() labcorr	• Oncology	Client Services TN: (800) 874-8532 fax: (615) 370-8074		HEMATOLOGY/ONCOLOGY oncology.labcorp.com				
•		AZ: (800) 710-1800 fax: (800) 481-4151			Highlighted fields are REQUIRED			
CLIENT INFORMAT	<b>TION</b>	NPI #	COMPREHENSIV (Peripheral Blood or B	e Hematopathology	ANALYSIS			
TREATING PHYSICIAN		NP1 #		aluation: Morphologic evaluatio	n Flow Cytometry			
		NPI #	Cytogenetics, and	Other Relevant Diagnostic and/c	or Prognostic Tests per Opinion			
PHYSICIAN/AUTHORIZED SIGNATURE	E			ologist (see reverse for prognosti				
				aluation as above without Cytog				
Client#				EVALUATION (include o	a copy of CBC report)  Peripheral Blood Morphology			
Client Name			Bone Marrow Morp	0.				
				RY <sup>®</sup> (see reverse for antiboo				
Address			Add diagnostic per IO Reflex Cr Add prognostic per IO Reflex Cr DNA Ploidy/S-Phas	iteria (see reverse) tests iteria (see reverse)	<ul> <li>BAL CD4:CD8 Assessment<sup>1</sup></li> <li>ZAP70/CD38/CD49d Assessment</li> <li>PNH ◆</li> <li>Stem Cell Enumeration<sup>1</sup></li> <li>CLL MRD<sup>2</sup></li> <li>ALL MRD<sup>2</sup> (meets COG requirements)</li> <li><sup>1</sup> Send to TN <sup>2</sup> Send to CT</li> </ul>			
Phone Number		Fax Number						
Phone Number			CATOGENETICS		Constitutional Cytogenetics <sup>‡</sup>			
			1 0	es e state profile OR individual pro				
PATIENT INFORMA	TION			s (see reverse for panel comp				
Name (LAST, FIRST, MI):			ABC Lymphoma					
Date of Birth:		Sex: 🗌 Male 🔲 Female	□ Multiple Myeloma		/ Eosinophilia 🛛 MDS			
Address:			Pediatric (COG)	ALL (Std Risk) ALL (Hi	gh Risk) 🔲 AML			
City, State, Zip:			COG Single Probes	ABL1 ABL2	PDGFRb			
Phone Number:			Individual Probes (fo	r a complete list of probes vis				
Med. Rec. # / Patient #	t.		5q					
		nd copy of insurance card – both sides)	CCND1/IGH, t(11;1	eflex to JAK2 V617F Qual, If JAK	2 neg reflex to CALR and MPL CL2, t(14;18)			
	Insurance  Medicare In-Patient Ou			KM2TA (MLL)     PML/RARA     RUNX1/RUNX1T1, f(8;21)     TCRA/D      TP53 (17p-)				
		SECONDARY BILLING PARTY	<b>MOLECULAR</b> <sup>@</sup>					
INSURANCE CARRIER		INSURANCE CARRIER	IntelliGEN® NGS Asso	y (see reverse for gene list; b	one marrow or peripheral blood)			
ID #		ID #	IntelliGEN <sup>®</sup> Myeloi Indication:	d for AML, MDS, MPN				
GROUP #		GROUP #		v for MDD for Multiple Muclem	CLL DALL (Dillad by Adaptive Distashpalazias)			
INSURANCE ADDRESS		INSURANCE ADDRESS	clonoSEQ® NGS Assay for MRD for Multiple Myeloma, CLL, B-ALL (Billed by Adaptive Biotechnologies) Indication:					
			Specimen types: Blood or bone marrow. For blood or fresh bone marrow aspirate, use a lavender-top					
NAME OF INSURED PERSON		NAME OF INSURED PERSON	(EDTA) tube only. clonoSEQ ID test for Multiple Myeloma requires bone marrow.					
RELATIONSHIP TO PATIENT		RELATIONSHIP TO PATIENT	clonoSEQ ID. <u>Must be run first to establish baseline</u> . Performed using a high disease burden diagnostic specimen (fresh or archived). If diagnostic specimen is not accompanying this order complete Procurement information in SPECIMEN INFORMATION section. For CLL/SLL, IGHV mutation					
EMPLOYER NAME		EMPLOYER NAME						
*IF MEDICAID STATE PHYS	ICIAN'S PROVIDER #	WORKERS Ves No	status will be report	ed.				
SPECIMEN INFOR	MATION				llected during or after treatment. Patient must have			
Collection Date:	Time:	🗆 AM 🗖 PM	had a previous clon	oSEQ ID test performed. If not, pl	ease also check clonoSEQ ID above.			
Specimen ID #(s):					B-ALL, please complete and submit an ABN,			
Body Site/Descriptor:			Reveal® SNP Microar	eq.com/for-clinicians/ordering.	ed translocations, run cytogenetics and/or FISH			
Fixative: 10% Neutral Specimen Type:	Buffered Formalin Ot	ther: Hours Fixed: Smears:	SNP Microarray for	ALL, AML, CLL, MDS and other H	Hematologic Malignancies			
BM Aspirate	Fluid:	Peripheral Blood #	Indication:					
BM Clot	FNA:	BM Touch Preps #		rray for Multiple Myeloma	SNP Microarray for Multiple Myeloma			
BM Core	CSF	BM Aspirate #			11; 14), t(14; 16) are performed			
Dry Tap	Lymph Node:	Effusion #/Source	Acute Leukemia	Lymphoid Neoplasm B-cell Rearrangement IgH	MPN/CML/Mastocytosis			
Peripheral Blood	Slides #	Fresh Tissue #/Site	DH 1/2 Mutation	T-cell Rearrangement TRG/1				
If slide procurement require Facility Name:			CEBPA Mutation	B-cell Rearrangement IgH				
Address:			NPM1 Mutation PML/RARA	B-cell Rearrangement IgK T-cell Rearrangement TRG	JAK2 V617F Mutation			
Phone Number:		Fax Number:	(Quantitative)	T-cell Rearrangement TRB				
	TION FOR STUDY (at	tach clinical history and pathology reports)	CKIT Mutation	BCL1 Rearrangement	CALR CALR			
		eport with diagnosis, indication for study, and previous test results)	LeukoStrat® CDx FLT3 Mutation	BCL2 Rearrangement IqVH Mutation	JAK2 Exon 12-15			
			FLI'S MUIUIUI	p53 (CLL/B-cell ONLY)	MPL 515 JAK2 Exon 12-15 Mutation MPL 515 Mutation			
	For pediatric pat			MYD88 Mutation	MPL 515 Mutation CALR Mutation			
	vided by the ordering physician of in ICD-CM format in effect at Dat	or an authorized designee. te of Service (Highest Specificity Required)			KIT D816V Mutation Digital PCR			
ICD-CM	ICD-CM formal in ellect at Dat		🗖 Other Molecular, sp	,				
Acute Lymphoblastic Le			SPECIAL CHEM	ISTRY (Serum ONLY)				
B-cell T-cell	🗖 Leukemia, Un	specified Dyeloproliferative Neoplasm	Multiple Myeloma Di		*Meets IMWG Guidelines			
Lineage Uncertain	a Leukocytosis,	Unspecified Non-Hodgkin Lymphoma Polycythemia	120256 Immunofixation (sIFE), Protein Electrophoresis (SPE), Quant Free K/X Light Chains (sFLC)* 123200 Mutiple Myeloma Cascade, SPE Reflex to sIFE and sFLC					
🗖 Anemia								
Chronic Lymphocytic Le			001495 sIFE, SPE	001487 S 🗌 🗌 VIII A A A A A A A A A A A A A A A A A				
Chronic Myelogenous Leukemia     Myeloma, Plasma Cell     Thrombocytosis     Disease Stage/Clinical Course:     New Diagnosis     Relapse     Follow-Up     Other:								
	adiation Chemotherapy			SEQ is a registered trademark of Adaptive Biotechnologies www.adaptivebiotech.com				
Mana and size basis for 1111	Manifester on Manifestial and 1		<ul> <li>Peripheral blood only</li> <li>*If sending DNA, the lab only acc</li> </ul>	epts isolated or extracted nucleic acids for	which extraction or isolation is performed in an appropriately qualified			

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When ordering tests for which Medicare or Medicaid reimbursements will be sought, physicians should order only those tests that are medically necessary for the diagnosis or treatment of the patient.

				Prognostic Test Refle	ex Criteria				
Disease Category	Timing	Finding	(Morpholog	y, Flow cytometry, FISH and		Tests to Perfo	orm		
ALL	Initial Diagnosis	ALL				Pediatric FISH Profile (<22 years) or Adult FISH Profi		dult FISH Profile (>22	
4.841						years); Reveal® SNP Array			
AML	Initial Diagnosis	AML or b	orderline AML			FISH probes for RUNX1T1/RUNX1 t(8;21), CBFB inv(16), or PML/ RARA t(15;17) or MLL respectively, as indicated; NGS myeloid panel + FL73 testing for patients <60 years; discuss necessity of testing with client or place comment in report for patients >= 60 years			
AML	Relapse	Findings	indicative of rel	apse		NGS myeloid panel <60 years; discuss necessity of testing with client or place comment in report for patients >= 60 years			
CLL (peripheral blood/bone marrow)	Initial Diagnosis			ssic or variant CLL features; > 0% or more marrow based mo		CLL FISH profi	FISH probe for CCND1/IGH and IgVH mutation analysis		
CLL (peripheral blood/bone marrow)	Follow-up*	Features	of refractory dis	ease or disease progression/t	ransformation	FISH probe for SNP array	TP53 (17p-) deletion, T	P53 mutation analysis, and	
CML	Initial Diagnosis	Compatil	ole or diagnosti	nostic findings for CML Quantitative BCR/ABL1 assay an				genetics	
CML	Follow-up*	Prior dia	gnosis of CML			Quantitative BCR/ABL1 assay; if features of progression, discuss addition of NGS myeloid panel with client or place comment in report			
MPN	Initial Diagnosis	Morpholo		MPN, but negative for JAK2 V6	17F, CALR, and MPL	NGS myeloid panel for patients <60 years; discuss necessity of testing with client or place comment in report for patients >= 60 years			
MPN	Follow-up*		f MPN, currently c features)	with features of progression (	increased blasts or	Discuss addition of NGS myeloid panel with client or place comment in report			
MDS	Initial Diagnosis	Morphole	ogic diagnosis c	f MDS with normal cytogenetic	c karyotype	NGS myeloid panel for patients <60 years; discuss necessity of testing with client or place comment in report for patients >= 60 years			
Plasma cell neoplasia	Initial Diagnosis	5% or m by flow c		lasma cells by morphology or	1% or more	Myeloma FISH profile			
Plasma cell neoplasia	Follow-up*	Features	of disease prog	ression		FISH probes fo	or TP53 (17p-), CKS1B (	1q21), Monosomy 13/13q-	
SLL	Initial Diagnosis	neoplast	c cells	Imple by flow cytometry with 1	0% or more	CLL FISH profile or CLL SNP array with FISH probe for CCND1/IGH t(11;14), IgVH mutation analysis			
*recommendation for follow-u									
	Diagnostic	Test Reflex C	riteria Base	d on Flow Cytometry	y or Surgical Patholo	<u> </u>			
Disease Category	Timing	Finding				Tests to Perfo			
AML	Initial Diagnosis		ARA t(15;17), (	for AML with RUNX1T1/RUNX1 acute myelomonocytic, or acut		RARA t(15;17) + FLT3 testing	) or MLL respectively, as	21), CBFB inv(16), or PML/ indicated; NGS myeloid panel discuss necessity of testing with atients >= 60 years	
B-cell lymphoma	Initial Diagnosis	with rega	rd to subclassific	ignostic for B-cell lymphoma, b cation (for tissue cases 5% or m blood/bone marrow cases, 10%	nore abnormal B-cells by flow	NHL FISH probes and molecular assays as indicated			
Large B-cell lymphoma or Burkitt lymphoma			Abnormal B-cells diagnostic or suspicious for large B-cell lymphoma or Burkitt lymphoma				FISH probes for MYC, BCL6, and BCL2 translocations and cytogenetic karyotyping, as indicated; reflex to 11q FISH probe (BCL1 and ATM) for MYC, BCL6, BCL2 negative cases suspicious for Burkitt lymphoma, as indicated		
Eosinophilia	Initial Diagnosis	Periphero	I blood with 1.0	)K/µL or more eosinophils			or PDGFRA, PDGFRB, and	d FGFR1	
Hairy Cell Leukemia (HCL)	Initial Diagnosis	CD103+	monoclonal B-(	cells (5% or more) inconclusiv	e for HCL	BRAF mutation	n		
Lymphoplasmacytic	Initial Diagnosis	Monoclo	nal B-cells (10%	6 or more) with features indica	ting LPL in	MYD88 mutat	ion		
Lymphoma (LPL)	Initial Dimensio		al diagnosis			FIGU and a fee	000001/0004/11.140		
Mantle cell lymphoma (MCL)	Initial Diagnosis				more) diagnostic or suspicious of MCL		FISH probe for CCND1/IGH t(11;14)		
Mastocytosis CML	Initial Diagnosis	/1	mast cells by flo	1 1		High-sensitivity KIT D816V mutation analysis for m FISH for BCR/ABL1		narysis for mast cell alsease	
MDS/MPN	Initial Diagnosis		· ·	suspicious for CML IDS/MPN (CMML, aCML, etc.)			panel for patients <60 ye	are discuss necessity of	
	Initial Diagnosis	- I munigs						report for patients >= 60 years	
T-cell lymphoma/leukemia	Initial Diagnosis			c or suspicious for T-cell lymp		TCR gene rearrangement ; ALK FISH probe for CD30+ cases, as indicated; cytogenetic karyotyping if material adequate			
+LeukoStrat® CDx FLT3 Mutation	on performed by The	e Laboratory for Pe	rsonalized Mol	ecular Medicine (LabPMM®)	<sup>+</sup> Informed consent is	s required for no	n-oncology genetics tes	sting for New York state patients.	
		Morphologic		Common Compone	nts (Please include po	tient CBC re	port)		
Peripheral Blood Interpreto	· ·	• Clot (88305)	Core (88)					(88313) – Iron and Reticulin	
Bone Marrow Aspirate Sm	ear & Interpretation	(85097)	Decalcition	ation (88311)		• IHC Global	marker number (8834)	<ol> <li>varies but typically 0-4</li> </ol>	
				Flow Cytomet	ry*	1			
Peripheral blood/bone 24 *® antibodies	marrow panel	(HNA)	Tissue/flu 21 *⊕ anti	ids panel (HNA) bodies		PNH Evaluc	ation		
CD2, CD3, CD4, CD5, CD7,	CD8, CD10, CD11b	), CD13, CD14,	14, CD2, CD3, CD4, CD5, CD7, CD8, CD10, CD11b, CD19, CD20,		CD14, CD15, CD24, CD45, CD64, FLAER.				
CD16, CD19, CD20, CD23, CD56, CD64, HLA-DR, kapp			CD45, CD23, CD30, CD38, CD43, CD45, CD56, CD57, FMC-7,		CD59 and CD235a may be added at discretion of reviewing pathologist				
	-	-		nder a diagnosis in the opini			⊕Markers performe	ed determined by testing facility.	
Additional antibodies may be		sa to be meatering		H (disease state profile OR i		9151.		sa acionimica by losning racimy.	
ALL (Adult) ALI	(Pediatric)	AML	116	CLL	MPN/CML	Multi	iple Myeloma	NHL	
BCR/ABL1, t(9;22) E	CR/ABL1,t(9;22)	PML/RARA, t(15	17)	TP53 (17p-)	20q	Mo	onosomy 13/13q- 53 (17p-)	ALK	
MYC 1	0	CBFB, inv(16) RUNX1T1/RUN>	(1, t(8;21)	ATM (11q-) CCND1/IGH, t(11;14)	8 9	7 9	00(170)	BCL6 CCND1/IGH, t(11;14)	
	7 MT2A (MLL)	5q 7q		13q14 (DLEU) 12	13q14 (DLEU) BCR/ABL1, t(9;22)	15		IGH/BCL2, t(14;18) IGH/MYC, t(8;14)	
, C	DKN2A (p16)	KMT2A (MLL)		12	DOIGNDET, ((0,22)	CK	ND1/IGH, t(11;14) (S1B (1q21) (FR3/IGH, t(4;14)	MALT1	
E	CF3 (E2A) TV6/RUNX1, t(12;21)	Aggressive B-cell (	ABC) Lymphoma	MDS	MPN with Eosinophilia	IGH	H/MAF, t(14;16)	TCRAVD	
		BCL2 BCL6 MYC		5q 7q 20q	FGFR1 PDGFRA PDGFRB				
				8					
SERUM - Mu	Itiple Myelom	a Cascade,		ctrophoresis (SPE) re or interpretation, refer to ww		ition (sIFE)	and Free Light C	Chain (sFLC)	
			IntelliGE	${ m I}^{ m I\! B}$ (for genes evaluated, refer	to oncology.labcorp.com)				
						Darzalex® is a re	egistered trademark of J	ohnson & Johnson Corporation	
LeukoStrat® is a registered trademark of Invivoscribe Technologies, Inc.									
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				Lab Location	าร				
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