A Direct Path to Therapy in Ovarian Cancer

FoundationFocus™ CDx_{BRCA} is an FDA-approved companion diagnostic test for BRCA-mutated ovarian cancer to aid in identifying women for whom treatment with the PARP inhibitor Rubraca™ (rucaparib) may be considered.

More Patients on Therapy Sooner

1 in 4 ovarian cancer patients are BRCA+.

By testing for both germline and somatic BRCA1/2 mutations, FoundationFocus™ CDx_{BRCA} can help identify up to twice as many women\(^{1,4,5}\) who may benefit from Rubraca, a PARP inhibitor therapy. Conventional tests only look for germline alterations in blood or saliva. FoundationFocus CDx_{BRCA} analyzes ovarian cancer tumor tissue to detect BRCA1/2 mutation types associated with response to PARP inhibition.\(^1,4\)

Uncover More BRCA Mutations with a Single Tissue Test

FoundationFocus™ CDx_{BRCA} is an FDA-approved tissue-based next generation sequencing test. Alterations detected in tumor may include both somatic and germline.

Hybrid capture-based, comprehensive genomic profiling is used to achieve high, uniform depth of coverage of all coding exons, including splice sites, and select intronic regions of BRCA1 and BRCA2.
When to Test

FoundationFocus™ CDx_BRCA uses recent or archived tumor tissue specimens to find BRCA mutations that may be missed by other methods of testing.1-4 Consider testing ovarian cancer patients...

who have:
• No prior tissue or germline BRCA1/2 testing or
• Prior BRCA1/2 testing with germline BRCA1/2 (-) results

who would like to:
• Determine eligibility for Rubraca™ therapy as a treatment option

Simplified Sample Collection and Ordering

• No need to re-biopsy; archived specimens accepted. Select the most recent specimen that meets size and tumor content criteria
• DNA extracted from 1mm³ formalin-fixed, paraffin embedded (FFPE) ovarian cancer tumor tissue specimens, which are at least 20% of malignant origin
• 14-day turnaround time from time specimen is received
• Reports available via fax or online using FoundationICE®
• Patient financial assistance and case management services available through FoundationACCESS™
• Contact Client Services for a FoundationFocus CDx_BRCA Specimen Shipping Kit

Intended Use

The FoundationFocus™ CDx_BRCA assay detects sequence alterations in BRCA1 and BRCA2 genes. Results of the test are used as an aid in identifying ovarian cancer patients for whom treatment with Rubraca™ (rucaparib) is being considered. If a patient is positive for any of the deleterious alterations specified in the BRCA1/2 classification, the patient may be eligible for treatment with Rubraca. This assay is to be performed at Foundation Medicine, Inc., a single laboratory site located at 150 Second Street, Cambridge, MA 02141. Contraindication None. Warnings and Precautions BRCA1/2 alterations reported include somatic (not inherited) or germline (inherited) alterations; however, the test does not distinguish between germline and somatic alterations. The test does not provide information about susceptibility. Biopsy may pose a risk to the patient when archival tissue is not available for use with the assay. The patient’s physician should determine whether the patient is a candidate for biopsy. Limitations For in vitro diagnostic use. For professional use only. This test must be ordered by a qualified medical professional in accordance with clinical laboratory regulations. Limited performance characteristics of the test were evaluated for insertion alterations > 4 nucleotides and deletions > 10 nucleotides. Performance of the FoundationFocus CDx_BRCA was not established for insertions > 10 nucleotides, deletions > 12 nucleotides, alterations residing in polyC homopolymer runs, homozygous deletions or large rearrangements. Alterations in polyT homopolymer runs may not be reliably detected. Alterations detected at allele frequencies below the established limit of detection are not detected consistently. Information generated by this test is an aid in the identification of patients who are most likely to benefit from the therapeutic product. Decisions on patient care and treatment must be based on the independent medical judgment of the treating physician, taking into consideration all applicable information concerning the patient’s condition, such as patient and family history, physical examinations, information from other diagnostic tests, and patient preferences, in accordance with the standard of care in a given community. The test is intended to be performed at a single site on specific serial number-controlled instruments at Foundation Medicine, Inc. Rubraca™ (rucaparib) is a product of Clovis Oncology. For additional information on the assay and detailed performance specifications, refer to the complete FoundationFocus™ CDx_BRCA label at www.foundationmedicine.com/focus.