

ONCOLOGY GENETIC TESTING REQUISITION

All Information Must Be Completed Before Sample Can Be Processed

PATIENT INFORMATION

Patient Name: _____, _____, _____
Last First MI

Address: _____

Home Phone: _____ MR#: _____

Date of Birth: ____/____/____ Gender: ☐ Male ☐ Female

INDICATIONS/DIAGNOSIS/ICD-9 CODE

- | | | | |
|---|--|--|---|
| ☐ Acute Myelogenous Leukemia | ☐ Glioma | ☐ Lymphoproliferative Disorder | ☐ Pancytopenia |
| ☐ Acute Promyelocytic Leukemia | ☐ Hodgkin Lymphoma | ☐ Malignant Melanoma | ☐ Polycythemia Vera (PV) |
| ☐ Adenopathy | ☐ Langerhans Cell Histiocytosis (LCH) | ☐ Medulloblastoma | ☐ Sarcoma |
| ☐ Anemia | ☐ Leukemia | ☐ Monoclonal Gammopathy | ☐ Thrombocytopenia |
| ☐ Burkitt Lymphoma | ☐ Leukocytosis | ☐ Multiple Myeloma | ☐ Thrombocytosis |
| ☐ Chronic Lymphocytic Leukemia | ☐ Leukopenia | ☐ Myelodysplastic Syndrome/Disease (MDS) | ☐ Wilms Tumor |
| ☐ Chronic Myelogenous Leukemia | ☐ Lung Cancer | ☐ Myeloproliferative Disease (MPS or MPD) | ☐ Other _____ |
| ☐ Colorectal Cancer | ☐ Lymphocytosis | ☐ Neutropenia | _____ |
| ☐ Ewing Sarcoma | ☐ Lymphoma | ☐ Non-Hodgkin Lymphoma (NHL) | _____ |

ETHNIC/RACIAL BACKGROUND (Choose All)

- | | |
|--|--|
| ☐ European American (White) | ☐ African-American (Black) |
| ☐ Native American or Alaskan | ☐ Asian-American |
| ☐ Pacific Islander | ☐ Ashkenazi Jewish ancestry |
| ☐ Latinx-Hispanic _____
(specify country/region of origin) | |
| ☐ Other _____
(specify country/region of origin) | |

REFERRING PHYSICIAN

Physician Name (print): _____

Address: _____

Phone: (_____) _____ Fax: (_____) _____

Email: _____

Genetic Counselor/Lab Contact Name: _____

Phone: (_____) _____ Fax: (_____) _____

Email: _____

Referring Physician Signature (REQUIRED) Date: ____/____/____

Contact Information for Results/Questions (if different than ordering provider) :

Name & Title: _____

Phone: (_____) _____ Fax: (_____) _____

Email: _____

BILLING INFORMATION (Choose ONE method of payment):

☐ REFERRING INSTITUTION

Institution: _____

Address: _____

City/State/Zip: _____

Accounts Payable Contact Name: _____

Phone: _____

Fax: _____

Email: _____

☐ COMMERCIAL INSURANCE

Insurance can only be billed if requested at the time of service.

Policy Holder Name: _____

Gender: _____ Date of Birth ____/____/____

Authorization Number: _____

Insurance ID Number: _____

Insurance Name: _____

Insurance Address: _____

City/State/Zip: _____

Insurance Phone Number: _____

See page 3 for more billing information

SAMPLE/SPECIMEN INFORMATION

Has patient received a bone marrow transplant? ☐ Yes ☐ No

If yes, date of bone marrow transplant _____

Percent engraftment _____

Specimen Date: _____ Time: _____

DRAWN BY: _____

*Phlebotomist must initial tube of specimen to confirm sample identity

DISEASE STATUS:

☐ New diagnosis ☐ Remission ☐ Relapse ☐ E(COG) study ☐ COG patient

SPECIMEN TYPE — SEE PAGE 3 FOR SPECIMEN REQUIREMENTS

☐ Bone marrow ☐ Oncology blood ☐ Lymph node

☐ Formalin fixed paraffin embedded tissue ☐ Touch prep ☐ Smear

Estimated percent of tumor in sample: _____

☐ Solid tumor (specify): _____ ☐ If in media, type: _____

Estimated percent of tumor in sample: _____

Other: _____ WBC _____ % Blasts _____

TEST(S) REQUESTED

SEE PAGE 3 FOR SPECIMEN AND TEST DETAILS

Cytogenetic Chromosome and Microarray Analysis

☐ Oncology Chromosome Analysis

☐ Constitutional (blood) Chromosome Analysis

☐ Oncology Microarray

[Additional 3 mL blood or bone marrow (NaHep) if ordered without chromosomes] — % Tumor: _____

FISH

(Additional FISH probes available. See page 3 for details.)

☐ t(9;22) [BCR/ABL1]

☐ 11q23 [KMT2A]

☐ X/Y [Opposite sex BMT]

☐ t(15;17) [PML/RARα]

☐ Other (please call lab) _____

Hematologic FISH Panels

(All probes available individually. Please see page 3 for panel descriptions)

☐ ALL Hyperdiploid

☐ Fanconi anemia

☐ ALL Risk Stratification

☐ Multiple myeloma

☐ B-Cell ALL

☐ MDS

☐ Ph-like ALL

☐ Myeloid Malignancy

☐ AML

☐ MPD

☐ APL

☐ SDS

☐ Burkitt Lymphoma

☐ Large cell NHL

☐ CLL

☐ Small cell NHL

☐ Double Hit Lymphoma

☐ Combined NHL

☐ Eosinophilia

☐ T-Cell Lymphoma/Leukemia

Non-Hematologic FISH

Fresh tumor or FFPE slides (include 1 marked H&E slide with FFPE)

See page 3 for specimen requirements

☐ BRAF(7q34) FISH

☐ Ependymoma FISH Panel

☐ High-Grade Glioma FISH Panel

☐ Low-Grade Glioma FISH Panel

☐ Lung Cancer FISH Panel

☐ Medulloblastoma FISH Panel

☐ Melanocytic Tumor FISH Panel

☐ Pilocytic Astrocytoma FISH Panel

Molecular Genetic Analysis (RNA assays)

Samples must be received within 24 hours of collection.

☐ BCR/ABL - QUANTITATIVE (p210)

☐ BCR/ABL - QUANTITATIVE (p190)

☐ BCR/ABL - RT-PCR (QUALITATIVE)

Molecular Genetic Analysis (DNA assays)

Samples must be received within 48 hours of collection.

☐ JAK2 QUANTITATIVE (V617F)

☐ PTEN sequencing

☐ Bone marrow engraftment (BME) by STR (Same sex donor & recipient)

☐ Pre-transplant host sample

☐ Post transplant sample

☐ Donor sample

☐ WBC sub-populations engraftment study* [Select cell type(s) and prioritize below]

☐ STR (same sex) ☐ X/Y FISH (opposite sex)

☐ T cells (CD3+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

☐ Myeloid cells (CD15+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

☐ B cells (CD19+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

☐ NK cells (CD56+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

☐ Cell Separation (for non-engraftment testing)* [Select cell type(s) and prioritize below]

☐ T cells (CD3+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

☐ Myeloid cells (CD15+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

☐ B cells (CD19+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

☐ NK cells (CD56+) - Priority: ☐ 1st ☐ 2nd ☐ 3rd ☐ 4th

You MUST call the GENETICS LAB at 513-636-4474 to schedule this test prior to sample submission.

Molecular Pathology PCR Analysis (contact 513-636-9820 with questions)

☐ EBV Quantitative PCR

Non-Hematologic Genetic Analysis

☐ MAP2K1 full gene sequence analysis (Langerhans cell histiocytosis, colon, lung, melanoma) — % Tumor: _____

Please see the **Pediatric/Adult Requisition** for
Chromosome Breakage Study for Fanconi Anemia or contact the lab at:
www.cincinnatichildrens.org/cytogenetics or 513-636-4474

SPECIMEN REQUIREMENTS

Cytogenetic Analysis (Chromosome, FISH, and Microarray analysis):

3 mL blood or bone marrow (NaHep)

Chromosome analysis
Cell culture only
FISH probes and FISH panels

3 mL blood or bone marrow (EDTA)

Oncology microarray

Fresh Tumor or Lymph Nodes (1cm³ in sterile saline or sterile transport media)

Chromosome analysis
Cell culture
FISH probes and FISH panels

Molecular Genetic Analysis (RNA Assays): 5-10 mL blood or 3–5 mL bone marrow (EDTA) — Samples must be received within 24 hours of collection.

BCR/ABL — Quantitative (p210), *BCR/ABL* — Quantitative (p190), *BCR/ABL*

Molecular Genetics Analysis (DNA Assays): 3 mL bone marrow or blood (EDTA) — Samples must be received within 48 hours of collection.

JAK2 Quantitative (V617F), *PTEN* Seq, Bone marrow engraftment by STR, WBC sorted sub-populations engraftment study (by STR or FISH)

Non-Hematologic Genetic Analysis

MAP2K1 full gene sequencing: 3 mL blood or bone marrow (EDTA), or 1 cm³ fresh tumor.

Molecular Pathology PCR Analysis (contact 513-636-9820 with questions)

EBV Quantitative PCR: For plasma: 1 mL blood in lavender top (EDTA) , For serum: 1 mL blood in gold top, 1 mL bone marrow in lavender top (EDTA), 1 mL CSF in sterile container, 0.3 g fresh tissue in sterile container

FISH (Fluorescence In Situ Hybridization)

NOTE: All FISH probes are available for individual testing

Hematologic FISH Panels — 3 mL blood or bone marrow (NaHep)

- ALL Hyperdiploid: trisomy 4, 10, 17
- ALL Risk Stratification: 4, 10, 17, t(1;19), t(12;21), t(9;22), *KMT2A*
- B-Cell ALL: *CRLF2*, t(1;19), *ABL2*, 4/10/17, *CSF1R*, *PDGFRB*, *JAK2*, *ABL1*, t(9;22), *KMT2A*, t(12;21), *IGH*, *EPOR*
- Ph-like ALL: *CRLF2*, *ABL2*, *PDGFRB*, *CSF1R*, *JAK2*, *ABL1*, *EPOR*
- AML: t(6;9), t(8;21), *NUP98*, *KMT2A*, inv(16)
- APL: t(15;17), *RARα*
- Burkitt Lymphoma: t(8;14), *MYC*
- CLL: 13q14.3, 13q34, 12 centromere, *ATM*, *TP53*, t(11;14)
- Double Hit Lymphoma: *BCL6*, *MYC*, t(8;14), t(14;18)
- Eosinophilia: 4q12, *PDGFRB*, *FGFR1*, *CBFB*
- Fanconi Anemia: 1q25, 3q27, mono 7 / del(7q)
- Multiple Myeloma (CD138+): 1p32.3/1q21, t(4;14), t(11;14), monosomy 13/ del 13q, t(14;16), t(14;20), *TP53*
- MDS: mono 5/del 5q, mono 7/del 7q, tri 8, *TP53*, del (20q)
- Myeloid Malignancy: mono 5/del 5q, t(6;9), mono 7/del 7q, tri 8, t(8;21), *NUP98*, *KMT2A*, *ETV6*, t(15;17), inv(16), *TP53*, del (20q)

- MPD: 4q12, *PDGFRB*, *FGFR1*, *BCR/ABL1*
- SDS: mono 7/del 7q, tri 8, del (20q)
- Large cell NHL: t(11;14), t(14;18), *TP53*, *BCL6*, *ALK*
- Small B-cell NHL: t(11;14), t(14;18), 18q21 (*MALT1*), CLL Panel
- Combined NHL: (large and small cell NHL panels)
- T-Cell Lymphoma: *TRA/TRD*, *TRB*, *TRG*, *BCR/ABL1*, *KMT2A*

Non-Hematologic FISH Panels — 4-8 FFPE slides cut to 4 micron thickness and 1 marked H & E slide** — Fresh Tumor (1cm³)

- Ependymoma: *ABL2*, *CDKN2A*, *C11orf95*, *RELA*, *YAP1*
- High-Grade Glioma: *PDGFRA*, *CDKN2A*, *NTRK2*, *MYCN*
- Low-Grade Glioma: *TP73/ABL2*, *FGFR1*, *MYB*, *BRAF*, *MYBL1*
- Lung Cancer: *ALK*, *ROS1*, *MET*, *RET*
- Medulloblastoma : *MYB*, *LIS1/RARα*, *MYC*, *MYCN*
- Melanocytic Tumor: *RREB1*, *MYC*, *CDKN2A*, *CCND1*
- Pilocytic Astrocytoma: *BRAF*, *CDKN2A**

*For each probe ordered, send 2 unstained slides with one section cut to 4 micron thickness and mounted on a charged slide. Blocks are also accepted for processing.

*Pilocytic Astrocytoma FISH Panel only needs 2-4 FFPE slides and 1 marked H&E slide

SHIPPING INFORMATION

Local courier is available; please call 513-636-4474 for information.

Shipping for samples that arrive Monday–Saturday:

Cincinnati Children's
Genetics and Genomics Diagnostic Laboratory
3333 Burnet Ave. TCHRF 1042
DOCK 5
Cincinnati, OH 45229-3039

BILLING INFORMATION

* PLEASE NOTE:

- We will not bill Medicaid, Medicaid HMO, or Medicare except for the following: CCHMC Patients, CCHMC Providers, or Designated Regional Counties.
- If you have questions, please call 1-866-450-4198 for complete details.

☐ Patient Signed Completed ABN:

Medical Necessity Regulations: At the government's request, the Molecular Genetics Laboratories would like to remind all physicians that when ordering tests that will be paid under federal health care programs, including Medicare and Medicaid programs, that these programs will pay only for those tests the relevant program deems to be (1) included as covered services, (2) reasonable, (3) medically necessary for the treatment and diagnosis of the patient, and (4) not for screening purposes.