TEST REQUISITION FORM

For fastest order processing, order online at http://www.foundationmedicine.com Other options: Email (*recommended*) client.services@foundationmedicine.com or Fax to 617.418.2290 IF REQUIRED FIELDS ARE NOT PROVIDED, TESTING MAY BE DELAYED.

For Found	ation N	Andicir	مالم	Only
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									Sex	
First Name	MI	Last Name			Medical Record	#	DOB	(MM/DD/YYYY)	F	Ом
Address			City		State Postal Cod	le Country		Primar	y Phone	
CURRENT DIAGNOSI	S/PATIEN	T HISTOR	Y							
Primary ICD-10 (C&D codes o	only)	Stage	Diagnosis:	inoma () Melanoma		a se Statu Aetastatio	s (select all that a		
Prior/Current Targeted Thera			⊖ Breast ○ N	ISCLC (Ovarian		Refractory	Relapse		
Patient has received transpl	lant? 🔿 Ye	s 🔿 No	○ Prostate ○ O	ther		- OI	lone	O Progress	ion	
Attachments: O Copy of				f available)				ar Diagnostic Ass		C, or
· · ·			, FAB classification.		-		g., ER, PR, I	HER2, EGFR, KRAS,	etc.	
TREATING PHYSICIA	N INFORM	MATION (P	lease provide best	contact in	formation for case	e follow-up)				
Facility Name				Treating F	Physician (full legal n	ame)				
Facility Address				City			State	Postal Code Co	ountry	
Foundation Medicine Accoun				Email				Phone	Fax	
Additional Physician to be Co			ity Name (optional)			Email (option			Fax (option	al)
TEST MENU Test/Lab	oratory De	veloped Tes	t (LDT) Selection							
Genomic Test	Descriptio		Accepted Specir	nen Type G	enomic Test	Des	cription		Accepted Specin	nen Type
FoundationOne®CDx FoundationOne®Liquid C	diagnos	proved compa tic for solid tu proved compa	tumors panion Peripheral		Specimen has or is for sar		or heme malignancies, Blood, E sarcomas or solid tumors Aspirat		Peripheral W Blood, Bone M Aspirate, FFP Extracted Nu	Marrow E Tissue,
 If specimen submitted i 	diagnos	tic for solid tu	umors Whole Blood	A	NGS testing					
use portfolio reflex opt				(IHC Testing PD-L1	I (see back fo	r details)		FFPE tissue	
SPECIMEN RETRIEVA	L Provide	all informat	ion required per sa	mple type						
Submitting Pathologist Na	m o	Pathology La	ah Namo	 Ema			Phone		Fax	
I am requesting a spe			ill let the pathologist			🗌 I am prov		= E block return add		f form
Date of Collection (∧	MM/DD/YYY	(Y) Specin	nen ID		Site of Biopsy			 Alternate Choi	ce (optional)	
○ FFPE Tissue			O Periphera	al Whole B		С		arrow Aspirate/I		
 I will arrange for spec Contact the patholog 			Mobile	-	specimen shipment ny requested ebsite)	t	Orde	ring Facility respo	onsible for ship	ment
BILLING INFORMATIO	ON Select	one of the t	hree payment optic	ons and co	mplete all fields in	ndicated (As	erisk indic	ates Medicare requi	rement)	
Insurance (check one):	🗌 * Medic	are 🔿 * ABI	N Attached 🛛 🗆 M	Aedicare A	dvantage 🗌 Othe	er Plan Na	ame			
					*			e of specimen coll		
Policy #	Group #		Prior Authorizati	on #		□ Office (•	ital) 🗌 Outpa s Discharge Date be		YYY)
Self-Pay: Contact Name	e	Email		Phone	e			Not yet discha		
○ Facility: Address		City			Postal Code Cou	Intry		— 🗌 Same as	Treating Physic	ian:
CERTIFICATE OF MED	DICAL NE	CESSITY/C	ONSENT/TEST	AUTHO	RIZATION AND	PHYSICI	AN SIG	NATURE		
My signature constitutes a Certi physician. I have explained to th Medicine to (a) perform the test or disclose such de-identified re	e patient the n ting specified I	ature and purp nerein, (b) retai	ose of the testing to be n the test results for an	performed a indefinite pe	nd have obtained inform riod for internal quality	med consent, t y assurance/or	o the exten perations p	t legally required, to urposes, (c) de-iden	permit Foundation tify the test result	on ts and use
My signature also authorizes Four										
Treating Physician Signature					Printed Name (Full le	eaal name)			Date (MM/DD/)	(

FFPE BLOCK RETURN INFORMATION

FFPE Block Return Address			
City	State	Postal Code	Country
Email		Phone	

TECHNICAL INFORMATION

FOUNDATION**ONE®CD**x

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 http://www.FICDxLabel.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 available. For the complete label, including companion diagnostic indications and complete risk information, please visit *http://www.FILCDxLabel.com*

OTHER INFORMATION

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

Portfolio Reflex Option:

If the "Portfolio Reflex" checkbox 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 detailed below 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 https://www.foundationmedicine.com/genomic-testing/order for more information.

Additional Case Information (optional)

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FOUNDATION**ONE®HEME**

About the Test 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 sensitive fusion detection 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 http://www.foundationmedicine.com/heme

IHC Testing

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

MEDICARE COVERAGE SUMMARY (Foundation Medicine tests may be covered by Original Medicare¹ and Medicare Advantage²)

TEST	CONDITIONS FOR MEDICARE COVERAGE	PATIENT COVERAGE CRITERIA		
		i) Patient has been diagnosed with a solid malignant neoplasm; AND		
FoundationOne [®] CDx	Covered ³ if all patient coverage criteria are met. ABN required if patient does not meet the patient	 ii) Patient has either recurrent, relapsed, refractory, metastatic, or advanced stage III or IV cancer (only requires one of these to be met); AND 		
	coverage criteria or if person ordering the test is not a treating physician ⁴ .	iii) Patient has not been previously tested with the same test using NGS for the same cancer genetic content ⁶ ; AND		
FoundationOne®Liquid CDx		 iv) Patient has decided to seek further cancer treatment (e.g., therapeutic chemotherapy) 		
FoundationOne [®] Heme		i) Patient has been diagnosed with acute myeloid leukemia (AML), myelodysplastic syndrome (MDS) or myeloproliferative neoplasms (MPN); OR		
	Covered ⁵ 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 ⁴ .	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 $^{\rm 6}$		

References

1. Medicare administered by federal government.

2. Medicare administered by private insurers.

3. Decision Memo for Next Generation Sequencing (NGS) for Medicare Beneficiaries with Advanced Cancer (CAG-00450R - reference appendix B)

4. 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.

MolDx Local Coverage Determination (LCD): Next-Generation Sequencing Lab-Developed Tests for Myeloid Malignancies and Suspected Myeloid Malignancies (L38047)
 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) may be covered under the NCD for qualifying Medicare beneficiaries.

