

2012 PERIPHERAL BLOOD REFERRAL FORM FOR CHROMOSOME ANALYSIS

Medical Genetics – Wake Forest University School of Medicine – Winston Salem NC

www.wfubmc.edu/medical genetics

Phone: 336-716-4321 Fax: 336-716-2554

Collection Date: _____ **Time:** _____ am/pm **WFU LAB #:** _____

Name: _____ / _____ / _____ / _____
 (Please print) Last First Middle Maiden

Address: _____ / _____ / _____ / _____ **Daytime Phone:** (____) _____
 Mailing Address City State Zip

Birth Date: _____ **SS#:** _____ **Sex:** male / female

Hospital: _____ **Hospital/Unit #:** _____ **Patient's Mother's first name:** _____

Type of Specimen: Blood PUBS Blood (POC) Cancer

Collection Technique: 5 – 10 mls blood collected in a green stoppered sodium heparin tube. **Keep at room temperature**

Physician/Provider Order	Statement of Financial Responsibility
Physician: Last, First / Phone / Beeper	I authorize any holder of medical or other information about me to release to my healthcare provider, third party processor, the Centers for Medicare and Medicaid Services or its intermediaries or carriers any information needed for this health care encounter or related claim. I permit a copy of this authorization to be used in place of the original, and request payment of authorized insurance benefits be made on my behalf to the WFU Physicians. I understand I am responsible for payment of these charges. I am also responsible for payment if my insurance carrier decides this is a non-covered service or requires prior authorization, which I did not obtain.
1. _____	
X Physician Signature Required	
2. _____	
3. _____	Patient Signature: _____ Date: _____

Billing Information
Bill: <input type="checkbox"/> Forsyth Hospital/Novant <input type="checkbox"/> Moses Cone - Spectrum <input type="checkbox"/> Solstas <input type="checkbox"/> Women's Hospt. of Greensboro <input type="checkbox"/> Wesley Long Hospital <input type="checkbox"/> Other : _____ <input type="checkbox"/> Medicare # _____ <input type="checkbox"/> Medicaid #: _____ Carolina Access #: _____ <input type="checkbox"/> Insurance: _____ Employer: _____ Policy #: _____ (Enclose copy of both sides of insurance card)

SIGNS/SYMPTOMS/INDICATION (ICD-9 CODES) FOR CHROMOSOME STUDY

- Indicate all that apply. Codes here do not represent entire listing of ICD-9 codes available. Please consult current ICD-9 code book for complete listing.
- | | | |
|---|--|---|
| <input type="checkbox"/> Other chromosome conditions (758.9) | <input type="checkbox"/> Anomalies face/neck (744.89) | <input type="checkbox"/> Lack of nml physiological dev (783.40) |
| <input type="checkbox"/> Other sex chromosome anomaly (758.81) | <input type="checkbox"/> Eye anomaly (743.9) NOS | <input type="checkbox"/> Brain anomalies (742.4) NOS |
| <input type="checkbox"/> Ambiguous genitalia (752.9) | <input type="checkbox"/> Cleft lip/palate (749.20) NOS | <input type="checkbox"/> Failure to thrive (783.41) |
| <input type="checkbox"/> Multiple congenital anomalies (759.7) | <input type="checkbox"/> Hypotonia, cong. (779.89) | <input type="checkbox"/> Mixed develop disorder (315.5) NOS |
| <input type="checkbox"/> Unspec anom congen anomaly (759.9) | <input type="checkbox"/> Obesity (278.00) | <input type="checkbox"/> Other spec cond. perinatal (779.89) |
| <input type="checkbox"/> Other Specified anomalies (759.89) congenital | <input type="checkbox"/> Delay in sexual development (259.0) | <input type="checkbox"/> Fam hx of other cong anomal (v19.5) |
| <input type="checkbox"/> Developmental delay (315.9) NOS | <input type="checkbox"/> Abnormal ears (744.29) | <input type="checkbox"/> Family hx, mental retardation (v18.4) |
| <input type="checkbox"/> Delayed learning (315.2) NOS | <input type="checkbox"/> Short Stature (783.43) | <input type="checkbox"/> Fam hx other other conditions (v19.8) |
| <input type="checkbox"/> MR Moderate (318.0) <input type="checkbox"/> Severe (318.1) <input type="checkbox"/> Profound (318.2) | | <input type="checkbox"/> Congen heart anomaly _____ |
| <input type="checkbox"/> Infertility -> <input type="checkbox"/> female (628.9) NOS / <input type="checkbox"/> male (606.9) NOS | | <input type="checkbox"/> Language Delay (315.31) |
| <input type="checkbox"/> Multiple Miscarriages (habitual abortion) {G ____ P ____ A ____} [SAB ____ TAB ____] | | |
| <input type="checkbox"/> Additional Clinical Information/ICD-9 codes: _____ | | |

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. For 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.)

SUSPECTED DIAGNOSIS OR CHROMOSOME ABNORMALITY (not for billing purposes)

- | | | |
|---|--|--|
| <input type="checkbox"/> Trisomy 21 (Down Syndrome) | <input type="checkbox"/> Habitual Aborter | <input type="checkbox"/> Smith Magenis Synd. (17p-) |
| <input type="checkbox"/> Trisomy 13 | <input type="checkbox"/> Prenatal confirmation | <input type="checkbox"/> Di-George/ VCF Synd. (22q-) |
| <input type="checkbox"/> Trisomy 18 | <input type="checkbox"/> Prader Willi Synd. (15q-) | <input type="checkbox"/> Miller-Dieker Synd. (17p-) |
| <input type="checkbox"/> Turner Syndrome (45,X) | <input type="checkbox"/> Angelman Synd. (15q-) | <input type="checkbox"/> Klinefelter Syn. (47,XXY) |
| <input type="checkbox"/> Mental Retardation (for Fragile X Syndrome – purple top also required) | | |
| <input type="checkbox"/> Other genetic/chromosome abnormality: _____ | | |

TEST REQUESTED	FISH Specific Probes
----------------	----------------------

- | | |
|---|---|
| <input type="checkbox"/> Routine Chromosome / karyotype (7 - 10 days)
(88230, 88262, 88280, 88285)
<input type="checkbox"/> STAT Chromosomes (48 hrs. - additional charges)
(88230 88262, 88280, 88285, 88261)
<input type="checkbox"/> Chromosome / karyotype +Stat FISH (13/18/21/X/Y)
(88230, 88262, 88280, 88285, 88271x5, 88274x5)
<input type="checkbox"/> Routine Chromosome / karyotype +FISH select → →
<input type="checkbox"/> FISH Only - chromosome previously done select → → →
<input type="checkbox"/> Culture Only – Microarray testing | <input type="checkbox"/> Angelman 15q12 <input type="checkbox"/> +13 <input type="checkbox"/> +18 <input type="checkbox"/> +21
<input type="checkbox"/> Prader-Willi 15q12 <input type="checkbox"/> Sex - X&Y
<input type="checkbox"/> DiGeorge/VCF 22q11.2 <input type="checkbox"/> SRY Yp11.3
<input type="checkbox"/> Miller-Dieker 17p13 <input type="checkbox"/> 4p-
<input type="checkbox"/> STS Xp22.3 <input type="checkbox"/> 5p-
<input type="checkbox"/> Kallmann Xp22.3 <input type="checkbox"/> _____
<input type="checkbox"/> Smith-Magenis 17p11 <input type="checkbox"/> Subtelomere
<input type="checkbox"/> Williams 7q11 <input type="checkbox"/> SHOX Xp22.3 |
|---|---|

2012 PERIPHERAL BLOOD REFERRAL FORM FOR CHROMOSOME ANALYSIS
Medical Genetics – Wake Forest University School of Medicine – Winston Salem NC
www.wfubmc.edu/medical genetics Phone: 336-716-4321 Fax: 336-716-2554

Collection Date: _____ **Time:** _____ am/pm **WFU LAB #:** _____

CYTOGENETIC LAB USE ONLY

Name: _____ **Lab #:** _____

Date Received: _____/_____/_____ **Time Received:** _____

Additional Specimen Evaluation: _____

Additional Samples Received: DNA: _____ FRAX FISH other _____

LABORATORY REPORT SUMMARY

Date culture initiated: _____/_____/_____ **Tech:** _____

Sample type: PB FISH FRAX PUBS BM stat

Culture: 48h. 72h. col. 72h.EtBr DBM 24h 96h.

Media: RPMI 1640 less folic acid RPMI 1640

REPORT OF RESULTS / SPECIMEN SUMMARY

Final Preliminary Read Back **Date** _____ **Tech** _____

To: _____

KARYOTYPE: 46,XY 46,XX

INTERPRETATION: normal male normal female

Abnormal: _____

Additional Studies / Results: NOR C-band R-band

FISH: normal male normal female

+13 +18 +21 -X +X +Y

Williams 7q22
 DiGeorge/VCF 22q11
 Prader-Willi 15q12
 Angelman 15q12
 Smith-Magines 17p11
 Miller-Dieker 17p13
 STS Xp22.3
 chromosome paint # _____

To: _____ **By:** _____ **Date:** _____