

CLIENT INFORMATION
ORDERING PHYSICIAN
TREATING PHYSICIAN
PHYSICIAN/AUTHORIZED SIGNATURE

PATIENT INFORMATION
Name (LAST, FIRST, MI):
Date of Birth:
Sex: Male Female
Address:
City, State, Zip:
Phone Number:
Med. Rec. # / Patient #:

BILLING INFORMATION (attach face sheet and copy of insurance card - both sides)
Bill: My Account Insurance Medicare Medicaid Patient Workers Comp
Patient Hospital Status: In-Patient Out-Patient Non-Patient
Insurance Information: See attached Authorization #
PRIMARY BILLING PARTY SECONDARY BILLING PARTY
INSURANCE CARRIER
ID #
GROUP #
INSURANCE ADDRESS
NAME OF INSURED PERSON
RELATIONSHIP TO PATIENT
EMPLOYER NAME
*IF MEDICAID STATE PHYSICIAN'S PROVIDER # WORKERS COMP Yes No

SPECIMEN INFORMATION
Collection Date: Time: AM PM
Specimen ID #(s):
Body Site/Descriptor:
Fixative: 10% Neutral Buffered Formalin Other: Hours Fixed:
Specimen Type: Smears:
BM Aspirate Fluid: Peripheral Blood #
BM Clot FNA: BM Touch Preps #
BM Core CSF BM Aspirate #
Dry Tap Lymph Node: Effusion #/Source
Peripheral Blood Slides # Fresh Tissue #/Site
If slide procurement required, indicate below:
Facility Name:
Address:
Phone Number: Fax Number:

CLINICAL INDICATION FOR STUDY (attach clinical history and pathology reports)
Narrative Diagnosis/Clinical Data (please include Pathology report with diagnosis, indication for study, and previous test results)
For pediatric patients ONLY: COG Study COG Post Treatment
All diagnoses should be provided by the ordering physician or an authorized designee.
Diagnosis/Signs/Symptoms in ICD-CM format in effect at Date of Service (Highest Specificity Required)
ICD-CM ICD-CM ICD-CM
Acute Lymphoblastic Leukemia B-cell T-cell Lineage Uncertain
Acute Myeloid Leukemia Anemia
Chronic Lymphocytic Leukemia Chronic Myelogenous Leukemia
Hodgkin Lymphoma Leukemia, Unspecified Leukocytosis, Unspecified Leukopenia Lymphadenopathy Monoclonal Gammopathy Myeloma, Plasma Cell
Myelodysplastic Syndrome Myeloproliferative Neoplasm Non-Hodgkin Lymphoma Polycythemia Suspected malignant neoplasm Thrombocytopenia Thrombocytosis
Disease Stage/Clinical Course: New Diagnosis Relapse Follow-Up Other:
Post Treatment: Radiation Chemotherapy BM Transplantation Donor: M F

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.

Patient, client, and billing information is requested for timely processing of this case. Medicare and other third party payors require that services be medically necessary for coverage, and generally do not cover routine screening tests.

When ordering tests that are subject to ABN guidelines, refer to the policies published by your Medicare Administrative Contractor (MAC), CMS, or www.Labcorp.com/MedicareMedicalNecessity.
Symbols Legend
@ = Subject to Medicare medical necessity guidelines
^ = Medicare deems investigational. Medicare does not pay for services it deems investigational.

SPECIMEN LABEL INSTRUCTIONS

- 1. Complete the requisition with all requested information.
2. Label specimen with two unique identifiers.
3. Remove the required number of labels from the front of this sheet.
4. Place one (1) label on each specimen container (not on the lid).
Please dispose of unused labels.

Name _____ Name _____ Name _____ Name _____

Name _____ Name _____ Name _____ Name _____

LABCORP HEMEPATH CASE (Comprehensive Analysis Services & Expertise) (Peripheral Blood or Bone Marrow)

- Comprehensive Evaluation: Morphologic Evaluation, FLOW Cytometry, Cytogenetics, and Other Relevant Diagnostic and/or Prognostic Tests per Opinion of Reviewing Pathologist (see reverse for reflex criteria)
Comprehensive Evaluation as above without Cytogenetics

MORPHOLOGIC EVALUATION (include a copy of CBC report)

- Bone Marrow Morphology Peripheral Blood Morphology

FLOW CYTOMETRY (see reverse for antibody list)

- Leukemia/lymphoma phenotyping Add diagnostic/prognostic tests - per LCO reflex criteria (see reverse)
DNA Ploidy/S-Phase Assessment Leukocyte Adhesion Deficiency Assessment BAL CD4:CD8 Assessment
PNH Stem Cell Enumeration CLL MRD B- ALL MRD (meets COG requirements)

CYTOGENETICS

- Cancer Cytogenetics Constitutional Cytogenetics

FISH (select disease state profile OR individual probes)

Disease State Profiles (see reverse for panel components)

- ALL (Adult) ALL (Pediatric) ALL (Philadelphia-like) CLL
High Grade Multiple Myeloma AML MDS
B-cell Lymphoma MPN/CML MPN w/ Eosinophilia

Pediatric (COG) COG Single Probes

- ALL (Std Risk) ALL (High Risk) AML
ABL1 ABL2 PDGFRb

Individual Probes (for a complete list of probes visit oncology.labcorp.com)

- 5q ALK (2p23) BCL6 (3q27) BCR/ABL1, t(9;22) JAK2 (9p24)
BCR/ABL1, if neg reflex to JAK2 V617F Qual, If JAK2 neg reflex to CALR and MPL
CBFB (inv16) CCND1/IGH, t(11;14) IGH/BCL2, t(14;18) IGH/MYC, t(8;14)
MALT1 (18q21) PML/RARA, t(15;17) KMT2A (MLL; 11q23) RUNX1/RUNX1T1, t(8;21)
TRAFD (14q11.2) TP53 (17p-) MYC (8q24)
Other FISH, specify: _____

MOLECULAR

Labcorp NGS Tests (bone marrow aspirate, peripheral blood or fresh tissue)

- Labcorp Myeloid NGS
Labcorp Lymphoid NGS
Labcorp Pan-Heme NGS

This patient meets the following medical necessity criteria for this test (please mark/complete all that apply):

- Undefined cytopenia for greater than ___ months and other possible causes have been reasonably excluded
The working/clinical diagnosis is (mark all that apply):
AML MDS MPN Other (i.e., CLL, ALL) _____

See oncology.labcorp.com for a full gene list for each panel

Reveal SNP Microarray (If suspect balanced translocations, run cytogenetics and/or FISH)

- SNP Microarray for ALL, AML, CLL, MDS and other Hematologic Malignancies

Indication: _____

- FISH + SNP Microarray for Multiple Myeloma SNP Microarray for Multiple Myeloma
If MM (FISH+SNP) is ordered, probes t(4;14), t(11;14), t(14;16) are performed

Acute Leukemia Lymphoid Neoplasm MPN/CML/Mastocytosis

- Rapid AML Panel+ IDH 1/2 Mutation CEBPA Mutation NPM1 Mutation PML/RARA (Quantitative) cKIT Mutation FLT3 Mutation
B-cell Rearrangement IGH/IGK T-cell Rearrangement TRG/TRB B-cell Rearrangement IGH B-cell Rearrangement IGK T-cell Rearrangement TRG T-cell Rearrangement TRB IGHV Somatic Hypermutation p53 (CLL/B-cell ONLY) BRAF Mutation MYD88 Mutation
BCR/ABL1 Quantitative ABL Kinase Domain Mutation (BCR/ABL1 will be run) JAK2 V617F Mutation
Qualitative Quantitative if negative reflex to: CALR JAK2 Exon 12-15 MPL 515
JAK2 Exon 12-15 Mutation MPL 515 Mutation CALR Mutation KIT D816V Mutation Digital PCR-Systemic Mastocytosis

Other Molecular, specify: _____

SPECIAL CHEMISTRY (Serum ONLY)

Multiple Myeloma Diagnostic: *Meets IMWG Guidelines

- 120256 Immunofixation (sIFE), Protein Electrophoresis (SPE), Quant Free K/Lambda Light Chains (sFLC)
123200 Multiple Myeloma Cascade, SPE Reflex to sIFE and sFLC

Multiple Myeloma Monitoring:

- 001495 sIFE, SPE 001487 SPE 001685 sIFE
123218 sIFE DARZALEX (daratumumab patients ONLY) 123062 sIFE SARCLISA (isatuximab patients ONLY)
121137 sFLC, Quantitative Free Light K/Lambda Chains plus Ratio

Peripheral blood only
*If sending DNA, the lab only accepts isolated or extracted nucleic acids for which extraction or isolation is performed in an appropriately qualified CLIA or CAP/CMS equivalent laboratory.
†Informed consent is required for non-oncology genetics testing for New York state patients
+Rapid AML Panel includes FLT3 mutation, IDH 1/2 mutation and NPM1 mutation analyses
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Phone Number:
Med. Rec. # / Patient #:

BILLING INFORMATION (attach face sheet and copy of insurance card - both sides)
Bill: My Account Insurance Medicare Medicaid Patient Workers Comp
Patient Hospital Status: In-Patient Out-Patient Non-Patient
Insurance Information: See attached Authorization #
PRIMARY BILLING PARTY SECONDARY BILLING PARTY
INSURANCE CARRIER ID # GROUP # INSURANCE ADDRESS NAME OF INSURED PERSON RELATIONSHIP TO PATIENT EMPLOYER NAME

SPECIMEN INFORMATION
Collection Date: Time: AM PM
Specimen ID #(s):
Body Site/Descriptor:
Fixative: 10% Neutral Buffered Formalin Other: Hours Fixed:
Specimen Type: Smears:
BM Aspirate Fluid: Peripheral Blood #
BM Clot FNA: BM Touch Preps #
BM Core CSF BM Aspirate #
Dry Tap Lymph Node: Effusion #/Source
Peripheral Blood Slides # Fresh Tissue #/Site
If slide procurement required, indicate below:
Facility Name:
Address:
Phone Number: Fax Number:

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High Grade Multiple Myeloma AML MDS
B-cell Lymphoma MPN/CML MPN w/ Eosinophilia

Pediatric (COG) COG Single Probes

- ALL (Std Risk) ALL (High Risk) AML
ABL1 ABL2 PDGFRb
Individual Probes (for a complete list of probes visit oncology.labcorp.com)
5q ALK (2p23) BCL6 (3q27) BCR/ABL1, t(9;22) JAK2 (9p24)
BCR/ABL1, if neg reflex to JAK2 V617F Qual, If JAK2 neg reflex to CALR and MPL
CBFB (inv16) CCND1/IGH, t(11;14) IGH/BCL2, t(14;18) IGH/MYC, t(8;14)
MALT1 (18q21) PML/RARA, t(15;17) KMT2A (MLL; 11q23) RUNX1/RUNX1T1, t(8;21)
TRAFD (14q11.2) TP53 (17p-) MYC (8q24)
Other FISH, specify:

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Undefined cytopenia for greater than ___ months and other possible causes have been reasonably excluded
The working/clinical diagnosis is (mark all that apply):
AML MDS MPN Other (i.e., CLL, ALL)
See oncology.labcorp.com for a full gene list for each panel

- Reveal SNP Microarray: If suspect balanced translocations, run cytogenetics and/or FISH
SNP Microarray for ALL, AML, CLL, MDS and other Hematologic Malignancies
Indication:
FISH + SNP Microarray for Multiple Myeloma SNP Microarray for Multiple Myeloma
If MM (FISH+SNP) is ordered, probes t(4;14), t(11;14), t(14;16) are performed

- Acute Leukemia: Rapid AML Panel+, IDH 1/2 Mutation, CEBA Mutation, NPM1 Mutation, PML/RARA (Quantitative), cKIT Mutation, FLT3 Mutation
Lymphoid Neoplasm: B-cell Rearrangement IGH/IGK, T-cell Rearrangement TRG/TRB, B-cell Rearrangement IGH, B-cell Rearrangement IGK, T-cell Rearrangement TRG, T-cell Rearrangement TRB, IGHV Somatic Hypermutation, p53 (CLL/B-cell ONLY), BRAF Mutation, MYD88 Mutation
MPN/CML/Mastocytosis: BCR/ABL1 Quantitative, ABL Kinase Domain Mutation (BCR/ABL1 will be run), JAK2 V617F Mutation, Qualitative Quantitative if negative reflex to: CALR, JAK2 Exon 12-15, MPL 515, JAK2 Exon 12-15 Mutation, MPL 515 Mutation, CALR Mutation, KIT D816V Mutation Digital PCR-Systemic Mastocytosis
Other Molecular, specify:

SPECIAL CHEMISTRY (Serum ONLY)

- Multiple Myeloma Diagnostic: 120256 Immunofixation (sIFE), Protein Electrophoresis (SPE), Quant Free K/Lambda Light Chains (sFLC)*
123200 Multiple Myeloma Cascade, SPE Reflex to sIFE and sFLC
Multiple Myeloma Monitoring: 001495 sIFE, SPE 001487 SPE 001685 sIFE
123218 sIFE DARZALEX (daratumumab patients ONLY) 123062 sIFE SARCLISA (isatuximab patients ONLY)
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Peripheral blood only
*If sending DNA, the lab only accepts isolated or extracted nucleic acids for which extraction or isolation is performed in an appropriately qualified CLIA or CAP/CMS equivalent laboratory.
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B2A

| Test Reflex Guidelines | | | |
|---|----------------------------------|--|---|
| Disease Category | Timing | Findings (Morphology, Flow cytometry, FISH and/or karyotyping) | Tests to Perform |
| ALL | Initial Diagnosis; Follow-up* | ALL | Pediatric FISH Profile (<= 18 yrs or up to 30 yrs if treated in pediatric oncology setting) or Adult FISH Profile (>22 years); Reveal® SNP Array; Labcorp Lymphoid NGS depending on clinical presentation |
| AML | Initial Diagnosis | AML or borderline AML (MDS/AML) | FISH probes for RUNX1T1/RUNX1 t(8;21), CBFB inv(16), or PML/RARA t(15;17) or KMT2A/MLL respectively, as indicated; Labcorp Myeloid NGS + FLT3 and IDH1/2 testing |
| AML | Relapse | Findings indicative of relapse | Labcorp Myeloid NGS |
| CLL (peripheral blood/bone marrow) | Initial Diagnosis; Follow-up* | CD5+ neoplasm with classic or variant CLL features; features of refractory disease or disease progression/transformation# | CLL FISH profile; TP53 mutation analysis; and IGHV Somatic Hypermutation#; Labcorp Lymphoid NGS depending on clinical presentation |
| CML | Initial Diagnosis | Compatible or diagnostic findings for CML | FISH for BCR/ABL1 and/or RT-PCR Quantitative and cytogenetics |
| CML | Follow-up* | Prior diagnosis of CML | Quantitative BCR/ABL1 assay; add ABL Kinase mutation analysis if features of progression, discuss addition of Labcorp Myeloid NGS panel with client or place comment in report |
| MPN | Initial Diagnosis; Follow-up* | Morphologic features of MPN, but negative for JAK2 V617F, CALR, and MPL mutations; History of MPN with features of progression | Labcorp Myeloid NGS |
| MDS/MPN | Initial Diagnosis | Findings suspicious for MDS/MPN | Labcorp Myeloid NGS |
| MDS | Initial Diagnosis | Morphologic diagnosis of MDS | Labcorp Myeloid NGS |
| Plasma cell neoplasia | Initial Diagnosis; Follow-up* | evidence of abnormal/monotypic plasma cells | Myeloma FISH profile |
| SLL | Initial Diagnosis; Follow-up* | SLL identified in tissue sample by flow cytometry with 5% or more neoplastic cells | CLL FISH profile; Labcorp Lymphoid NGS depending on clinical presentation |
| B-cell lymphoma | Initial Diagnosis; Follow-up* | Findings suspicious or diagnostic for B-cell lymphoma, but with equivocal findings with regard to subclassification (for tissue cases 5% or more abnormal B-cells by flow cytometry; for peripheral blood/bone marrow cases, 10% or more abnormal B-cells) | FISH probes from NHL FISH panel and molecular assays as indicated; SNP micro-array to detect 11q abnormalities as needed; Labcorp Lymphoid NGS depending on clinical presentation |
| Large B-cell lymphoma or Burkitt lymphoma | Initial Diagnosis; Follow-up* | 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; Labcorp Lymphoid NGS depending on clinical presentation |
| Eosinophilia | Initial Diagnosis | peripheral blood or bone marrow with increased eosinophils | FISH probes for PDGFRA (4q); PDGFRB (5q); and FGFR1 (8q) |
| Hairy Cell Leukemia (HCL) | Initial Diagnosis; Follow-up* | abnormal/monotypic B-cells with features indicating HCL in the differential diagnosis | BRAF mutation; Labcorp Lymphoid NGS depending on clinical presentation |
| Lymphoplasmacytic Lymphoma (LPL) | Initial Diagnosis; Follow-up* | abnormal/monotypic B-cells with features indicating LPL in the differential diagnosis | MYD88 mutation; Labcorp Lymphoid NGS depending on clinical presentation |
| Mantle cell lymphoma (MCL) | Initial Diagnosis; Follow-up* | abnormal/monotypic B-cells with features indicating MCL in the differential diagnosis | FISH probe for CCND1/IGH t(11;14); TP53 mutation analysis; Labcorp Lymphoid NGS depending on clinical presentation |
| Mastocytosis | Initial Diagnosis | Atypical mast cells | High-sensitivity KIT D816V mutation analysis for mast cell disease |
| T-cell lymphoma/leukemia | Initial Diagnosis; Follow-up* | Atypical T-cells diagnostic or suspicious for T-cell lymphoma/leukemia | TCR gene rearrangement; ALK FISH probe for CD30+ cases, as indicated; cytogenetic karyotyping if material adequate; Labcorp Lymphoid NGS depending on clinical presentation |

Testing may vary from this table depending on clinical and morphologic context.

* recommendation for follow-up evaluation requires that prior material was evaluated in an Labcorp Oncology (LCO) facility

#IGHV will not be performed on follow-up

¹ AZ/TN ² CT

| Morphologic Evaluation Common Components (Please include patient CBC report) | | | |
|--|----------------|---------------------------|--|
| • Peripheral Blood Interpretation (85060) | • Clot (88305) | • Core (88305) | • Additional Studies/Special Stains (88313) – Iron and Reticulin |
| • Bone Marrow Aspirate Smear & Interpretation (85097) | | • Decalcification (88311) | • IHC Global marker number (88342) varies but typically 0-4 |

| Flow Cytometry* | | |
|---|---|--|
| Leukemia/lymphoma phenotyping panel (peripheral blood/bone marrow) 21 * [Ⓞ] antibodies CD2, CD3, CD4, CD5, CD7, CD8, CD10, CD11b, CD13, CD14, CD16, CD19, CD20, CD23, CD57, CD33, CD34, CD38, CD45, CD56, CD64, HLA-DR, kappa light chain, lambda light chain | Tissue/fluids panel 21 * [Ⓞ] antibodies CD2, CD3, CD4, CD5, CD7, CD8, CD10, CD11b, CD19, CD20, CD23, CD30, CD38, CD43, CD45, CD56, CD57, FMC-7, HLA-DR, kappa light chain, lambda light chain | PNH Evaluation CD14, CD15, CD24, CD45, CD64, FLAER. CD59 and CD235a may be added at discretion of reviewing pathologist |

*Additional antibodies may be added if determined to be medically necessary to render a diagnosis in the opinion of the reviewing pathologist

[Ⓞ]Antibodies performed determined by testing facility and may vary from the list above. Performed antibodies will appear in the patient report.

| FISH (disease state profile OR individual probes) | | | | | | |
|---|---|--|--|--|--|--|
| ALL (Adult) BCR/ABL1, t(9;22) KMT2A (MLL; 11q23) MYC (8q24) 6 21q | ALL (Pediatric/Std Risk) BCR/ABL1, t(9;22) 4 10 17 KMT2A (MLL; 11q23) CDKN2A (p16) TCF3 (E2A) ETV6/RUNX1, t(12;21) | AML PML/RARA, t(15;17) CBFB, inv(16) RUNX1T1/RUNX1, t(8;21) 5q 7q KMT2A (MLL) | CLL TP53 (17p-) ATM (11q-) CCND1/IGH, t(11;14) 13q14 (DLEU) 12 | MPN/CML 20q 8 9 13q14 (DLEU) BCR/ABL1, t(9;22) | Multiple Myeloma Monosomy 13/13q- TP53 (17p-) 7 9 15 CCND1/IGH, t(11;14) CKS1B (1q21) FGFR3/IGH, t(4;14) IGH/MAF, t(14;16) | NHL (Individual Probes) ALK (2p23) BCL6 (3q27) CCND1/IGH, t(11;14) IGH/BCL2, t(14;18) IGH/MYC, t(8;14) MALT1 (18q21) TRA/D (14q11.2) MYC (8q24) BCL2 (18q21) |
| ALL (Philadelphia-like) CRLF2 ABL1 ABL2 JAK2 PDGFRB | ALL (High Risk) includes the above probes PLUS: ABL1 ABL2 PDGFRB | High grade B-cell Lymphoma BCL2 (18q21) BCL6 (3q27) MYC (8q24) | MDS 5q 7q 20q 8 | MPN with Eosinophilia FGFR1 PDGFRA PDGFRB JAK2 | | |

Note: *1 in genotype results denotes detection of the normal (reference) sequence at all the variant sites assessed.

| SERUM - Multiple Myeloma Cascade, Protein Electrophoresis (SPE) reflex to Immunofixation (sIFE) and Free Light Chain (sFLC) for interpretation, refer to www.labcorp.com |
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onc-711-v29 02142025

| Lab Locations | | |
|---|---|---------------------------------------|
| Accupath Diagnostic Laboratories, Inc. | | Esoterix Genetic Laboratories, LLC |
| 201 Summit View Drive, Suite 100 Brentwood, TN 37027 | 5005 South 40th Street Phoenix, AZ 85040 | 3 Forest Parkway Shelton, CT 06484 |