

# Test Requisition Form

Please fax to: (617) 418-2290 Email: client.services@foundationmedicine.com

All fields required | For more information or to order online, visit [www.foundationmedicine.com/genomic-testing/order](http://www.foundationmedicine.com/genomic-testing/order)

## Patient Demographics

Last Name \_\_\_\_\_ First Name \_\_\_\_\_ MI \_\_\_\_\_ Medical Record # \_\_\_\_\_ DOB (MM/DD/YYYY) \_\_\_\_\_ Sex  F  M  
Address \_\_\_\_\_ City/State/Postal Code \_\_\_\_\_ Country \_\_\_\_\_ Phone (primary) \_\_\_\_\_

## Treating Physician Information

Facility Name \_\_\_\_\_ Treating Physician (full legal name) \_\_\_\_\_  
Facility Address \_\_\_\_\_ City/State/Postal Code \_\_\_\_\_ Country \_\_\_\_\_  
Phone \_\_\_\_\_ Fax \_\_\_\_\_ Email \_\_\_\_\_ Account # \_\_\_\_\_

Additional Physician to be Copied (optional) Facility Name \_\_\_\_\_ Email \_\_\_\_\_ Fax \_\_\_\_\_

## Current Diagnosis/Patient History

Diagnosis:  NSCLC  Melanoma  Colorectal Carcinoma  Ovarian  Breast  Other \_\_\_\_\_

Disease Status (select all that apply):  Metastatic  Recurrent  Refractory  Relapse  None of these options

Additional Details \_\_\_\_\_ Stage \_\_\_\_\_ ICD Codes (only codes beginning C or D accepted) \_\_\_\_\_

Transplant Information \_\_\_\_\_ Targeted Therapies \_\_\_\_\_

Attachments:  Copy of recent pathology/cytology reports including (if available), CBC/differential, BMA differential, FAB classification.  
 Test results from all other Molecular Diagnostic Assays by FISH, IHC, or other genetic assays, e.g., ER, PR, HER2, EGFR, KRAS, etc.

## Test Selection | Select one

Genomic test	Description	Accepted Specimen Type	Genomic test	Description	Accepted Specimen Type
<input type="radio"/> <b>FoundationOne®CDx</b>	FDA-approved companion.....	FFPE Tissue	<input type="radio"/> <b>FoundationOne®Heme</b>	For hematologic.....	Peripheral Whole Blood, Bone malignancies & Marrow Aspirate, FFPE Tissue, sarcomas
<input type="checkbox"/> If tissue submitted does not meet the criteria for.....	Peripheral successful testing, reflex to FoundationOne®Liquid (option for mobile phlebotomy below)	Whole Blood	Additional Option: -----		
<input type="radio"/> <b>FoundationOne®Liquid</b>	Liquid biopsy for all.....	Peripheral solid tumors	<input type="radio"/> <b>IHC Testing PD-L1.....</b>		FFPE tissue
				(Scoring and clone utilization based on FDA-approved indications. See back of this document for information.)	

## Specimen Retrieval | Only one specimen can be tested per order

Submitting Pathologist Name \_\_\_\_\_ Pathology Lab Name \_\_\_\_\_ Phone \_\_\_\_\_ Fax \_\_\_\_\_ Email \_\_\_\_\_

Specific specimen requested  Let the submitting pathologist choose specimen

Date of Collection (MM/DD/YYYY) \_\_\_\_\_ Specimen ID \_\_\_\_\_ Specimen Site \_\_\_\_\_ Alternate Choice \_\_\_\_\_  
(FFPE or BMA) (optional)

**FFPE Tissue:**  **Peripheral Whole Blood:**  **Bone Marrow Aspirate/Extracted Nucleic Acid:**

I will arrange for specimen shipment  I will arrange for specimen shipment  Ordering Facility responsible for shipment

Contact the pathology lab to obtain specimen  Mobile Phlebotomy requested (see guidelines on website)

## Billing Information | Select one of the three payment options and complete all fields indicated

**Insurance (check one):**  Medicare  Medicare Advantage  Other Health Insurance Plan Name \_\_\_\_\_

Policy # \_\_\_\_\_ Group # \_\_\_\_\_ Prior Authorization # \_\_\_\_\_  ABN Attached

Patient status at time of collection:  Office (non-hospital)  Outpatient  Inpatient (requires discharge date MM/DD/YYYY): \_\_\_\_\_  
(required for all Medicare patients) OR  Not yet discharged

**Facility:** \_\_\_\_\_ Address \_\_\_\_\_  Same as Treating Physician

**Self-Pay:** Contact Name \_\_\_\_\_ Phone \_\_\_\_\_ Email \_\_\_\_\_

## Certificate of Medical Necessity/Consent/Test Authorization and Physician Signature

My signature constitutes a Certificate of Medical Necessity, certifies that this test information will inform the patient's ongoing treatment plan, and certifies that I am the patient's treating physician. I have explained to the patient the nature and purpose of the testing to be performed and have obtained informed consent, to the extent legally required, to permit Foundation Medicine to (a) perform the testing specified herein, (b) retain the test results for an indefinite period for internal quality assurance/operations purposes, (c) de-identify the test results and use or disclose such de-identified results for future unspecified research or other purposes, and (d) release the test results to the patient's third-party payer as needed for reimbursement purposes.

My signature also authorizes Foundation Medicine to select the most appropriate test (pursuant to Foundation Medicine's Change in Test Authorization Policy) based on requisition/pathology information.

Treating Physician Signature \_\_\_\_\_ Printed Name (Full legal name) \_\_\_\_\_ Date (MM/DD/YYYY) \_\_\_\_\_

## Technical Information

### FOUNDATIONONE® CDx

**Intended Use** FoundationOne®CDx is a next-generation sequencing based *in vitro* diagnostic device for detection of substitutions, insertion and deletion alterations, and copy number alterations in 324 genes and select gene rearrangements, as well as genomic signatures including microsatellite instability (MSI) and tumor mutational burden (TMB) using DNA isolated from formalin-fixed, paraffin-embedded (FFPE) tumor tissue specimens. For the complete Intended Use statement, including companion diagnostic indications, please see the FoundationOne CDx Technical Information page: [www.foundationmedicine.com/f1cdx](http://www.foundationmedicine.com/f1cdx).

### FOUNDATIONONE® HEME

**About the Test** FoundationOne®Heme is a comprehensive genomic profiling test for hematologic malignancies and sarcomas. The test is designed to provide physicians with clinically actionable information to help with diagnostic sub-classification, prognosis assessment, and targeted therapeutic selection. Test results provide information about clinically significant alterations, potential targeted therapies, available clinical trials, and quantitative markers that may support immunotherapy clinical trial enrollment. FoundationOne Heme is validated to detect all classes of genomic alterations in more than 400 cancer-related genes. In addition to DNA sequencing, FoundationOne Heme employs RNA sequencing across more than 250 genes to capture a broad range of gene fusions, common drivers of hematologic malignancies and sarcomas.

### FOUNDATIONONE® LIQUID

**About the Test** FoundationOne®Liquid is a blood-based circulating tumor DNA (ctDNA) liquid biopsy test for solid tumors that identifies clinically relevant genomic alterations and provides an assessment of high microsatellite instability, across 70 genes known to be drivers of cancer. This test can assist physicians in identifying treatment options by providing clinically actionable information relevant to diagnosis, risk-stratification and prognosis. Test results provide information about potential targeted therapies and/or clinical trials to better inform treatment decisions.

### IHC Testing

Foundation Medicine performs PD-L1 IHC testing utilizing the appropriate platform and clone which may be informed by FDA approved companion diagnostic status for the submitted tissue type and diagnosis. More information available at this web link: [www.foundationmedicine.com/genomic-testing/order](http://www.foundationmedicine.com/genomic-testing/order).

## Medicare Coverage Summary

Foundation Medicine tests may be covered by Original Medicare<sup>1</sup> and Medicare Advantage<sup>2</sup>.

TEST	CONDITIONS FOR MEDICARE COVERAGE	PATIENT COVERAGE CRITERIA
FoundationOne®CDx	Covered <sup>3</sup> if all patient coverage criteria are met. ABN required if patient does not meet the patient coverage criteria or if person ordering the test is not a treating physician <sup>4</sup> .	i) Patient has been diagnosed with a solid malignant neoplasm; <b>AND</b> ii) Patient has either recurrent, relapsed, refractory, metastatic, or advanced stages III or IV cancer (only requires one of these to be met); <b>AND</b> iii) <b>Either</b> Patient has not been previously tested using the same NGS test for the same primary diagnosis of cancer <b>OR</b> Patient is undergoing repeat testing using the same NGS test for a new primary cancer diagnosis made by the treating physician; <b>AND</b>
FoundationOne®Liquid	Coverage <sup>5</sup> may be available if all patient coverage criteria are met. ABN required if patient does not meet the patient coverage criteria, or if person ordering the test is not a treating physician <sup>4</sup> .	iv) Patient has decided to seek further cancer treatment (e.g., therapeutic chemotherapy)
FoundationOne®Heme	Not covered at this time. Foundation Medicine is working toward securing future coverage. ABN required for every case.	N/A

### References

- Medicare administered by federal government.
- Medicare administered by private insurers.
- Per the "Decision for Next Generation Sequencing (NGS) for Medicare Beneficiaries with Advanced cancer – CAG-00450N."
- A "treating physician" is a physician, as defined in §1861(r) of the Social Security Act, who furnishes a consultation or treats a beneficiary for a specific medical problem, and who uses the results of a diagnostic test in the management of the beneficiary's specific medical problem. More information is available at <https://www.cms.gov/Regulations-and-Guidance/Guidance/Transmittals/Downloads/R80BP.pdf>.
- National Government Services, the local Medicare Administrative Contractor with jurisdiction over testing performed by Foundation Medicine at its Cambridge, MA laboratory for Original Medicare beneficiaries, does not have a Local Coverage Determination (LCD) for liquid biopsy next generation sequencing >50 genes. Coverage is determined by National Government Services on a case-by-case basis.