

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 Sc	ource: Specim	nen Collection:						
Fax to:		Date:	Provider Signature:					
Call to:		Time:						
Copy to:		Tech:	Date of Signature (required):					
Ethnicity:	African American Asian Caucas	isian Hispanic Other- please specify:						
Clinical Deso	Clinical Description (required):							
ICD10 Code	ICD10 Code (Required for DCH Billing):							
conditions.		ain commercial insurances do not reimburse for screenin llowing test(s) meet relevant medical necessity criteria or						
Physician or	Authorized Signature:		Date:					
STAT	ASAP							
		CYTOGENETICS (Test Code)	Phone 937-641-3801 • Fax 937	7-641-5956				
Routin	ne Chromosome Analysis (CHROB1)		Chromosome	Chromosome Analysis, Turner syndrome (TURNER)				
** Paguiras 1 cs parinharal blood in Na Haparin tuba			** Paguiros 2 cs paripharal bload in Na Haparin tuba					

Routine Chromosome Analysis (CHROB1) ** Requires 1 cc peripheral blood in Na-Heparin tube	Chromosome Analysis, Turner syndrome (TURNER) ** Requires 3 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 Chromoso Analysis (RAFSH1)	me				
Velocardiofacial/DiGeorge syndrome: 22q11.2 (FISHMD) Williams syndrome: 7q11.23 (FISHMD)	Smith-Magenis syndrome: 17p11.2 (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				
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 (SFLE)				
Prader-Willi Syndrome (PRW) Hereditary Hemochromatosis C282Y/H63D (HHC)	Prothrombin G20210A (PRT)				
	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) Phenylalanie + 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 GLOOF_Rev_04/23