

6. PATHOLOGY LABORATORY & PROCUREMENT SERVICES (*Indicates a required field if applicable to test order)

*Pathology Lab Name		Submitting Pathologist Name (optional)	
*Phone	Email (preferred)	Fax (if email not provided)	
*Specimen Retrieval Type		Physician is requesting a specific specimen (add specimen details below)	
*Specimen ID		*Date of Collection (MM/DD/YYYY)	*Specimen Site
*Alternate Specimen ID		*Date of Collection (MM/DD/YYYY)	*Alternate Specimen Site

7. FFPE BLOCK RETURN INFORMATION (*Indicates a required field if applicable to test order)

*Return Address			
*City	*State	*Postal Code	*Country
Email (preferred)	Phone (optional)	Fax (optional)	

8. RELEVANT CLINICAL HISTORY (All Required For Medical Coverage Determination)

a. Is a tissue specimen from a recent procedure available?	Yes	No
b. Tissue specimen is insufficient for testing or tissue testing resulted as a Quantity Not Sufficient (QNS)	Yes	No
c. Is the requested test assessing for tumor mutation burden (TMB) to identify if the patient is a candidate for checkpoint inhibitor immunotherapy?	Yes	No

9. FDA COMPANION DIAGNOSTIC INDICATIONS¹ FOR FOUNDATIONONE CDX AND FOUNDATIONONE LIQUID CDX* (*Required Section: Select or write in indication for testing)

TUMOR TYPES	BIOMARKERS ² (See complete gene list on our website)	FDA-APPROVED THERAPY ³ <small>Last Updated 10/25/2023, please use "If other" box below to include additional</small>
Solid tumors	TMB ≥ 10 mutations per megabase	Keytruda® (pembrolizumab)
	NTRK1/2/3 fusions	Vitrakvi® (larotrectinib) or Rozlytrek® (entrectinib)
	MSI-H	Keytruda® (pembrolizumab)
	RET	Retevmo (selpercatinib)
Non-Small Cell Lung Cancer (NSCLC)	EGFR exon 19 deletions and EGFR exon 21 L858R alterations	EGFR Tyrosine Kinase Inhibitors (TKI) approved by FDA ¹
	EGFR exon 20 T790M alterations	Tagrisso® (osimertinib)
	ALK rearrangements	Alecensa® (alectinib), Alunbrig® (brigatinib), Xalkori® (crizotinib), or Zykadia® (ceritinib)
	MET single nucleotide variants (SNVs) and indels that lead to MET exon 14 skipping	Tabrecta® (capmatinib)
	BRAF V600E	Tafinlar® (dabrafenib) in combination with Mekinist® (trametinib) or BRAFTOVI® (encorafenib) in combination with MEKTOVI® (binimetinib)
	EGFR exon 20 insertion mutations	EXKIVITY® (mobocertinib)
	ROS1 fusions	Rozlytrek® (entrectinib)
Melanoma	BRAF V600E	BRAF Inhibitors approved by FDA ¹
	BRAF V600E and V600K	Mekinist® (trametinib) or BRAF/MEK Inhibitor Combinations approved by FDA ¹
	BRAF V600 mutation-positive	Tecentriq® (atezolizumab) in combination with Cotellic® (cobimetinib) and Zelboraf® (vemurafenib)
Breast Cancer	ERBB2 (HER2) amplification	Herceptin® (trastuzumab), Kadcyla® (ado-trastuzumab-emtansine), or Perjeta® (pertuzumab)
	PIK3CA C420R, E542K, E545A, E545D [1635G>T only], E545G, E545K, Q546E, Q546R, H1047L, H1047R, and H1047Y alterations	Piqray® (alpelisib)
Colorectal Cancer	KRAS wild-type (absence of mutations in codons 12 and 13)	Erbix® (cetuximab)
	KRAS wild-type (absence of mutations in exons 2, 3 and 4) and NRAS wild-type (absence of mutations in exons 2, 3 and 4)	Vectibix® (panitumumab)
	BRAF V600E	BRAFTOVI® (encorafenib) in combination with cetuximab
Ovarian Cancer	BRCA1/2 alterations	Lynparza® (olaparib)
Cholangiocarcinoma	FGFR2 fusions and select rearrangements	Pemazyre™ (pemigatinib) or Truseltiq™ (infigratinib)
Prostate Cancer	Homologous Recombination Repair (HRR) gene (BRCA1, BRCA2, ATM, BARD1, BRIP1, CDK12, CHEK1, CHEK2, FANCL, PALB2, RADS1B, RADS1C, RADS1D and RADS4L) alterations	Lynparza® (olaparib)
	BRCA1/2 alterations	Rubraca® (rucaparib) or AKEEGATM (niraparib and abiraterone acetate dual action tablet)
If other indications for testing apply, please indicate here:		

10. OTHER INFORMATION

For information on ICD codes, visit this website: <https://icd10cmtool.cdc.gov/>

PORTFOLIO REFLEX OPTION:
If the reflex option is selected, we will proceed with the initial NGS test selected and if the specimen does not meet the criteria for successful testing, we will automatically reflex to the other test (in Section 5) and procure a new specimen. The failed test is not billed, and the successful test will be billed according to our standard practices. Please see foundationmedicine.com/order for more information.

11. PHYSICIAN CERTIFICATION OF MEDICAL NECESSITY AND CONSENT (*Indicates a required field)

My signature below certifies that (1) I am the patient's treating physician and am authorized under applicable law to order the tests on this test requisition, (2) each test ordered on this test requisition is medically necessary for the patient, (3) the patient has decided to seek further cancer treatment, (4) the results of each test will inform the patient's ongoing treatment plan, (5) I have explained to the patient the nature and purpose of each test to be performed pursuant to this test requisition, and the patient has had the opportunity to ask questions regarding each test and the collection, use, and disclosure of his/her samples and data, (6) I have obtained informed consent from the patient using the consent form available at <https://foundationmedicine.com/asset/patient-consent> to have each test performed, including the collection, use, and disclosure of his/her samples and data, and (7) I have informed the patient that he/she may receive a copy of the signed consent and have also included a signed copy in his/her medical record. I understand that Foundation Medicine may reach out to me to request a copy of the signed consent, in which case I will furnish Foundation Medicine a signed copy of the consent. * (or the patient's legal guardian or representative)

*Treating Physician Signature	*Printed Full Name (Full legal name)	*Date (MM/DD/YYYY)
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Notice for CA HCPs: Please review our privacy policy, available at <https://www.foundationmedicine.com/california-privacy-notice>, for more information about how we collect, use and disclose personal information about ordering physicians.

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TECHNICAL INFORMATION

→ Visit Our Testing Portfolio Here: <https://www.foundationmedicine.com/portfolio>

FOUNDATIONONE® CDx

FoundationOne®CDx is a qualitative next-generation sequencing based *in vitro* diagnostic test for advanced cancer patients with solid tumors and is for prescription use only. The test analyzes 324 genes as well as genomic signatures including microsatellite instability (MSI) and tumor mutational burden (TMB) and is a companion diagnostic to identify patients who may benefit from treatment with specific therapies in accordance with the approved therapeutic product labeling. Additional genomic findings may be reported and are not prescriptive or conclusive for labeled use of any specific therapeutic product. Use of the test does not guarantee a patient will be matched to a treatment. A negative result does not rule out the presence of an alteration. Some patients may require a biopsy. For the complete label, including companion diagnostic indications and important risk information, please visit www.F1CDxLabel.com

FOUNDATIONONE® LIQUID CDx

FoundationOne®Liquid CDx is for prescription use only and is a qualitative next-generation sequencing based *in vitro* diagnostic test for advanced cancer patients with solid tumors. The test analyzes 324 genes utilizing circulating cell-free DNA and is FDA-approved to report short variants in 311 genes and as a companion diagnostic to identify patients who may benefit from treatment with specific therapies (listed in Table 1 of the Intended Use) in accordance with the approved therapeutic product labeling. Additional genomic findings may be reported and are not prescriptive or conclusive for labeled use of any specific therapeutic product. Use of the test does not guarantee a patient will be matched to a treatment. A negative result does not rule out the presence of an alteration. Patients who are negative for companion diagnostic mutations should be reflexed to tumor tissue testing and mutation status confirmed using an FDA-approved tumor tissue test, if feasible. For the complete label, including companion diagnostic indications and complete risk information, please visit www.F1LCDxLabel.com

FOUNDATIONONE® HEME

FoundationOne®Heme is a laboratory developed test that combines DNA sequencing of 406 genes and RNA sequencing of 265 genes for patients with hematologic malignancies, sarcomas or solid tumors where RNA sequencing is desired. The test can be used by physicians to identify potential targeted therapy options, detect alterations in prognostic genes, and sub-classify sarcoma diagnoses. For more information on FoundationOne Heme, please see its Technical Specifications at www.foundationmedicine.com/heme

IHC Testing

Scoring and clone utilization for PD-L1 testing is based on FDA-approved indications. Refer to www.foundationmedicine.com/ihc for information.

- Dako 22C3 with Combined Positive Score (CPS) scoring (KEYTRUDA®): Cervical Cancer, HNSCC, ESCC, TNBC
- Dako 22C3 with Tumor Proportion Score (TPS) scoring (KEYTRUDA®, LIBTAYO®): NSCLC
- Dako 28-8 with Tumor Cell Expression scoring (OPDIVO®, YERVOY®): NSCLC
- VENTANA SP142 with Tumor Cell (TC) and Immune Cell (IC) scoring (TECENTRIQ®): NSCLC
- VENTANA SP263 with Tumor Cell (TC) scoring (TECENTRIQ®, LIBTAYO®): NSCLC
- Dako 22C3 with TPS/CPS for other tumors
- VENTANA FOLR1 (ELAHERE™): epithelial ovarian, fallopian tube, or primary peritoneal cancer

CERTIFICATION AND ACCREDITATION

<https://www.foundationmedicine.com/resource/licenses>

FACILITY INFORMATION

This information will be used by Foundation Medicine to determine if the test(s) performed may result in a bill that is affected by surprise billing laws.

MEDICARE COVERAGE SUMMARY (Foundation Medicine tests may be covered by Original Medicare⁴ and Medicare Advantage⁵)

TEST	CONDITIONS FOR MEDICARE COVERAGE	PATIENT COVERAGE CRITERIA
FoundationOne®CDx	Covered ⁶ if all patient coverage criteria are met. ABN required for an Original Medicare beneficiary if they do not meet the patient coverage criteria or if person ordering the test is not a treating physician ⁷ .	i) Patient has been diagnosed with a solid malignant neoplasm; AND ii) Patient has either recurrent, relapsed, refractory, metastatic, or advanced stage III or IV cancer (only requires one of these to be met); AND iii) Patient has not been previously tested with the same test using NGS for the same cancer genetic content ⁸ ; AND iv) Patient has decided to seek further cancer treatment (e.g., therapeutic chemotherapy)
FoundationOne®Liquid CDx		
FoundationOne®Heme	Covered ⁸ if all patient coverage criteria are met. ABN required for an Original Medicare beneficiary if they do not meet the patient coverage criteria or if person ordering the test is not a treating physician ⁷ .	i) Patient has been diagnosed with acute myeloid leukemia (AML), myelodysplastic syndrome (MDS) or myeloproliferative neoplasms (MPN); OR ii) Patient has a suspected myeloid malignancy with an undefined cytopenia for greater than 4 months, and other possible causes have been reasonably excluded AND (both criteria iii and iv below) iii) Patient has not previously received or is not currently receiving NGS testing on the specimen for which the test is currently being ordered iv) Patient has not been tested with the same test for the same genetic content ⁹

References

1. For the most current information about the therapeutic products in this group, go to: <https://www.fda.gov/medical-devices/in-vitro-diagnostics/list-cleared-or-approved-companion-diagnostic-devices-in-vitro-and-imaging-tools>
2. Please reference the US Food & Drug Administration website for a current list of cleared or approved companion diagnostic devices and associated therapies: <https://www.fda.gov/medical-devices/in-vitro-diagnostics/list-cleared-or-approved-companion-diagnostic-devices-in-vitro-and-imaging-tools>
3. Inclusive of the targeted therapies listed and others for which FoundationOne CDx and/or FoundationOne Liquid CDx may be an FDA-approved companion diagnostic in the future
4. Medicare administered by federal government.
5. Medicare administered by private insurers.
6. Decision Memo for Next Generation Sequencing (NGS) for Medicare Beneficiaries with Advanced Cancer (CAG-00450R - reference appendix B).
7. 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>.
8. MolDx Local Coverage Determination (LCD): Next-Generation Sequencing Lab-Developed Tests for Myeloid Malignancies and Suspected Myeloid Malignancies (L38047).
9. Repeat testing (FoundationOne®CDx, FoundationOne®Liquid CDx, or FoundationOne®Heme) after disease progression (i.e., there is evidence of a new malignant growth despite response to a prior targeted therapy) or for additional primary cancer diagnosis may be covered under the NCD for qualifying Medicare beneficiaries.