

Last Name: \_\_\_\_\_ First Name: \_\_\_\_\_

Sex: \_\_\_\_\_ Date of Birth: \_\_\_\_\_ PT#: \_\_\_\_\_

SSN: \_\_\_\_\_ Phone (Day): \_\_\_\_\_ Phone (Eve): \_\_\_\_\_

Address: \_\_\_\_\_

Charge to (please attach admissions face sheet, completed insurance forms, and/or a copy of insurance card.  
If not available, please complete the following):

Insurance Company: \_\_\_\_\_ Phone: \_\_\_\_\_

Address: \_\_\_\_\_

Policy Holder: \_\_\_\_\_ ID#: \_\_\_\_\_ Group#: \_\_\_\_\_

Inpatient  Outpatient Hospital Name \_\_\_\_\_  Nonpatient

## SPECIMEN TYPE

DATE SAMPLE DRAWN: \_\_\_\_\_ TIME SAMPLE DRAWN: \_\_\_\_\_

Bone Marrow Aspirate      Bone Core Biopsy      Peripheral Blood, Leukemic      Lymph Node

Other (please describe): \_\_\_\_\_

**CLINICAL INFORMATION/ICD-9\$** (required): \_\_\_\_\_

New Diagnosis       Relapse       Monitoring

Radiation Therapy \_\_\_\_\_  Chemotherapy \_\_\_\_\_

Bone Marrow Transplant       Autologous       Allogenic       Sex Mismatch

## TEST REQUESTED

Chromosome Analysis      Hold Culture for Testing

Flow Cytometry\*\*       Molecular Analysis\*\* \_\_\_\_\_

FISH (Please mark below)\* For paraffin embedded tissue, please see our FFPE request form.

ALL AML CLL CML MDS MM MPD Eosinophilia NHL Other:

REFLEX FISH IF CHROMOSOME RESULTS ARE NORMAL (Please mark below)\*

ALL AML CLL CML MDS MM MPD Eosinophilia NHL Other:

OTHER \_\_\_\_\_

\*Please complete second page for other testing

\*\*Sent out to reference laboratory

Physician: \_\_\_\_\_

Referring Hospital/Lab: \_\_\_\_\_

Address: \_\_\_\_\_

Phone: \_\_\_\_\_ Fax: \_\_\_\_\_

**ADDITIONAL INFORMATION FOR FISH** (please see our website for our most current FISH offerings)

Patient Last Name: \_\_\_\_\_ First Name: \_\_\_\_\_

**FISH**

● **MDS**

- MDS Panel:** -5/del(5q), -7/del(7q), +8, del(20q)
- Monosomy 5/deletion 5q (EGR1)
- Monosomy 7/deletion 7q (D7S522/D7Z1)
- Trisomy 8 (D8Z2)
- Deletion 20q (D20S108)
- Other \_\_\_\_\_

● **CLL**

- CLL Panel:** del(11q), del(17p), +12, -13/del(13q)
- Deletion 11q (ATM)
- Deletion 17p (TP53)
- Trisomy 12 (D12Z3)
- Monosomy 13/Deletion 13q (D13S319/LAMP1)
- Translocation 11;14 (IGH/CCND1)
- Other \_\_\_\_\_

● **MM/Plasma cell dyscrasia**

**MM Panel:** 1p/1q, t(4;14), -13/del(13q), del(17p)

- Deletion 1p32.3/gain of 1q21 (CKS1B/CDKN2C)
- Monosomy 13/Deletion 13q (D13S319/LAMP1)
- Deletion 17p (TP53/CEP17)
- 14q32 rearrangements (IGH breakapart)
- Translocation 4;14 (IGH/FGFR3)
- Translocation 11;14 (IGH/CCND1)
- Translocation 14;16 (IGH/MAF)
- Hyperdiploidy of chromosomes 5, 9, 15
- Other \_\_\_\_\_

● **CML**

- Translocation 9;22 (BCR/ABL1)
- Deletion 9q34 (ASS)
- Other \_\_\_\_\_

● **AML**

**AML Panel:** -7/del(7q), +8, t(8;21), 11q23 rearrangements, t(15;17), inv(16)/t(16;16)

- Monosomy 7/deletion 7q (D7S522/D7Z1)
- Trisomy 8 (D8Z2)
- Translocation 8;21 (ETO/RUNX1)
- 11q23 rearrangements (MLL)
- Translocation 15;17 (PML/RARA)
- Inversion 16/translocation 16 (CBFB)
- Other \_\_\_\_\_

● **ALL**

- ALL Panel:** t(9;22), 11q23 rearrangements, t(12;21), del(9p),t(8;14)
- Translocation 9;22 (BCR/ABL1)
- 11q23 rearrangements (MLL)
- Translocation 12;21 (ETV6/RUNX1)
- Translocation 8;14 (IGH/MYC/D8Z2)
- Deletion 9p21 (CDKN2A)
- Other \_\_\_\_\_

● **LYMPHOMA**

**NHL Panel:** 3q27 rearrangements, 8q24 rearrangements, t(11;14), t(11;18), t(14;18)

- 3q27 rearrangements (BCL6)
- 8q24 rearrangements (MYC)
- Translocation 11;14 (IGH/CCND1)
- Translocation 11;18 (BIRC3/MALT1)
- ↳ Translocation 14;18 (IGH/BCL2)
- Other \_\_\_\_\_

● **MPD**

**MPD Panel:** 4q12 rearrangements, 5q33rearrangements,+8, t(9;22), del(13q), del(20q) 4q12 rearrangements (PDGFRA-FIP1L1) 5q33 rearrangements (PDGFRB) Trisomy 8 (D8Z2) Translocation 9;22 (BCR/ABL1) Deletion 13q (D13S319/LAMP1) Deletion 20q (D20S108) Other \_\_\_\_\_

● **EOSINOPHILIA**

**Eosinophilia Panel:** 4q12 rearrangements, 5q33 rearrangements, 8p11.23-p11.22 rearrangements 4q12 rearrangements (PDGFRA-FIP1L1) 5q33 rearrangements (PDGFRB) 8p11.23-p11.22 rearrangements (FGFR1) Other \_\_\_\_\_

● **OTHER**

- 2p23 rearrangements (ALK)
- 3q26.2 rearrangements (EVI1)
- 6q23 deletion (MYB)
- Translocation 8;9 (PCM1/JAK2)
- 12p13.1 rearrangements/deletions (ETV6)
- 13q14 rearrangements (FOXO1)
- 14q11.2 rearrangements (TCR alpha/delta)
- 17q21 rearrangements (RARA)
- 22q12 rearrangements (EWSR1)

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## SPECIMEN REQUIREMENTS AND SHIPPING

All specimens must be labeled with at least two patient identifiers and be accompanied by completed request form. All samples should be kept at room temperature and transported to the laboratory with minimum delay. Please call (800) 328-2026 if you have any questions.

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### NEOPLASIA CHROMOSOME ANALYSIS

**Bone Marrow:** Aspirate 1-2 ml bone marrow into a sterile syringe containing 0.1 ml preservative free sodium heparin. Invert syringe to mix. Transfer to a 3 ml preservative free sodium-heparin (green top) vacutainer

tube or sterile tube of transport media (we provide).

**Leukemic Peripheral Blood:** Patient should have WBC of 15,000 or higher with approximately 10% circulating immature myeloid or lymphoid blast cells. Collect 5 ml of peripheral blood in a preservative free sodium-heparin (green top) vacutainer tube.

**Solid Tumor Tissue:** >5 mm<sup>3</sup> representative tumor tissue collected under aseptic conditions and transported in sterile tissue culture media.

**Lymph Node Biopsy:** >5 mm<sup>3</sup> tumor biopsy collected under aseptic conditions and transported in sterile tissue culture media.

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### SHIPPING INFORMATION

Sample should be securely packaged and sent at room temperature to:

[Diagnostic Cytogenetics](#)

[2360 West Commodore Way Suite 201, Seattle, WA 98199](#)

To arrange pick up in the local Puget Sound area, please call (206) 328-2026 / (800) 328-2026. For overnight delivery service: Federal Express (800) 463-3339. Please call for our FedEx account number. Please send specimens by Standard Overnight Service. Specimens sent on Friday MUST be marked with a "Saturday Delivery" sticker. Please call the lab at (800) 328-2026 with the airbill number so that we may track your specimen.

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### FLUORESCENCE IN SITU HYBRIDIZATION (FISH)

FISH studies are indicated when classic cytogenetics alone cannot resolve an abnormality. Specimen collection is as described previously for the tissue to be studied.

**Paraffin-embedded tissue slides:** For each probe requested, a minimum of three slides of four-micron tissue sections from formalin-fixed, paraffin-embedded blocks on positively-charged slides. The area of interest should be marked. Pathology report and H&E slide of the tissue should be submitted.

If possible, please provide slides of normal tissue of the same type to be used as a control.

If paraffin blocks are submitted, there will be additional fees for processing.