



# Genetics Laboratory Outpatient - Order Form

Central Scheduling

PH:937-641-4000 Fax: 937:641-4500

One Children's Plaza • Dayton, OH 45404 937-641-5100, FAX 641-5403

PATIENT NAME (LAST, FIRST, MI)				Office Contact Person:				
Address:		Home Phone:		Referring Provider:				
City:	State:	Zip:	Work Phone:	Address:		City:	State:	Zip:
Sex:	Date of Birth:	Medical Record #		Phone Number:		Fax Number:		
Specimen Source:		Specimen Collection:		Provider Signature: _____				
Fax to:		Date:		Date of Signature (required): _____				
Call to:		Time:						
Copy to:		Tech:						
Ethnicity: African American Asian Caucasian Hispanic Other- please specify: _____								
<b>Clinical Description (required):</b>								
<b>ICD10 Code (Required for DCH Billing):</b>								

**NOTICE TO PHYSICIANS:** Medicare, Medicaid and certain commercial insurances do not reimburse for screening or other tests that are not medically necessary to diagnose and treat the patient's current symptoms and conditions. The ordering physician certifies that the following test(s) meet relevant medical necessity criteria or have been identified as screening. Advance Beneficiary Notices (ABNs) must be obtained for non-covered tests. ABN has been obtained and sent to the hospital.

Physician or Authorized Signature: \_\_\_\_\_ Date: \_\_\_\_\_

STAT ASAP

## CYTOGENETICS (Test Code) Phone 937-641-3801 • Fax 937-641-5956

### Routine Chromosome Analysis (CHROB1)

\*\* Requires 1 cc peripheral blood in Na-Heparin tube

### Chromosomal Microarray (MASNPB)

\*\* Requires 3 cc peripheral blood in EDTA tube

Parental Studies CHILD'S NAME: \_\_\_\_\_

### Chromosomal Microarray (PARMCR)

\*\* Requires 3 cc peripheral blood in EDTA tube

### FISH (PARF)

\*\* Requires 2-3 cc peripheral blood in Na-Heparin tube

### Bone Marrow Chromosome Analysis (BMMGB)

\*\* Requires 3 cc bone marrow

### Leukemic Blood Chromosome Analysis (BMMGB)

\*\* Requires 5-7 cc peripheral blood in Na-Heparin tube

### Fluorescence in situ hybridization (Specify below)

\*\* Requires 3 cc peripheral blood in Na-Heparin tube

Rapid Aneuploidy FISH (13, 18, 21, X & Y) with Routine Chromosome

### Analysis (RAFSH1)

Velocardiofacial/DiGeorge syndrome: 22q11.2 (FISHMD)

Smith-Magenis syndrome: 17p11.2 (FISHMD)

Williams syndrome: 7q11.23 (FISHMD)

XY (XYFSHB)

Miller-Dieker syndrome: 17p13.3 (FISHMD)

Special/Oncology FISH analysis (FMISC): \_\_\_\_\_

Call laboratory to arrange prior to order at 937-641-3801

### Chromosome Analysis, Turner syndrome (TURNER)

\*\* Requires 3 cc peripheral blood in Na-Heparin tube

## MOLECULAR GENETICS (Test Code) Phone 937-641-3262 • Fax 937-641-5872

\*\* 3 - 10 cc peripheral blood in EDTA tube (lavendar top)

Fragile X syndrome (FMR1)

DNA Isolation Only (DNAI)

Angelman syndrome (ANGEL)

Factor V Leiden R506Q (5FLE)

Prader-Willi Syndrome (PRW)

Prothrombin G20210A (PRT)

Hereditary Hemochromatosis C282Y/H63D (HHC)

Thrombophilia Panel (THROM)- includes 5FLE, PRT

## Biochemical Genetics (Test Code) Phone 937-641-3262 • Fax 937-641-5872

\*\* 0.5 - 2 cc peripheral blood in Na-Heparin tube

Amino Acid Analysis, plasma (AAPP)

Phenylalanine level, plasma (GPKU)

Phenylalanine + Tyrosine levels, plasma (GPKUB)

**Other:** Please indicate test name and test code or GMISC1 (gene test) or GMISCB (biochemical test) if known. Please indicate specimen type, tube type, and volume if known.

**BILLING:**  
Insurance Billing - Attach demographic and insurance information  
Institutional Billing