## **Helix Lynch Syndrome Panel**



Item	Description		
Test Name	Helix Lynch Syndrome Panel		
Test Type	Hereditary Cancer		
Catalog Number	LYNS1		
Procedure Code	79570-8 (LOINC)		
Test Description	This panel evaluates 5 genes associated with Lynch Syndrome and constitutional mismatch repair deficiency syndrome (CMMR-D).		
Genes Tested	EPCAM, MLH1, MSH2, MSH6, PMS2		
Genetics Information	This test utilizes next-generation sequencing to detect single nucleotide variants, insertions and deletions up to 20 bp, and copy number variants in genes associated with Lynch Syndrome and constitutional mismatch repair deficiency syndrome (CMMR-D).		
Indications For Testing	A relevant personal and/or family history suggestive of a hereditary form of cancer.		
Clinical Descriptions	This panel includes genes that have an established association with Lynch Syndrome and constitutional mismatch repair deficiency syndrome (CMMR-D).  The genes on this panel were specifically selected for their established association with hereditary cancer predisposition. Identification of a pathogenic variant may facilitate early and more frequent screening to help detect cancer at a more treatable stage. Certain medications and preventive surgeries can help to reduce the risk of developing cancer. Identification of a pathogenic variant also helps identify at-risk family members, who can pursue genetic testing and preventive measures.  The genes on this panel are associated with conditions that have autosomal dominant and/or autosomal recessive inheritance. Note that some of these genes may also be associated with other unrelated conditions; this means that when undergoing this test, there is a possibility of incidentally detecting carrier status for, or predisposition to, one of these conditions.  Analyzing a wide range of genes in a single test can provide an efficient, cost-effective method of testing for several hereditary cancer conditions. This approach increases the chance of identifying the underlying diagnosis responsible for an individual's or family's cancer predisposition.		
Conditions	Constitutional mismatch repair deficiency (MLH1, MSH2, MSH6, PMS2 and EPCAM) Lynch syndrome (MLH1, MSH2, MSH6, PMS2 and EPCAM) Autosomal recessive Congenital diarrhea 5 with tufting enteropathy (EPCAM)		
Interpretation	All detected variants are evaluated according to American College of Medical Genetics and Genomics recommendations. Variants are classified based on known, predicted, or possible pathogenicity and reported with interpretive comments detailing their potential or known significance.		

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Reclassification Of Variants	Helix does not systematically review their variant database looking for classification changes. Helix will review the classification of previously reported variants upon request of the ordering physician/provider. Ordering physicians/providers may contact Helix Customer Support or their Dedicated Advisor and request a review of the variant classification to be performed. At the discretion of the laboratory director, the frequency of reclassification requests may be limited to once per year, no earlier than 12 months after initial variant interpretation has been performed.		
Variant Evaluation	Variant classification is performed using the guidelines set forth by the American College of Medical Genetics and Genomics and the Association for Molecular Pathology, with modifications as suggested by domain specific Expert Panels of the Clinical genome Resource (ClinGen) when available. Variant pathogenicity is categorized as benign, likely benign, variant of uncertain significance (VUS), likely pathogenic, or pathogenic.		
Turnaround Time	7 to 24 days		
Available In NY State	No		
Test Classification	This test was developed, and its performance characteristics determined, by Helix, Inc. in a manner consistent with CLIA requirements. This test has not been cleared or approved by the US Food and Drug Administration.		
Performing Laboratory Information			
Regulatory Information	CLIA Complexity: High Test Classification: Non-Waived/ Laboratory Developed Test		
CLIA Category	Chemistry / Routine Chemistry		

# Methods & Limitations for Helix Lynch Syndrome Panel



Extracted DNA is enriched for targeted regions and then sequenced using the Helix Exome+ (R) assay on an Illumina DNA sequencing system. Data is then aligned to a modified version of GRCh38 and all genes are analyzed using the MANE transcript and MANE Plus Clinical transcript, when available. Small variant calling is completed using a customized version of Sentieon's DNAseq software, augmented by a proprietary small variant caller for difficult variants. Copy number variants (CNVs) are then called using a proprietary bioinformatics pipeline based on depth analysis with a comparison to similarly sequenced samples. Reportable variants in *PMS2* exons 12-15 are confirmed by PacBio long reads. The *MSH2* Boland inversion (exons 1-7) is detected by identifying discordant read-pairs spanning the presumed breakpoint. Interpretation is based upon guidelines published by the American College of Medical Genetics and Genomics (ACMG) and the Association for Molecular Pathology (AMP) or their modification by ClinGen Variant Curation Expert Panels when available. Interpretation is limited to the transcripts indicated on the report, +/- 10 bp into intronic regions, except as noted below. Helix variant classifications include pathogenic, likely pathogenic, variant of uncertain significance (VUS), likely benign, and benign. Variants classified as pathogenic, likely pathogenic, or VUS are included in the report. All reported variants (except for VUSs with limited evidence of pathogenicity) are confirmed through secondary manual inspection of DNA sequence data or orthogonal testing. Benign and likely benign variants are not reported but are available upon request. Risk estimations and management guidelines included in this report are based on analysis of primary literature and recommendations of applicable professional societies, and should be regarded as approximations.

Based on validation studies, this assay delivers > 99% sensitivity and specificity for single nucleotide variants and insertions and deletions (indels) up to 20 bp. Larger indels and complex variants are also reported but sensitivity may be reduced. Based on validation studies, this assay delivers > 99% sensitivity to multi-exon CNVs and > 90% sensitivity to single-exon CNVs. This test may not detect variants in challenging regions (such as short tandem repeats, homopolymer runs, and segment duplications), sub-exonic CNVs, chromosomal aneuploidy, or variants in the presence of mosaicism. Phasing will be attempted and reported, when possible. Structural rearrangements such as inversions, translocations, and gene conversions are not tested in this assay unless explicitly indicated. Additionally, deep intronic, promoter, and enhancer regions may not be covered. It is important to note that this assay cannot detect all variants known to increase disease risk, and that a negative result does not guarantee that the tested individual does not carry a rare, undetectable variant in genes analyzed. Any potential incidental findings outside of these genes and conditions will not be identified, nor reported. The results of a genetic test may be influenced by various factors, including bone marrow transplantation, blood transfusions, or in rare cases, hematolymphoid neoplasms.

### **Gene Specific Notes:**

*BRCA1*: sequencing analysis extends to CDS +/-20 bp; *BRCA2*: sequencing analysis extends to CDS +/-20 bp; *EPCAM*: analysis is limited to CNV of exons 8-9; *HOXB13*: analysis is limited to the c.251G>A (p.Gly84Glu) variant; *MLH1*: analysis includes CNV of the promoter; *MSH2*: analysis includes detection of the Boland inversion (inversion of exons 1-7) and detection of c.942+3A>T; *TP53*: analysis includes CNV of the promoter.

#### Disclaimer:

This test was developed and validated by Helix, Inc. This test has not been cleared or approved by the United States Food and Drug Administration (FDA). The Helix laboratory is accredited by the College of American Pathologists (CAP) and certified under the Clinical Laboratory Improvement Amendments (CLIA #: 05D2117342) to perform high-complexity clinical tests. This test is used for clinical purposes. It should not be regarded as investigational or for research.

# Targeted Genes & Methodology for Helix Lynch Syndrome Panel



The following applies to the Helix Lynch Syndrome Panel. Testing is performed to evaluate for the presence of variants in coding regions and extending to +/- 10 base pairs of adjacent intronic sequences on either side of the coding exons of the genes analyzed. In addition, the analysis will cover select non-coding variants, as listed below. Next-generation sequencing is performed to test for the presence of small variants and copy number variants in the genes analyzed. Confirmation of select reportable variants may be performed by alternate methodologies based on internal laboratory criteria.

This list is current from November 2024 to the present. This document is intended to highlight additional evaluations for variants of high clinical interest as well as technical limitations. For questions regarding genes, reference transcripts, or specific regions covered, contact Helix Customer Service at (844) 211-2070.

Genomic Build: GRCh38 Catalog Number: LYNS1

Gene	Transcript	Additional Evaluations	Technical Limitations
EPCAM	NM_002354.3	_	Results limited to CNV and limited to exons 8 and 9
MLH1	NM_000249.4	Includes CNV detection in the promoter	-
MSH2	NM_000251.3	Includes detection of the exon 1-7 rearrangement known as the Boland Inversion and of c.942+3A>T	-
MSH6	NM_000179.3	-	-
PMS2	NM_000535.7	-	-