoma
- Duodenal or pancreatic neuroendocrine tumor
- Primary hyperparathyroidism. (p. NE-E 3 of 8)

MULTIPLE ENDOCRINE NEOPLASIA TYPE 2 (MEN2)

***RET* Targeted Variant Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Neuroendocrine and Adrenal Tumors guidelines (1.2023) recommend that targeted genetic testing for MEN2 be performed for individuals with a close blood relative with a known pathogenic variant/likely pathogenic variant in a cancer susceptibility gene. (p. NE-E 3 of 8).

Additionally, NCCN states that testing is recommended when a mutation is identified on tumor genomic testing that has clinical implications if also identified in the germline. (p NE-E 3 of 8)

***RET* Sequencing and/or Deletion/Duplication Analysis**

GeneReviews: Multiple Endocrine Neoplasia Type 2

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online. The clinical description and testing indications for multiple endocrine neoplasia type 2 are as follows:

Germline testing for *RET* mutations is indicated for any individual with a diagnosis of primary C-cell hyperplasia or medullary thyroid cancer. Multiple endocrine neoplasia type 2 (MEN2) should be also suspected in any individual with pheochromocytoma or parathyroid adenoma/hyperplasia. GeneReviews also notes that pheochromocytomas in individuals with MEN2 are almost always adrenal.

National Comprehensive Cancer Network (NCCN)

NCCN Neuroendocrine and Adrenal Tumors guidelines (1.2023) also recommends MEN2 testing when there is clinical suspicion of MEN2 due to the presence of medullary thyroid cancer or other combination of MEN2-related features. GeneReviews also states that genetic testing is indicated for a first degree relative meeting this criteria, where the relative is not available for testing. (p. NE-E 3 of 8)

NEVOID BASAL CELL CARCINOMA SYNDROME (aka Gorlin syndrome)

***PTCH1* and/or *SUFU* Targeted Variant Analysis**

GeneReviews: Nevoid Basal Cell Carcinoma Syndrome

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online.

GeneReviews states that it is appropriate to evaluate apparently asymptomatic older and younger at-risk relatives (including children) of an affected individual in order to identify as early as possible those who would benefit from surveillance for complications of NBCCS (most notably medulloblastoma in children and jaw cysts and BCCs in adults) and avoidance of x-rays and sun exposure. Evaluations can include molecular genetic testing if the pathogenic variant in the family is known.

***PTCH1* and/or *SUFU* Sequencing and/or Deletion/Duplication Analysis**

GeneReviews: Nevoid Basal Cell Carcinoma Syndrome

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online. The clinical

description and testing indications for nevoid basal cell carcinoma syndrome/Gorlin syndrome are as follows:

Nevoid basal cell carcinoma syndrome (NBCCS) should be suspected in individuals with the following findings, which constitute major or minor diagnostic criteria. The diagnosis of NBCCS is established in a proband with either:

- Two major diagnostic criteria and one minor diagnostic criterion, **OR**
- One major and three minor diagnostic criteria

Major criteria

- Lamellar (sheet-like) calcification of the falx or clear evidence of calcification in an individual younger than age 20 years. Falx calcification is nearly always present and is visible on anteroposterior (AP) x-rays of the skull after age 20 years (see Notes regarding radiographs).
- Jaw keratocyst. Odontogenic keratocyst histologically; seen on orthopantomogram as an area of translucency
- Palmar/plantar pits (at least 2); particularly useful in diagnosis and more pronounced when the hands and feet are soaked in warm water for up to ten minutes. Pits may appear as white "punched-out" or pink "pin-prick" lesions.
- Multiple basal cell carcinomas (BCCs) (more than 5 in a lifetime) or a BCC before age 30 years. Provision needs to be made for decreased risk of BCC in individuals with dark skin and increased risk in those with light skin living in hot sunny climates, particularly those with type 1 Celtic skin and red hair, and of this group, particularly those with the common MC1R variant (rs1805007), which can modify age of onset for NBCCS.
- First-degree relative with NBCCS

Minor criteria

- Childhood medulloblastoma (also called primitive neuroectodermal tumor)
Lympho-mesenteric or pleural cysts
- Macrocephaly (OFC greater than 97th centile)
- Cleft lip/palate
- Vertebral/rib anomalies observed on chest x-ray and/or spinal x-ray (see Notes regarding radiographs): bifid/splayed/extra ribs; bifid vertebrae
- Preaxial or postaxial polydactyly
- Ovarian/cardiac fibromas
- Ocular anomalies (e.g., cataract, developmental defects, and pigmentary changes of the retinal epithelium).

HEREDITARY PARAGANGLIOMA/PHEOCHROMOCYTOMA SYNDROME (PGL/PCC)

MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, or TMEM127 Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Kidney Cancer (1.2024) include Hereditary paraganglioma/pheochromocytoma (PGL/PCC) syndrome in their overview of hereditary renal cell carcinoma syndromes, and state that this testing is indicated for an individual with a close blood relative with a known pathogenic/likely pathogenic variant. (p. HERED-RCC-1 and HERED-RCC-2)

MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, or TMEM127 Sequencing and/or Deletion/Duplication Analysis

GeneReviews: Hereditary Paraganglioma-Pheochromocytoma Syndromes

GeneReviews is an expert-authored review of current literature on a genetic disease, and goes through a rigorous editing and peer review process before being published online. The clinical description and testing indications for hereditary paraganglioma-pheochromocytoma syndromes are as follows:

Hereditary paraganglioma-pheochromocytoma (PGL/PCC) syndromes should be suspected in any individual with a paraganglioma or pheochromocytoma. Other tumors associated with these conditions are gastrointestinal stromal tumors (GIST), pulmonary chondromas, and renal clear cell carcinoma. In addition, individuals with a family history of paraganglioma or pheochromocytoma should also be suspected to have hereditary paraganglioma-pheochromocytoma syndromes.

The diagnosis of hereditary PGL/PCC should be strongly suspected in an individual with multiple, multifocal, recurrent, or early-onset paraganglioma or pheochromocytoma and/or a family history of paraganglioma or pheochromocytoma.

PEUTZ-JEGHERS SYNDROME (PJS)

STK11 Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline testing criteria for the evaluation of Lynch syndrome. NCCN recommends analysis of *STK11* in individuals with a family history of PJS. (p. PJS-1)

Additionally, NCCN states that tumor testing can be complementary to germline testing and can assist in interpretation of results. Although germline origin can sometimes be inferred with a high degree of confidence, confirmatory germline testing is indicated for pathogenic/likely pathogenic variants with a reasonable clinical suspicion of being a germline origin (based on patient/family history or clinical characteristics, presence of founder mutation, and in some cases variant allele frequency). (p. HRS-A 4 of 7)

***STK11* Sequencing and/or Deletion/Duplication Analysis**

National Comprehensive Cancer Network (NCCN)

NCCN Genetic/Familial High-Risk Assessment: Colorectal guidelines (1.2023) outline clinical criteria for PJS genetic testing in individuals with a personal and/or family history suggestive of PJS, as a majority of cases occur due to pathogenic variants in the *STK11* (*LKB1*) gene. These criteria include: two or more PJS-type hamartomas in the GI tract, hyperpigmentation in mucocutaneous membranes (such as the mouth, lips, nose, eyes, genitals, or fingers) and a family history of PJS. (p. PJS-1)

RETINOBLASTOMA

***RBI* Targeted Variant Analysis**

The AAOOP with support of the American Association for Pediatric Ophthalmology and Strabismus and the American Academy of Pediatrics (AAP) developed expert consensus guidelines for children at risk for development of retinoblastoma that included the following recommendations:

Genetic counseling and testing clarify the risk for retinoblastoma in children with a family history of the disease and improve outcomes at reduced cost, justifying making testing available to all patients with a personal or family history of retinoblastoma. Genetic evaluation should be initiated whether the affected relative demonstrated unilateral or bilateral disease because both have a substantial risk of being heritable (grade C). (p. 456)

***RBI* Sequencing and/or Deletion/Duplication Analysis**

American Association of Ophthalmic Oncologists and Pathologists (AAOOP)

The AAOOP with support of the American Association for Pediatric Ophthalmology and Strabismus and the American Academy of Pediatrics (AAP) developed expert consensus guidelines for children at risk for development of retinoblastoma that included the following recommendations:

Genetic counseling and testing clarify the risk for retinoblastoma in children with a family history of the disease and improve outcomes at reduced cost, justifying making testing available to all patients with a personal or family history of retinoblastoma. Genetic evaluation should be initiated whether the affected relative demonstrated unilateral or bilateral disease because both have a substantial risk of being heritable (grade C). (p. 456)

VON HIPPEL-LINDAU SYNDROME (VHL)

VHL Targeted Variant Analysis

National Comprehensive Cancer Network (NCCN)

NCCN guidelines for Kidney Cancer (1.2024) include von Hippel-Lindau (VHL) syndrome in their overview of hereditary renal cell carcinoma syndromes, and state that this testing is indicated for an individual with a close blood relative with a known pathogenic/likely pathogenic variant. (p. HERED-RCC-1 and HERED-RCC-2)

VHL Sequencing and/or Deletion/Duplication Analysis

National Comprehensive Cancer Network (NCCN)

NCCN Kidney Cancer guidelines (1.2024) outline clinical features seen in Von Hippel-Lindau syndrome including: hemangioblastomas (in the retina, spine, or brain), clear cell RCC (diagnosed before age 40 years or multiple/bilateral RCC diagnosed at any age), pheochromocytomas, paragangliomas (in the abdomen, thorax, or neck), retinal angiomas, endolymphatic sac tumors, epididymal or broad ligament papillary cystadenomas, multiple pancreatic serous cystadenomas, pancreatic neuroendocrine tumors, or multiple cysts in the pancreas. While these clinical features are categorized within the categories “major” and “minor,” the NCCN guidelines do not provide a scoring system required for patients to meet testing criteria. (p. HERED-RCC-A)

[back to top](#)

Reviews, Revisions, and Approvals	Revision Date	Approval Date
Policy developed.	03/23	03/23
Semi-annual review. Updated title to reflect V1.2024 version. Overview, coding, reference-table, background and references updated. Throughout policy: replaced “coverage criteria” with “criteria. For Overview: added “Of note, the National Society of Genetic Counselors...”. For Policy Reference Table; under Pre-Cancer Hereditary Cancer Susceptibility Panels: removed “Breast and GYN Cancers Panel (Invitae)””; under Hereditary Breast Cancer Susceptibility Panels: added “VistaSeg” and “Fulgent Genetics” and “part of Exact Sciences” and “plus PALB2” and “81307, 81321, 81351”; under Hereditary GI/Colon Cancer Panel Tests: added “0162U”; under Hereditary Pancreatic Cancer Susceptibility Panels: removed “Primary Panel”; under Hereditary Polyposis Panels: added “part of Exact Sciences”; under BRCA1 and BRCA2 Sequencing and/or Deletion/Duplication Analysis: removed “Breast and Ovarian Cancer Panel” and replaced with “BRCA1/2 Panel”; under PALB2 Targeted Variant Analysis: removed “Mutation Tests...” and replaced with “Variant (GeneDx)”; under PALB2 Sequencing and/or Deletion/Duplication Analysis: added “(Quest)”; under ATM or CHEK2 Targeted Variant Analysis: removed “Targeted Variants” and replaced with “Targeted Variant- Single Test” and removed “PreventionGenetics” and replaced with “(GeneDx)”; under ATM or CHEK2 Sequencing...added “part of Exact Sciences”; under MLH1, MSH2...replaced “Mutation Tests” with “Variant”; removed “Mutation Analysis” and replaced with “Variant (GeneDx)”; removed “81403” and replaced with “81479”; for FLCN Targeted Variant Analysis: removed “Targeted Variant” and added “Targeted Variant-Single Test (GeneDx)” removed “(Prevention Genetics); under PTEN Targeted Variant Analysis: removed “Targeted Variant” and replaced with “Targeted Variant-Single Test (GeneDx)”; under PTEN Sequencing and/or Deletion/Duplication Analysis: removed “Genomic Unity PTEN Analysis (Variantx Inc) and removed “0235U”; under Familial Adenomatous Polyposis...added “Adenomatous Polyposis Conditions” and “and MUTYH-Associated Polyposis Syndrome (MAP)”; under APC Sequencing and/or Deletion/Duplication Analysis: added “+RNAInsight...” and added “0157U”; under MUTYH Sequencing...added “MUTYH Full Gene...” and added “81406, 81479”; under CDKN2A Targeted Variant Analysis: removed “Targeted Variant” and replaced with “Targeted Variant-Single Test (GeneDX)” and removed “81403” and replaced with “81479”; under CDH1 Targeted Variant Analysis: removed “Targeted Variant” and replaced with “Targeted Variant-Single Test (GeneDX)” and removed “81403” and replaced with “81479”; under SMAD4 and/or BMPR1A Targeted Variant Analysis: added “part of Exact Sciences”; under FH Targeted Variant Analysis: removed “Miraca”, added “LLC” and removed “Laboratories”; under FH Sequencing and/or Deletion/Duplication Analysis: removed “FH Sequencint Tests...”; under TP53 Targeted Variant Analysis: removed “Targeted Variant” and replaced with “Targeted Variant-Single Test (GeneDX)”; under TP53 Sequencing and/or Duplication/Deletion Analysis: added “TP53 Full Gene Sequencing and Deletion/Duplication” and removed “Li Fraumeni Syndrome Test”; under MEN1 Targeted Variant Analysis: removed “Targeted Variant: MEN1 (Prevention Genetics)” and replaced with “Targeted Variant-Single Test (GeneDX)”	10/23	10/23

Reviews, Revisions, and Approvals	Revision Date	Approval Date
and removed “81403” and replaced with “81479”; for RET Targeted Variant Analysis: removed “Targeted Variant: RET (Prevention Genetics)” and replaced with “Targeted Variant-Single Test (GeneDX)” and removed “81405”; removed MUTYH-associated Polyposis (MAP)...”; under MAX, SDHA...: added “SDHB, SDHD...”; and replaced “81403” with “81479”; under STK11 Targeted Variant Analysis: added “-Single Test (GeneDx) PreventionGenetics”; under VHL Targeted Variant Analysis: removed “Miraca”, added “, LLC” and removed “Laboratories”. For Hereditary Breast Cancer Susceptibility Panels: under I. added “81307, 81321, 81351” and removed “0102U”; under I.A.1. removed “has a personal history...”; under I.B. removed “The member/enrollee has a probability...” and added “meets sequencing and/or...”; under II.A. replaced “all” with “any”; under II.B. removed “decisions”; under III. Added “81307, 81321, 81351” and removed “0102U”; removed “IV. Hereditary breast cancer...”. For Hereditary GI/Colon Cancer Panel Tests: under I. removed “0130U”; under I.2.a. removed “The member/enrollee’s tumor has deficient...”; under I.C. added “ant TP53,”; under II. removed “0130U”; under III. Added “0162U”. For Hereditary Gastric Cancer Panels: under I. and II. added “81201, 81203, 81404, 81405, 81406, 81408”. For Hereditary Pancreatic Cancer Susceptibility Panels: under I. added “81201” and “81351, 81433”; under II. added “81201,” and “81351, 81433”. For Hereditary Polyposis Panels: under I.A. removed “at least one of the following.”; under I.A.1. added “Adenomatous Polyposis Conditions”. For Hereditary Prostate Cancer Susceptibility Panels: under I. removed “0133U”; under I.B. added “The patient has a personal history...”; added I.C. “A personal history of prostate...”; under II. removed “0133U”. For BRCA1 and BRCA2 Sequencing and/or Deletion/Duplication Analysis: under I. removed “0138U”; added I.A.g. “Multiple primary breast cancers...”; under I.A.2.c. removed “Multiple primary breast cancers...”; under I.A.3. removed “meet any of the above criteria” and added “have a personal history of...”; under I.A.4. removed “decisions”; under I.A.6. removed “member/enrollee has a probability of greater than 5%... and added “member/enrollee’s probability of having” and added “is greater than 5%”; under II. removed “0138U”. For PALB2 Sequencing and/or Deletion/Duplication Analysis: under I. removed “0137U”; under I.A.1. removed breast cancer AND”; under I.A.1.a. replaced “Female” with “Male” and removed “diagnosed at age 50 years...”; under I.A.1.b. removed “male” and added “Triple-negative”; under I.A.1.c. removed “Ashkenazi Jewish...”; under I.A.1.c. removed “Triple negative breast” and added “Epithelial ovarian cancer...” under I.A.1.d. added “Pancreatic”; removed I.A.1.f. “Epithelial ovarian cancer...” removed I.A.1.g. “Pancreatic cancer...”; under I.A.2. removed “At least one close relative” and added “The member/enrollee has a personal history...”; under I.A.2.c. added “One or more close relatives”; under I.A.3. removed “meet the above criteria” and added “have a personal history of...”; under I.A.5. removed “member/enrollee has a probability of greater...” and added “member/enrollee’s probability of having” and added “is greater than 5%”. For ATM AND/OR CHEK2 Gene Testing: replaced “81403” with “81479” throughout. For Lynch Syndrome/Hereditary Nonpolyposis Colorectal Cancer (HNPCC) Testing: : replaced “81403” with “81479” throughout. For MLH1, MSH2, MSH6, PMS2, or EPCAM Sequencing and/or Deletion/Duplication Analysis: under		

Reviews, Revisions, and Approvals	Revision Date	Approval Date
I.A. removed “(i.e., colorectal, endometrial...”; under I.B. removed “colorectal cancer...” and added “Lynch syndrome...”; under I.B.2. removed (i.e., colorectal, endometrial...”removed I.B.3. “Diagnosed at any age...” removed I.B.4. “Diagnosed at any age...”; removed I.C. “The member/enrollee has a family history...”; added I.C.3. “Diagnosed at any age...”; added I.C.4. “Diagnosed at any age...”. For FLCN Sequencing and/or Deletion/Duplication Analysis: under I.A. added “any of the following...”; removed I.A.2. “Two of more of the following”; under I.A.5. removed “histology” and added “clear cell...”; added I.A.6. “Onocytoma, OR”; added I.A.7. “Angiomyolipoma”. For PTEN Sequencing and/or Deletion/Duplication Analysis: under I. removed “0235U”; removed “I.A.2. “Meets clinical criteria...”; and added I.A.2. “Autism-spectrum disorder...”. For Adenomatous Polyposis Conditions...: removed “Familial” from the title; under I. added “APC (81202)...”; added “81401, 81403”, removed “familial...” and added “adenomatous polyposis”, removed “(FAP)”; added “testing”; under I.A. and I.B. added “or MUTYH”; under APC and/or MUTYH Sequencing...: under I. removed “for familial” and added “and/or MUTYH sequencing...”; under I.A.1. replaced “20” with “10”; under I.A.2. removed “Multifocal/bilateral...” and added “congenital”; under I.A.3. added “Desmoid tumor...”; added II. APC sequencing...”; added III. “APC mRNA sequencing analysis...”. For Familial Atypical Multiple Mole Melanoma (FAMMM) Syndrome: under I. and II. replaced “81403” with “81479”. For Hereditary Diffuse Gastric Cancer: under I. and II. replaced “81403” with “81479”. For CDH1 Sequencing and/or Deletion/Duplication Analysis: under I.B.7. removed “The member/enrollee has a personal history...” and added “Two cases of lobular...”. For TP53 Sequencing and/or Deletion/Duplication Analysis: under I.D.3.b. removed “A member/enrollee has a diagnosis...”. For Multiple Endocrine Neoplasia Type 1 (MEN1): under I. and II. replaced “81403” with “81479”. For MEN1 Sequencing and/or Deletion/Duplication Analysis: under I.B. removed “diagnosis of cancer with a pathogenic...” and added “personal history of at least one...”. For Multiple Endocrine Neoplasia Type 2 (MEN2): under I. and II. replaced “81405” with “81404”. For Nevroid Basal Cell Carcinoma Syndrome (NBCCS) (aka Gorlin syndrome): under I. removed “MUTYH” and added “PTCH1 or SUFU”; removed “81403, 81404...” and added “81479...”; under I.A. replaced “blood relative” with “close relative” and removed “MUTYH” and added “PTCH1 or SUFU”; under I.B. removed “MUTYH” and added “PTCH1 or SUFU”; for II. removed “MUTYH” and added “PTCH1 or SUFU” and removed “81403, 81404...” and added “81479...”. For PTCH1 and SUFU Sequencing and/or Deletion/Duplication Analysis: under I. removed removed “MUTYH” and added “PTCH1 or SUFU”;removed “81406”; removed “MYH associated polyposis...”. Removed Nevroid Basal Cell Carcinoma Syndrome (NBCCS) (aka Gorlin syndrome). For Peutz-Jeghers Syndrome (PJS): under I. and II. replaced “81403” with “81479”. For Retinoblastoma; RB1 Sequencing and/or Deletion/Duplication Analysis: under I.B. removed “and has not previously undergone RB1 sequencing...”. For Notes and Definitions: added “11. Adenomatous polyposis...”; added “12. Lynch Syndrome related cancer...”. For Background and Rationale: removed “NCCN guidelines...”; removed “or a pathogenic variant with uncertain clinical management...”; added “in a well established gene...”; added		

Reviews, Revisions, and Approvals	Revision Date	Approval Date
“NCCN Guidelines...”; for Hereditary GI/Colon Cancer Panel Tests: removed “multigene panel testing...” and added “assessment for hereditary...”; added “history of”; removed “cancer has a known...”; removed “HRS” and added “LS-1”; removed “Lynch syndrome related...”; added “NCCN also states that the minimum...”; for Hereditary Pancreatic Cancer Panels: replaced “2.2022” with “1.2023”; for Hereditary Prostate Cancer Susceptibility Panels: added “NCCN Prostate Cancer guidelines...” and added “, triple-negative breast cancer...”; for BRCA1 and BRCA2 Sequencing and/or Deletion/Duplication Analysis: removed “American Society of Clinical Oncology (ASCO)”; for MLH1, MSH2, MSH6, PMS2, or EPCAM Sequencing and/or Deletion/Duplication Analysis: replaced “colorectal or endometrial” with “Lynch Syndrome...”; removed “including greater than...” and added “one of whom was diagnosed...”; added “An individual with a personal history of CRC...”; added “NCCN states that the minimum...”; for BAP1 Sequencing and/or Deletion/Duplication Analysis: removed “In addition to BAP1...”; added “BAP1-TBDS...”; added “*Excluding”; added “In addition to BAP1...”; for FLCN Sequencing and/or Deletion/Duplication Analysis: added “Commonly seen histologies...”; added “Identification of a heterozygous...”; for PTEN Sequencing and/or Deletion/Duplication Analysis: removed “or” multiple times throughout; added “*Revised Clinical Diagnostic Criteria...”; for Adenomatous Polyposis Conditions...: added “Of note, NCCN recommends...”; for APC and/or MUTYH Sequencing and/or Deletion/Duplication Analysis: removed “clinical criteria for the genetic testing...”; added “Adenomatous Polyposis testing criteria”; added “The guidelines also note...”; for Multiple Endocrine Neoplasia Type 1 (MEN1): removed “states that testing is recommended...” and added “recommends genetic risk evaluation...”; for MEN1 Sequencing and/or Deletion/Duplication Analysis: removed “be evaluated...” and added “or 1 AND a family history...”; removed “and parathyroidism”; added “Primary hyperparathyroidism”; removed MUTYH-Associated Polyposis (MAP)...; for PTCH1 and SUFU Sequencing and/or Deletion/Duplication Analysis: added “The diagnosis of NBCCS...”; removed “The diagnosis of NBCCS...”; for MAX, SDHA, SDHAF2, SDHB, SDHC, SDHD, or TMEM127 Sequencing and/or Deletion/Duplication Analysis: added “The diagnosis of hereditary...”.		
In hereditary breast cancer susceptibility panel criteria, changed the “or” after I.B.2. to “and.” In PTCH1 and SUFU Sequencing and/or Deletion/Duplication Analysis criteria, moved indication I.1.f)1)- 9) to I.A.2.a)-i); after criteria newly numbered as I.A.3)e), changed the “or” to “and”; moved indications previously listed as I.A.3.f)1)- 9) to new number I.A.4.a)-i).	01/24	01/24

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[back to top](#)

Important Reminder

This clinical policy has been developed by appropriately experienced and licensed health care professionals based on a review and consideration of currently available generally accepted standards of medical practice; peer-reviewed medical literature; government agency/program approval status; evidence-based guidelines and positions of leading national health professional organizations; views of physicians practicing in relevant clinical areas affected by this clinical policy; and other available clinical information. The Health Plan makes no representations and

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The purpose of this clinical policy is to provide a guide to medical necessity, which is a component of the guidelines used to assist in making coverage decisions and administering benefits. It does not constitute a contract or guarantee regarding payment or results. Coverage decisions and the administration of benefits are subject to all terms, conditions, exclusions and limitations of the coverage documents (e.g., evidence of coverage, certificate of coverage, policy, contract of insurance, etc.), as well as to state and federal requirements and applicable Health Plan-level administrative policies and procedures.

This clinical policy is effective as of the date determined by the Health Plan. The date of posting may not be the effective date of this clinical policy. This clinical policy may be subject to applicable legal and regulatory requirements relating to provider notification. If there is a discrepancy between the effective date of this clinical policy and any applicable legal or regulatory requirement, the requirements of law and regulation shall govern. The Health Plan retains the right to change, amend or withdraw this clinical policy, and additional clinical policies may be developed and adopted as needed, at any time.

This clinical policy does not constitute medical advice, medical treatment or medical care. It is not intended to dictate to providers how to practice medicine. Providers are expected to exercise professional medical judgment in providing the most appropriate care, and are solely responsible for the medical advice and treatment of members/enrollees. This clinical policy is not intended to recommend treatment for members/enrollees. Members/enrollees should consult with their treating physician in connection with diagnosis and treatment decisions.

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Note: For Medicaid members/enrollees, when state Medicaid coverage provisions conflict with the coverage provisions in this clinical policy, state Medicaid coverage provisions take

precedence. Please refer to the state Medicaid manual for any coverage provisions pertaining to this clinical policy.

Note: For Medicare members/enrollees, to ensure consistency with the Medicare National Coverage Determinations (NCD) and Local Coverage Determinations (LCD), all applicable NCDs, LCDs, and Medicare Coverage Articles should be reviewed prior to applying the criteria set forth in this clinical policy. Refer to the CMS website at <http://www.cms.gov> for additional information.

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