

### **GENETICS AND GENOMICS DIAGNOSTIC LABORATORY**

For local courier service and/or inquiries, please contact 513-636-4474 • Fax: 513-636-4373 www.cincinnatichildrens.org/diagnosticlabs • Email: labgeneticcounselors@cchmc.org

**Shipping Address:** 

3333 Burnet Avenue, Room R1042 Cincinnati, OH 45229-3039

Deliveries accepted Monday-Saturday

# PEDIATRIC/ADULT GENETIC TEST REQUISITION

All Information Must Be Completed Before Sample Can Be Processed

PATIENT INFORMATION		SPECIMEN INFORMATION		
Patient Name:,	First MI	□ Bone Marrow □ Saliva □ Cyto *See Page 3 for a list of acceptable		
Home Phone:  MR# Date of Birth //  Gender:   Male  Female			/ Time:	
INDICATIONS/DIAGNOSIS/ICD-10 CODE				
□ ADD/ADHD □ Acute myelogenous leukemia (AML) □ Amenorrhea: 1' or 2' □ Aplastic Anemia □ Autism Spectrum Disorder □ Broad thumbs and/or halluces □ Congenital heart anomaly □ Developmental Delay □ Dysmorphic features □ Encephalopathy □ Eye anomaly □ Erythematous "butterfly" lesion on face	☐ Failure to thrive ☐ Hydrocephalus ☐ Hyper/Hypopigmentation ☐ Hypotonia ☐ Immune deficiency ☐ Intellectual disability ☐ Language disorder ☐ Limb malformation ☐ MRI, abnormal ☐ Macrocephaly ☐ Microcephaly ☐ Myelodysplastic syndrome (MDS)	□ PDD-NOS □ Pancytopenia □ Seizures, convulsions □ Short stature □ Other: □ Newborn Indications: □ Abnormal NIPT/prenatal screen □ Suspected trisomy 21 □ Suspected Turner's syndrome □ Ambiguous genitalia □ Other:	Family History  ☐ Family history of genetic condition:  ☐ Consanguinity (describe relationship):  ☐ Known Chromosome Abnormality:	
BILLING INFORMATION (Choose ONE payment method)		PROVIDER INFORMATION		
REFERRING INSTITUTION Institution: Address: City/State/Zip: Accounts Payable Contact Name: Phone: Fax:		Provider Name (print): Address:) Email:) Genetic Counselor/Lab Contact N Phone: () Email:)		
Email: COMMERCIAL INSURANCE* Insurance can only be billed if requested at the time of service.  Policy Holder Name: Date of Birth / /  Authorization Number:		Referring Physician Signature (RE		
Insurance ID Number: Insurance Name:		ETHNIC/RACIAL BA	ACKGROUND (Choose All)	
Insurance Address: City/State/Zip: Insurance Phone Number: * PLEASE NOTE: • We will not bill Medicaid, Medicaid HMO, CCHMC Patients, CCHMC Providers, or D	or Medicare except for the following: esignated Regional Counties.	☐ European American (White) ☐ Native American or Alaskan ☐ Pacific Islander ☐ Latino-Hispanic ☐ (specify country/region of origin) ☐ Other	☐ African-American (Black) ☐ Asian-American ☐ Ashkenazi Jewish ancestry	
<ul> <li>If you have questions, please call 1-866-450-4198 for complete details.</li> </ul>		(specify country/region of origin)		

#### ☐ Patient signed completed ABN

Medical Necessity Regulations: At the government's request, the Molecular Genetics Laboratories would like to remind all physicians that when ordering tests that will be paid under federal health care programs, including Medicare and Medicaid programs, that these programs will pay only for those tests the relevant program deems to be (1) included as covered services, (2) reasonable, (3) medically necessary for the treatment and diagnosis of the patient, and (4) not for screening purposes.



Proband's DOB:

If testing was **not** performed at CCHMC, please include proband's report

and at least 100ng of proband's DNA to use as a positive control.

Proband's mutation: \_\_\_\_\_

Patient's relation to proband: \_

REQUIRED: Patient Name: Date of Birth:	REQUIRED: Patient Name:	Date of Birth:
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TEST(S) REQUESTED			
Cytogenetic Testing	Molecular Genetic Testing		
Chromosome Analysis	☐ Fragile X DNA testing		
☐ Routine chromosome analysis*	☐ <i>MECP2</i> sequence analysis (Rett syndrome)		
☐ High resolution chromosome analysis*	☐ MECP2 deletion/duplication analysis by MLPA		
☐ Chromosome mosaicism study*	☐ Prader-Willi/Angelman - by methylation-sensitive MLPA		
☐ Reflex to SNP Microarray if chromosome results are normal <sup>†</sup>			
*For chromosome analysis: reflex STAT prelim results on infants <1 month. Reflex to mosaicism study when sex chromosome/mosaic aneuploidy abnormality suspected by laboratory based on indications provided.  *Additional charge for reflex testing. If SNP Microarray is denied by insurance,	<ul> <li>□ PTEN Autism Spectrum Disorder sequencing</li> <li>□ Rubinstein-Taybi and Related Syndromes Gene Panel</li> <li>(CREBBP, EP300, HNRNPH1, HNRNPH2, SIN3A, SIN3B, SRCAP with CREBBP and EP300 deletion/duplication analysis by MLPA)</li> </ul>		
Chromosome Analysis will be performed as the first test in the algorithm.	☐ REFLEX to Whole Exome Sequencing <sup>#</sup> (See additional details below)		
SNP Microarray  SNP Microarray - Constitutional	☐ Spinal Muscular Atrophy - <i>SMN1/SMN2</i> Copy Number Analysis by MLPA☐ Other:		
FISH (Fluorescent In Situ Hybridization)  □ 22q11.2 del (VCFS) (metaphase FISH)	Fanconi Anemia Testing		
□ SRY (Xp11.1q11.1/Yp11.2) (metaphase FISH) □ X/Y centromeres (Xp11.1q1.1/Yp11.1q11.1) (interphase FISH) □ Other FISH (please call lab):	Fanconi anemia (FA) testing will be run sequentially (chromosome breakage study then molecular sequencing if breakage study is normal) unless concurrent testing is selected here:   Concurrent FA testing requested		
U Other Fish (piease cali lab).	☐ Fanconi Anemia (FA) Chromosome Breakage study		
Other Testing  Special study: Cell Culture, storage & freezing Other:	☐ Fanconi Anemia Gene Sequencing Panel (BRCA1, BRCA2, BRIP1, ERCC4, FANCA, FANCB, FANCC, FANCD2, FANCE, FANCF, FANCG, FANCI, FANCL, MAD2L2, PALB2, RAD51, RAD51C, RFWD3, SLX4, UBE2T, XRCC2)		
	☐ FANCA full gene sequencing		
Neurodevelopmental Reflex Genetic Test**	☐ FANCC full gene sequencing		
Tests will be run sequentially based on your selection below:	☐ FANCG full gene sequencing		
$\square$ Patient is macrocephalic: <u>SNP Microarray</u> $\rightarrow$ <u>Fragile X</u> $\rightarrow$ <u>PTEN</u>			
☐ Male patient with normal or small head circumference:	Chromosome Breakage Disorders Testing		
SNP Microarray → Fragile X	☐ Bloom Syndrome - Sister Chromatid Exchange (SCE) analysis		
☐ Female patient with normal or small head circumference:  SNP Microarray → Fragile X → MECP2  **See page 3 for additional information	☐ Chromosome Breakage Disorders Gene Sequencing Panel (ATM, BLM, BRCA1, BRCA2, BRIP1, ERCC4, FANCA, FANCB, FANCC, FANCD2, FANCE, FANCF, FANCG, FANCI, FANCL, LIG4, MAD2L2, MYSM1, NBN, NHEJ1, NSMCE3, PALB2, RAD51, RAD51C, RFWD3, SLX4, UBE2T, XRCC2)		
	d completion of the WES Test Requisition. Also, inclusion of biological parental samples d. Please visit our website at <a href="https://www.cincinnatichildrens.org/exome">www.cincinnatichildrens.org/exome</a> to obtain the required eived by the lab.		
CUSTOM GENE SEQUENCING	DELETION AND DUPLICATION ASSAY		
Gene(s) to be analyzed (specify):	Gene(s) to be analyzed (specify):		
Only genes with clear published functional relationship to rare diseases are accepted.	Please see list of available genes at: www.cincinnatichildrens.org/deldup		
Suspected syndrome/ condition:	Suspected syndrome/ condition:		
Please choose one of the following:	Please choose one of the following:		
☐ Full gene(s) sequencing	$\square$ Deletion and duplication analysis of gene(s) specified above		
☐ Full gene(s) sequencing with reflex to deletion and duplication analysis, if indicated (please see list of genes available for del/dup at	☐ Deletion and duplication analysis of gene(s) specified above with reflex to sequencing, if indicated		
www.cincinnatichildrens.org/deldup)	$\hfill\square$ Analysis of gene(s) specified above from previously analyzed deletion and		
☐ Familial mutation analysis	duplication		
Proband's name:	☐ Familial deletion analysis		

 $\square$  Familial deletion analysis

Proband's mutation: \_\_\_

Proband's DOB: \_\_

Proband's name: \_\_\_\_\_

Patient's relation to proband: \_\_\_

If testing was  $\underline{\textbf{not}}$  performed at CCHMC, please include proband's report

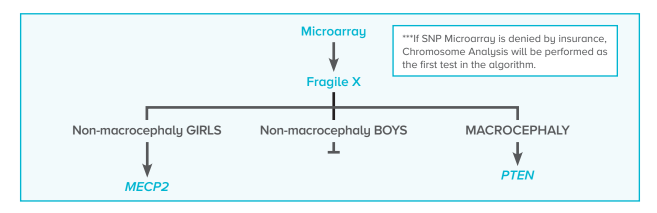
and at least 100ng of proband's DNA to use as a positive control. 2



## **ADDITIONAL INFORMATION**

### **NEURODEVELOPMENTAL REFLEX GENETIC TESTING ALGORITHM**

Tests will be performed sequentially based on the path that your patient follows in the Neurodevelopmental Reflex Test algorithm. Charges will apply to the tests <u>completed</u> in the patient's defined Neurodevelopmental Reflex Test algorithm. Testing will only proceed to the next step if the previous test result is uninformative.



### SPECIMEN REQUIREMENTS

Routine & High Resolution Chromosome Analysis:

3-5 mL blood (NaHep)

SNP Microarray:

3 mL blood (NaHep) and 3 mL blood (EDTA)

FISH Tests: 1-3 mL blood (NaHep)

Neurodevelopmental Reflex Genetic Testing:

3 mL blood (NaHep) and 3-5 mL blood (EDTA)

Fragile X DNA Testing:

3 mL blood (EDTA)

MECP2 del/dup by MLPA, Prader-Willi/Angelman - by MLPA, Spinal Muscular Atrophy - SMN1/SMN2 Copy Number Analysis & Deletion/Duplication Assay:
3 mL blood (EDTA)

Fanconi Anemia Chromosome Breakage Study:

5-10 mL blood (NaHep), 5-10 mL bone marrow (NaHep), or Skin biopsy (3-4 mm tissue in sterile transport media)

FANCA, FANCC, FANCG, MECP2, PTEN & Custom Gene Sequencing:

3 mL blood (EDTA), saliva collection kit $^{*}$ , or 6 cytobrushes

Bloom syndrome - Sister Chromatid Exchange(SCE) analysis:

3-5 mL blood (NaHep)

Chromosome Breakage Disorders Gene Seq, Fanconi Anemia Gene Seq & Rubinstein-Taybi and Related Syndromes Gene Panels:

3 mL blood (EDTA) or saliva collection kit\*

\*Call the office at 513-636-4474 to obtain saliva collection kits