

ALVERNO LABORATORIES

HEMATOPATHOLOGY Patient Information Sheet

CLIENT INFORMATION		PATIENT INFORMATION				
Account Name:		Last Name:	First Nar	ne:	M.I.	
Treating Physician (First, Last):		Date of Birth (mm/dd/yyyy):	Age:	Sex:		
Physician's Contact #:		Order Date:	r Date: Medical Record #:			
Physician's Signature:				1		
BILLING INFORMATION						
	Hospital Patient (out) ☐ Non-Hospital Patient are ☐ Medicaid ☐ Patient/Self-Pay ☐ Split Billing			g to Client		
SPECIMEN INFORMATION						
Col	lection Date mm/ dd/	yyyyCollection Time		·M		
	en Top(s) Purple Top(s) Other					
☐ Bone Marrow (attach CBC result): Green T	op(s) Purple Top(s) Core E	Biopsy Clot	☐ Fluid: CSF	Pleural Other:		
CLINICAL INFORMATION						
 □ Eosinophilia □ Monoclonal gammopa Diagnosis Under Consideration (C □ Chronic Lymphoproliferative Disorder 	Thrombocytopenia □ Leukopenia □ Ane thy □ Plasmacytosis □ Lymphadenopathy heck All That Apply): □ non-Hodgkin L □ Myelodysplastic Disorder □ Myeloprol □ Minimal residual disease □ Relapse □	y □ Extranodal mass □ Splenomegaly □ ymphoma □ Hodgkin Lymphoma □ A liferative Neoplasms □ Multiple Myelon □ BM Transplant	OtherAcute Leukemia		- -	
		TEST MENU				
Flow Cytometry - EDTA Global Tech-Only Lymphoma/Lymphocytosis Panel	Fluorescence in situ Hybridization (FISH) - Sodium Heparin *Process & hold RQFSH (all panels except Myeloma) *Plasma Cell Enrichment - process & hold RQPCE (all panels including Myeloma) *Call or email within 14 days of collection to add on panel.			Molecular Genetics - EDTA ** Prior insurance authorization may be required. □ Extract Nucleic Acid and Hold* (* Call or email within 14 days to add NGSMY panel.)		
(For CLL, MM, and NHL studies) FLWCY	_	r-pathologist for interp:	N	Myeloid Neoplasms Mutation Pa NGS (40 DNA /29 RNA genes)	anel by NGSMY	
□ Leukemia/Lymphoma Comprehensive Panel FLWCY (For ALL, AML, CML and MDS studies) Reflex panels if indicated □ Plasma Cell	Select a panel below: [FISH probes may be a probe by the probe by the probes may be a probe by the probe by the probes may be a probe by the probe b	□ CML Panel □ BCR/ABL t(9;22) □ Myeloma Panel □ Deletion 1p/1q Gain □ Deletion 13q/Monosomy 13 □ Trisomy 3, 5, 9	CMLFH MMYFH GTC	www.genomictestingcooperative Hematology Profile Plus Liquid Trace™Hematology st Diagnostics www.questdiagno		
□ Hairy Cell □ Acute Leukemia Intracellular Markers: (nTdT, cMPO, cCD3, cCD79a) □ T-cell Receptor □ PNH (QUEST) PNHHS	□ Trisomy 8 □ Deletion 20q □ Inv(3) 3q26 □ RUNX1/RUNX1T1 (AML/ETO) t(8;21) □ PML/RARA t(15;17) (APL) □ MYH11/CBFB; inv(16), t(16;16) □ MLL Rearrangement (11q23	□ Deletion 17p (<i>TP53</i>) □ IGH Rearrangement (14q32) Reflex to □ IGH/CCND1, t(11;14) □ IGH/FGFR3, t(4;14) □ IGH/MAF, t(14,16) □ NHL Panel		SCR/ABL1, Quantitative PCR AK2 V617F Mutation QL AK2 Exon 12 Mutation AK2 V617F Mutation QL, Casc Reflex CALR, JAK2 Exon (12, MF	PL, CSF3R	
Cytogenetics (Chromosome Analysis) (QUEST) - Sodium Heparin Chromosome Analysis, Hematologic Malignancy Myelogenous-MDS,MPD,ALL,CML (blood or bone marrow) Chromosome Analysis, CLL/LPD Lymphoproliferative CLL,LPD (blood or bone	□ CLL/SLL Panel □ Deletion 11q (ATM) □ Deletion 13q/Monosomy 13 □ Deletion 17p (TP53) □ Trisomy 12 □ IGH/CCND1, t(11;14) □ MDS Panel □ Deletion 5q/Monosomy 5 □ Deletion 7q/Monosomy 7 □ Trisomy 8 □ Deletion 20q	 □ BCL6 Rearrangement (3q27) □ MALT1 Rearrangement (18q21) □ MYC Rearrangement (8q24) □ IGH Rearrangement (14q32) □ IGH/BCL2, t(14;18) □ IGH/CCND1, t(11;14) 	MPNEF FR1, and CBFB	I-Cell Clonality by PCR (IgH/IgK -Cell Clonality by PCR (TCRB, 1908 L265P Mutation IPM1 Mutation (Exon 12) ILT3 ITD&TKD Mutation EBPA Mutation Detection P53 Somatic Mutation JVH Mutation Status (CLL)		
marrow)			— I 🗆 U	Other		

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TEST MENU DETAILS

Genomic Testing Cooperative (GTC)

GTC-Hematology Profile Plus

GTC-Hematology Profile Plus combines expression and fusion with mutation analysis in DNA and RNA. The test covers 179 DNA genes and 1408 RNA genes. This is a comprehensive evaluation of all hematologic neoplasms. However, it is especially recommended for:

- Acute Lymphoblastic Leukemia (ALL): This comprehensive assay is designed to confirm the diagnosis of Ph-ALL and Ph-like ALL and distinguish them from other types of ALL. It can be used for diagnosis as well as for monitoring. Ph-like ALL is detected in 20% to 25% of adult ALL and in 15% of pediatric ALL. Diagnosis of Ph+-ALL and Ph-like ALL is very important because TKI therapy can be helpful in most of these patients. This assay can determine most of the mutations, translocations, and expression of genes (CRLF2) associated with Ph+ ALL and Ph-like ALL.
- Diffuse Large B-cell Lymphoma (DLBCL) and other Types of Lymphoma: This assay can provide very valuable information for the management and monitoring of patients with DLBCL. It can distinguish between ABC and GCB and can help in the diagnosis of double hit lymphoma. The assay is also useful for follicular lymphoma and T-cell neoplasms
- Acute Myeloid Leukemia (AML): Translocations in AML are very important for diagnosis, prognosis and selecting therapy. This comprehensive testing can provide a complete evaluation of fusion mRNA and mutations. It also helps in determining a diagnosis in acute leukemia with ambiguous phenotype.
- Clonal Hematopoiesis of Indeterminate Potential (CHIP): Distinguish CHIP from clinically active and relevant hematologic neoplasm based on an internally developed algorithm using variant allele frequency, chromosomal structural abnormalities, clinical and laboratory data and longitudinal data. This distinction is particularly important when evaluating minimal residual disease and in the presence of other neoplastic process
- IgVH Mutation Status: IgVH mutation status is very important for prognosis and selecting therapy in patients with chronic lymphocytic leukemia (CLL).
- VEXAS Syndrome: Recently described VEXAS (vacuoles, E1 enzyme, X-linked, autoinflammatory, somatic) is caused by mutations in the UBA1 gene. This is an adults-onset fatal disease that may present as myelodysplastic syndrome, aplastic anemia or multiple myeloma, but characterized by fevers, low white cell count, vacuoles in bone marrow cells, dysplastic bone marrow, pulmonary inflammation, chondritis, and vasculitis. Detecting the presence of mutations in the UBA1 gene is the only way for confirming the diagnosis of this syndrome.

Liquid Trace[™] Hematology

Pan-Tumor Assay for Hematologic Malignancies

GTC's Liquid Trace Hematology is a pan-cancer highly sensitive test evaluating cfRNA and cfDNA providing highly informative data that can be used for diagnoses, evaluating the host immune response, and identifying biomarkers for predicting responses to various therapies

GTC's Liquid Trace can significantly reduce the need for bone marrow biopsies for hematology patients. Furthermore, the test can detect, chromosomal abnormalities, translocations, and gene amplifications. Liquid Trace can detect all types if hematologic cancers including:

- Multiple myeloma
- MPN MRD
- Lymphoma
- VEXAS syndrome
- Acute Lymphoblastic Leukemia
- EBV related neoplasms
- Acute myeloid leukemia
- MDS CMML
- Hypersinophilia

Liquid biopsy in its current form is dependent on cfDNA analysis; this method likewise presents multiple challenges. These include variations in DNA shedding between tumors as well as low sensitivity (especially in early-stage cancer), difficulty in detecting fusion genes (i.e., chromosomal translocations leading to the expression of chimeric mRNA from two genes), and inability to reflect the numerous biological processes that modify RNA expression levels, such as alternative splicing, stability, and allele-specific methylation. The latter limitation is critically important as recent studies have shown that RNA testing provides another level of biological information regarding the tumor and its microenvironment.

The Benefits of cfRNA

RNA sequencing has proven to be more sensitive for some types of mutations. Cancer cells typically contain one copy of mutated DNA but numerous copies of RNA. This research is consistent with GTC's findings that cfRNA has increased sensitivity over cfDNA alone. More specifically, cfRNA allowed GTC's Liquid Trace to detect more mutations and fusions in hematologic and solid tumor samples, which may be undetected by

Hematology **Profile Plus**

Genes: 179 /1408

TAT: 7-10 Days

Indications

All hematologic neoplasms including lymphoma Classification and diagnosis of lymphoma, multiple myeloma, acute lymphoblastic leukemia, and acute myeloid leukemia Includes IaVH

Chromosomal abnormalities, and gene amplifications

Sample Type

Bone marrow, Peripheral blood, Fresh tissue

Sample Requirements

Bone marrow: 2ml. Peripheral blood: 5 ml. EDTA tube preferred

FFPE: 1 H&E slide and 6-10 unstained slides, 5-7 microns of tissue fixed with 10% NBF fixative

Results Reported:

DNA + RNA

Liquid Trace™ Hematology

Genes: 284/1501

TAT: 5-7 Days

Indications

All hematologic neoplasms including lymphoma multiple myeloma, acute lymphoblastic leukemia, acute myeloid leukemia, MDS, CMML, MPN, MRD, VEXAS syndrome, and EBV Chromosomal abnormalities, and gene amplifications

Sample Type

Peripheral blood

Sample Requirements

8-10 mL. EDTA tube is required RNA stability is 48-72 hours from blood draw. DNA stability is 7 days from blood draw. Samples received beyond 72 hours may include only DNA results.

Results Reported:

DNA + RNA