## Ambry Genetics 🛛 💍 LifeStrands

# Testing for Hereditary Breast and Gynecologic Cancer REFERENCE GUIDE

Genetic Testing For Hereditary Breast and Gynecologic Cancer is Recommended By: American Society of Breast Surgeons (ASBrS)<sup>1</sup>

Society of Gynecologic Oncology (SGO)<sup>2</sup> American College of Obstetricians and Gynecologists (ACOG)<sup>2</sup>

NCCN Clinical Practice Guidelines In Oncology (NCCN Guidelines®)<sup>3</sup>

## GUIDELINES RECOMMEND GENETIC TESTING IF YOUR PATIENT HAS A PERSONAL OR FAMILY HISTORY OF ANY OF THE FOLLOWING SIGNS\* OF HEREDITARY CANCER:



## **MULTIPLE CANCERS**

- > 2 or more primary cancers in the same person
- > 3 or more cancers on the same side of the family



**RARE CANCERS** 

- > Male breast
- > Ovarian
- > Pancreatic
- > Metastatic prostate



- > Breast cancer diagnosed before 45
- > Triple negative breast cancer before 60
- > Uterine cancer diagnosed before 50

#### ABNORMAL TUMOR SCREENING

 Tumors with microsatellite instability (MSI) or loss of immunohistochemical (IHC) staining

#### ANCESTRY

> Ashkenazi Jewish with breast cancer



### GENETIC TESTING RESULTS MAY ALSO INFORM TREATMENT RECOMMENDATIONS

> PARP inhibitor therapy for BRCA1 or BRCA2 positive patients with ovarian cancer<sup>4-12</sup>

> Immunotherapy for patients with a germline mutation in a mismatch repair gene13

## HEREDITARY BREAST AND GYNECOLOGIC CANCER TESTING OPTIONS



## **CancerNext**®

36-gene test covering a range of cancers; NCCN Guidelines® provide recommendations regarding risk management for most genes in the panel<sup>3</sup>



## **BRCANext**<sup>®</sup>

18-gene test for hereditary breast and gynecologic cancers; NCCN Guidelines provide recommendations regarding risk management for **all genes** in the panel<sup>3</sup>



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GENES

## **BRCANext-Expanded®**

23-gene test for hereditary breast and gynecologic cancers; NCCN Guidelines provide recommendations regarding risk management for most genes in the panel<sup>3</sup>

## **BRCAplus**<sup>®</sup>

Ambry Genetics

8-gene STAT, breast cancer test; NCCN Guidelines provide recommendations regarding risk management for **all genes** in the panel<sup>3</sup>

# Add +RNAinsight<sup>®</sup> to Find More Mutations, Decrease Variants of Unknown Significance, and Provide More Accurate Results<sup>14-16</sup>

+RNAinsight is Available With Any Hereditary Cancer Panel\*



## SAMPLE REQUIREMENTS

One kit that includes 1 EDTA tube (DNA) and 1 PAXgene<sup>®</sup> tube (RNA)



## NO IMPACT ON TURNAROUND TIME OR COST

\*Exception: BRCAplus®

## **DID YOU KNOW?**

Adding RNA genetic testing decreases variants of unknown significance in splice sites by 70% and increases diagnostic yield by 1.3%<sup>16</sup>

References

1. Manahan ER,et al. Ann Surg Oncol. 2019 Oct;26(10):3025-3031. 2. Obstetrics & Gynecology. 2017 Sep;130(3):e110-e126. 3. Referenced with permission from the NCCN Clinical Practice Guidelines in Oncology (NCCN Guidelines<sup>®</sup>) for Genetic/Familial High-Risk Assessment: Breast, Ovarian, and Pancreatic. V1.2023. © National Comprehensive Cancer Network, Inc. 2019. All rights reserved. Accessed December 29, 2022. To view the most recent and complete version of the guideline, go online to NCCN.org. NCCN makes no warranties of any kind whatsoever regarding their content, use or application and disclaims any responsibility for their application or use in any way. 4. Banerjee S & Kaye S. Curr Oncol Rep. 2011 Dec;13(6):442-9. 5. Burgess M & Puhalla S. Front Oncol. 2014 Feb 27;4:19. 6. Yamamoto KN et al. PLoS One. 2014 Aug 26;9(8): e105724. 7. Moore et al. NEJM 2018 Dec 27; 379(26):2495–505. 8. Ledermann, et al. Lancet Oncol. 2014;15(8):852-861. 9. Pujade-Lauraine, et al. Lancet Oncol. 2017;18:1274-1284. 10. Mirza, et al. N Engl J Med. 2016;375:2154-2164. 11. Coleman RL, et al. Lancet. 2017 Oct 28;390(10106):1949-1961. 12. Swisher, et al. Lancet Oncology 2017;18: 75-87. 13. Zhao P, Li et al. J Hematol Oncol 2019 May;12:54. 14. Landrith T et al. npj Precision Oncology. 2020. 15. Karam et al. JAMA Network Open. 2019. 16. Horton, Carolyn, et al. "Mutational and splicing landscape in a cohort of 43,000 patients tested for hereditary cancer." NPJ genomic medicine 7.1 (2022): 1-6.

#### **BREAST AND GYNECOLOGIC GENES AND ASSOCIATED RISKS**

GENE(S)	ASSOCIATED CANCERS AND RISKS
ATM*	Breast, pancreatic, prostate
BRCA1*	Female breast, male breast, ovarian, pancreatic, prostate
BRCA2*	Female breast, male breast, ovarian, pancreatic, melanoma, prostate
CDH1*	Diffuse gastric, female breast
CHEK2*	Breast, colorectal, prostate, other
PALB2*	Female breast, male breast, pancreatic, ovarian, prostate
PTEN*	Breast, thyroid, uterine, colorectal, renal, melanoma
TP53*	Breast, sarcoma, brain, adrenocortical, leukemia, other
BRIP1*	Ovarian
EPCAM	Colorectal, uterine, stomach, ovarian, other
MLH1	Colorectal, uterine, stomach, ovarian, other
MSH2	Colorectal, uterine, stomach , ovarian, other
MSH6	Colorectal, uterine, other
NBN*	
NF1*	Female breast, malignant nerve sheath tumors, PGL/PCC, brain, other
PMS2*	Colorectal, uterine, other
RAD51C*	Ovarian, female breast
RAD51D*	Ovarian, female breast
BARD1	Female breast
DICER1	Pleuropulmonary blastoma, cystic nephroma, ovarian sex-cord tumors, brain, other
RECQL	
SMARCA4	Ovarian (small cell carcinoma, hypercalcemic type), brain, other
STK11*	Colorectal, stomach, female breast, pancreatic, other
APC*	Colorectal, small bowel, stomach, pancreatic, thyroid, other
AXIN2*	Colorectal
BMPR1A*/ SMAD4*	Colorectal, stomach
CDK4	Melanoma
CDKN2A*	Melanoma, pancreatic
GREM1	Colorectal
HOXB13	Prostate
MSH3*	Biallelic mutations: colorectal
MUTYH*	Biallelic mutations: colorectal, duodenal
NTHL1*	Biallelic mutations: Colorectal, other
POLD1*/POLE*	Colorectal

#### CONTACT INFORMATION

BRCAplus o-

BRCANext o.

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