

LABORATORY OF GENETICS AND GENOMICS

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

3333 Burnet Avenue, Room R1042 Cincinnati, OH 45229

HERITABLE LIVER DISEASE TESTING REQUISITION

All Information Must Be Completed Before Sample Can Be Processed

PATIENT INFORMATION	ETHNIC/RACIAL BACKGROUND (Choose All)	
Patient Name:,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	 European American (White) African-American (Black) Native American or Alaskan Asian-American Pacific Islander Ashkenazi Jewish ancestry 	
Home Phone: Date of Birth / / MR# Date of Birth / / Gender:	Latino-Hispanic (specify country/region of origin) Other (specify country/region of origin)	

BILLING INFORMATION (Choose ONE method of payment)

□ REFERRING INSTITUTION

COMMERCIAL INSURANCE*

Institution:	Insurance can only be billed if requested at the time of service.	
	Policy Holder Name:	
Address:	Gender: Date of Birth / /	
City/State/Zip:	Authorization Number:	
Accounts Payable Contact Name:	Insurance ID Number:	
Phone:	Insurance Name:	
Fax:	Insurance Address:	
Email:	City/State/Zip:	
	Insurance Phone Number:	

* PLEASE NOTE:

- We will not bill Medicaid, Medicaid HMO, or Medicare except for the following: Cincinnati Children's Patients, Cincinnati Children's Providers, or Designated Regional Counties.
- If you have questions, please call 1-866-450-4198 for complete details.

SAMPLE/SPECIMEN INFORMATION

SPECIMEN TYPE: \Box Amniotic fluid \Box Blood \Box Cytobrushes

Cord blood	□ CVS	□ Bone marrow	🗆 Saliva
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Tissue (specify): _____

Specimen Date: ____ / ____ / ____ Time: __

Specimen Amount:

Each test requires 3 mL of whole blood in EDTA tube. Please call before sending alternate tissue samples, and for free cytobrush or saliva collection kits.

DRAWN BY:

*Phlebotomist must initial tube of specimen to confirm sample identity

Physician Name (print):		
Address:		
Phone: ()	Fax: ()
Email:		
Genetic Counselor/Lab Contact Name:		
Phone: ()	Fax: ()
Email:		
		Date: / /

\Box 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.



INDICATIONS/DIAGNOSIS/ICD-9 CODE

Reason for Testing:

- □ Diagnosis in symptomatic patient
- □ Carrier testing
- □ Presymptomatic diagnosis of at-risk sibling
- □ Prenatal diagnosis (by previous arrangement only)
- □ Family history of disease
 - Please specify relationship (e.g.; cousin): ____

CLINICAL HISTORY

JaundiceLiver cystsCholestasisKidney cystsFailure to thriveLiver TransplantCharacteristic facial featuresElevated GTPCongenital heart diseaseNormal or low GTPVertebral body defectElevated bilirubinFat malabsorptionPaucity of bile ductsRicketsState State State

□ Other Symptoms (Please specify): ____

TEST(S) REQUESTED

Liver Panels by Next-Generation Sequencing (NGS)

Liver Diseases Panel

(ABCB11, ABCB4, ABCC12, ABCC2, ABCD3, ABCG5, ABCG8, AKR1D1, ALDOB, AMACR, ATP7B, ATP8B1, BAAT, CC2D2A, CFTR, CLDN1, CYP27A1, CYP7A1, CYP7B1, DCDC2, DGUOK, DHCR7, EHHADH, EPHX1, FAH, GPBAR1, HNF1A, HNF1B, HSD17B4, HSD3B7, INVS, JAG1, LIPA, MKS1, MPV17, MYO5B, NEUROG3, NOTCH2*, NPC1, NPC2, NPHP1, NPHP3, NPHP4, NR1H4, PEX1, PEX10, PEX11B, PEX12, PEX13, PEX14, PEX16, PEX19, PEX2, PEX26, PEX3, PEX5, PEX6, PEX7, POLG, SCP2, SERPINA1, SLC10A1, SLC10A2, SLC25A13, SLC27A5, SMPD1, TJP2, TMEM216, TRMU, UGT1A1, VIPAS39, VPS33B excluding exons 1, 2, and 4 in NOTCH2 due to high homologous regions)

□Reflex to deletion/duplication of entire panel

 \Box Reflex to deletion/duplication of single gene(s)⁺ (specify): ____

Note: "Reflex to Liver Diseases Panel" can be ordered with each of the Liver Panels **below.** If the primary test results are negative or they do not fully explain the patients clinical symptoms, the Liver Diseases Panel will automatically be performed when "Reflex to Liver Diseases Panel" is also selected.

□ Jaundice Panel

(ABCB4, ABCB11, ATP8B1, JAG1, TJP2)

 $\hfill\square$ Reflex to deletion/duplication of entire panel

 \Box Reflex to deletion/duplication of single gene(s)⁺ (specify): ____

□ Reflex to Liver Diseases Panel

□ Cystic Diseases of the Liver/Kidney Panel

(ALG8, DNAJB11, DZIP1L, GANAB, LRP5, PKD2, PKHD1, PRKCSH, SEC63)

□Reflex to deletion/duplication of entire panel[†]

□Reflex to deletion/duplication of single gene(s)⁺ (specify): ____

□Reflex to Liver Diseases Panel

Bile Acid Defects Panel

(ABCD3, AKR1D1, AMACR, CYP7B1, HSD3B7)

□ Reflex to deletion/duplication of entire panel

□ Reflex to deletion/duplication of single gene(s)⁺ (specify): ____

□ Reflex to Liver Diseases Panel

□ UGT1A1 Gene Sequencing (Gilbert, Crigler-Najjar Syndromes) □ Reflex to deletion/duplication of UGT1A1

Reflex to Liver Diseases Panel

□ ATP7B Gene Sequencing (Wilson Disease) □ Reflex to deletion/duplication of ATP7B □ Reflex to Liver Diseases Panel

Single Gene Testing

Note: Single gene sequencing is available for all genes in the Liver Panels. Please select a gene from the list below, or use the Custom Gene Sequencing section for any gene that is not specified below.

- □ Alagille syndrome (*JAG1*) full sequence analysis □ Reflex to deletion/duplication of *JAG1*
- □ PFIC1/FIC1 deficiency (ATP8B1) full sequence analysis □ Reflex to deletion/duplication of ATP8B1
- □ PFIC2/BSEP deficiency (*ABCB11*) full sequence analysis □ Reflex to deletion/duplication of *ABCB11*
- □ PFIC3/MDR3 deficiency (ABCB4) full sequence analysis □ Reflex to deletion/duplication of ABCB4
- □ PFIC4/Familial Hypercholanemia (*TJP2*) full sequence analysis □ Reflex to deletion/duplication of *TJP2*
- □ BAAT/Familial Hypercholanemia full sequence analysis □ Reflex to deletion/duplication of BAAT
- α1-antitrypsin deficiency (SERPINA1) full sequence analysis
 Reflex to deletion/duplication of SERPINA1
- □ α1-antitrypsin (SERPINA1) SNP assay for PI*Z and S alleles

□ Targeted (family specific) variant analysis of genes listed above

Gene of interest: _

Proband's name: ___

Proband's DOB: ____

Proband's variant:

Relationship to proband: ____

Please call 513-636-4474 to discuss any family-specific variant analysis with genetic counselor prior to shipment.

If testing was <u>not</u> performed at Cincinnati Children's, please include proband's report and at least 100ng of proband's DNA to use as a positive control.

⁺Deletion/Duplication analysis of *ABCC12* and *GANAB* is not available at this time.

Whole Exome Sequencing

If you are interested in Whole Exome Sequencing, test requisitions are available at: www.cincinnatichildrens.org/exome



CUSTOM GENE SEQUENCING

Gene(s) to be sequenced (specify): _

Only genes with clear published functional relationship to rare diseases are accepted.

Suspected syndrome/ condition: ____

Please choose one of the following:

□ Full gene(s) sequencing

□ Full gene(s) sequencing with reflex to deletion and duplication analysis, if indicated (please see list of genes available for del/dup at www.cincinnatichildrens.org/deldup)

□ Familial variant analysis

Proband's name:

Proband's DOB: ____

Proband's variant: ____

Patient's relation to proband: ____

If testing was <u>not</u> performed at Cincinnati Children's, please include proband's report and at least 100ng of proband's DNA to use as a positive control.

DELETION AND DUPLICATION ASSAY

Gene(s) to be analyzed (specify): ____

Please see list of available genes at: www.cincinnatichildrens.org/deldup

Suspected syndrome/ condition: _

Please choose one of the following:

Deletion and duplication analysis of gene(s) specified above

- Deletion and duplication analysis of gene(s) specified above with reflex to sequencing, if indicated
- □ Analysis of gene(s) specified above from previously analyzed deletion and duplication

□ Familial deletion analysis

Proband's name: ____

Proband's DOB: _____

Proband's variant: ____

Patient's relation to proband: _____

If testing was <u>not</u> performed at Cincinnati Children's, please include proband's report and at least 100ng of proband's DNA to use as a positive control.