

2020 **MOLECULAR CYTOGENETIC REFERRAL FORM FOR FISH ANALYSIS** 2020

Medical Genetics – Wake Forest Baptist Health – Winston-Salem NC
 wakehealth.edu/specialty/m/Medical-Genetics P: 336-716-4321 F: 336-716-2554

Collection Date: _____ Time: _____ am/pm Time of Fixation _____

Name: _____ / _____ / _____ / _____
 (Please print) Last First Middle Maiden
 Address: _____ / _____ / _____ / _____ Daytime Phone: (____) _____
 Mailing Address City State Zip
 Birth Date: _____ SS# or Patient ID #: _____ Sex: male female

Hospital: _____ Hospital/Unit #: _____
 Type of Specimen Amnio CVS Tissue PUBS POC Urine Blood Bone Marrow
 Solid Tumor Slides/Smears Paraffin Block Other: _____

Collection Technique: Green stoppered sodium heparin tube. Room temperature

Physician/Provider Order	
Physician: Last, First / Phone/beeper	
1.	
X.	
2.	
X.	

←ATTENDING SIGNATURE
 ←RESIDENT/OTHER SIGNATURE

Fill out form with at least 2 sample-matching patient identifiers and FISH probes needed. Email back to cytofish@wakehealth.edu Bring marked slides to Hanes Ground Floor G022 (FISH LAB)

Questions?
 Dr Pettenati: 3-7575
 Dr Lyalin: 3-7547
 FISH Lab: 6-2064

X. Physician Signature Required

Signs/Symptoms/Indication (ICD-10 Codes) for Chromosome Study

Indicate all that apply. Codes do not represent entire listing of ICD-109 codes available. Please consult current ICD-10 code book for complete listing.

LEUKEMIA		
<input type="checkbox"/> ALL (C91.00) <input type="checkbox"/> remission (.01)	<input type="checkbox"/> APL (C942.40) <input type="checkbox"/> remission (.01)	<input type="checkbox"/> CLL (C91.10)
<input type="checkbox"/> Acute Leukemia (C92.A0)	<input type="checkbox"/> Burkitt's Lymphoma unspec (C83.70)	<input type="checkbox"/> Hodgkin's Lymphoma unspec (C81.0)
<input type="checkbox"/> Leukemia unspec (C92.90)	<input type="checkbox"/> Non-Hodgkins (C85.50)	<input type="checkbox"/> Multiple Myeloma (C90.00) <input type="checkbox"/> remission (.01)
<input type="checkbox"/> AML (C92.00) <input type="checkbox"/> remission	<input type="checkbox"/> CML (C92.10) <input type="checkbox"/> remission (.01)	<input type="checkbox"/> Myeloproliferative Syndrome (C94.6)
<input type="checkbox"/>	<input type="checkbox"/> MDS (D46.Z)	<input type="checkbox"/> Thrombocythemia (D69.6)
	<input type="checkbox"/> Follicular lymphoma ((C82.0)	<input type="checkbox"/> Other (specify): _____

PRENATAL	POSTNATAL	OTHER
<input type="checkbox"/> Advanced Maternal Age (O09.519)	<input type="checkbox"/> Unspecified congenital anomaly (Q89.9)	<input type="checkbox"/> Malig of breast (C50.119)
<input type="checkbox"/> Inc. Down Syndrome Risk (028.9)	<input type="checkbox"/> Multiple congenital anomalies (Q89.7)	<input type="checkbox"/> Neoplasm of bladder (D49.4)
<input type="checkbox"/> Inc. Trisomy 18 Risk (028.9)	<input type="checkbox"/> Sex chromosome anomaly (Q52/55.9)	<input type="checkbox"/> Malign neoplasm of bladder wall NOS (C67.9)
<input type="checkbox"/> Suspect fetal chromosome abn (O35.1XX0)	<input type="checkbox"/> Cleft lip/palate (Q37.9)	<input type="checkbox"/> Post BMT transplant (Z94.81)
<input type="checkbox"/> Abn. Ultrasound – specify _____	<input type="checkbox"/> Ambiguous genitalia (Q51-55.9)	Specify original disease: _____
<input type="checkbox"/> Other (specify): _____	<input type="checkbox"/> Delay in sexual development (E30.0)	<input type="checkbox"/> Other (specify): _____

Test Requested Note: When ordering tests for which Medicare reimbursement will be sought, it is recommended that the Provider consult any Local Medical Review Policies (LMRP) or National Coverage Decisions (NCD) that may be applicable to the test(s) being ordered. Based on guidance issued in either of these policies it may be necessary to obtain an Advanced Beneficiary Notice (ABN) from the Medicare Patient Form Medicaid and other carriers a signed Statement of Financial Responsibility from the patient may be necessary. (See Statement of Financial Responsibility at top of form.)

FISH Probes – select as needed

Prenatal	Syndrome/Microdeletion	Leukemia / Tumor	Panels
<input type="checkbox"/> 13/16/18/21/XY	<input type="checkbox"/> Angelman 15q11	<input type="checkbox"/> inv(3q)	<input type="checkbox"/> B-ALL
<input type="checkbox"/> +13 LSI	<input type="checkbox"/> Cri-du-chat 5p15.2	<input type="checkbox"/> -5 / del 5q31	<input type="checkbox"/> Uro Vysion-Bladder
<input type="checkbox"/> +21 LSI	<input type="checkbox"/> DiGeorge/VCF 22q11	<input type="checkbox"/> -7 / del 7q31	<input type="checkbox"/> Her2/neu-breast / gastric
<input type="checkbox"/> +18 LSI	<input type="checkbox"/> Williams 7q11.23	<input type="checkbox"/> +8 (AML)	<input type="checkbox"/> CLL
<input type="checkbox"/> +/- X	<input type="checkbox"/> Kallmann Xp22.3	<input type="checkbox"/> (9;22-CML) BCR/ABL	<input type="checkbox"/> Mult Myeloma
<input type="checkbox"/> +/- Y	<input type="checkbox"/> Miller-Dieker 17p13.3	<input type="checkbox"/> BCR/ABL and ASS	<input type="checkbox"/> MDS 5q/7q/8
	<input type="checkbox"/> Prader-Willi 15q12	<input type="checkbox"/> inv (16)	<input type="checkbox"/> Eosinophilia
	<input type="checkbox"/> Retinoblastoma 13q14	<input type="checkbox"/> del 20q12.2	<input type="checkbox"/> Triple lymph
	<input type="checkbox"/> Smith-Magenis 17p11.2	<input type="checkbox"/> (15;17-APL) PML/RARA	
	<input type="checkbox"/> Wolf-Hirschhorn 4p16.3	<input type="checkbox"/> t(6;9) AML	
	<input type="checkbox"/> Trisomy 21	<input type="checkbox"/> t(8;21) AML RUNX1/RUNX1T1	
	<input type="checkbox"/> Trisomy 13	<input type="checkbox"/> 11q23 MLL	
	<input type="checkbox"/> Trisomy 18	<input type="checkbox"/> PDGFRA/CHIC2/FIPL1	
	<input type="checkbox"/> Turner 45, X	<input type="checkbox"/> PDGFRB	
	<input type="checkbox"/> Klinefelter 47, XXY	<input type="checkbox"/> X/Y (transplant)	
	<input type="checkbox"/> Sex Chromosome	<input type="checkbox"/> Other specify: _____	
	<input type="checkbox"/> SRY Yp11.3		
	<input type="checkbox"/> STS Xp22.3		