

## MOLECULAR GENETICS TEST REQUISITION

All Information Must Be Completed Before Sample Can Be Processed

### PATIENT INFORMATION

Patient Name: \_\_\_\_\_  
Last First MI

Address: \_\_\_\_\_  
\_\_\_\_\_

Home Phone: \_\_\_\_\_

MR# \_\_\_\_\_ Date of Birth \_\_\_\_\_ / \_\_\_\_\_ / \_\_\_\_\_

Gender:  Male  Female

### ETHNIC/RACIAL BACKGROUND (Choose All)

- European American (White)  African-American (Black)  
 Native American or Alaskan  Asian-American  
 Pacific Islander  Ashkenazi Jewish ancestry  
 Latino-Hispanic \_\_\_\_\_  
(specify country/region of origin)  
 Other \_\_\_\_\_  
(specify country/region of origin)

### BILLING INFORMATION (Choose ONE method of payment)

#### REFERRING INSTITUTION

Institution: \_\_\_\_\_

Address: \_\_\_\_\_

City/State/Zip: \_\_\_\_\_

Accounts Payable Contact Name: \_\_\_\_\_

Phone: \_\_\_\_\_

Fax: \_\_\_\_\_

Email: \_\_\_\_\_

#### COMMERCIAL INSURANCE\*

Insurance can only be billed if requested at the time of service.

Policy Holder Name: \_\_\_\_\_

Gender: \_\_\_\_\_ Date of Birth \_\_\_\_\_ / \_\_\_\_\_ / \_\_\_\_\_

Authorization Number: \_\_\_\_\_

Insurance ID Number: \_\_\_\_\_

Insurance Name: \_\_\_\_\_

Insurance Address: \_\_\_\_\_

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

Has patient received a bone marrow transplant?  Yes  No

If yes, date of bone marrow transplant \_\_\_\_\_

Percent engraftment \_\_\_\_\_

Please send saliva kit and two cytobrushes. Note: STR analysis at an additional charge is required on cytobrushes and saliva samples obtained on all patients post BMT.

SPECIMEN TYPE:  Amniotic fluid  Blood  Cytobrushes  Saliva

Cord blood  CVS  Bone marrow  Other \_\_\_\_\_

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

### REFERRING PHYSICIAN

Physician Name (print): \_\_\_\_\_

Address: \_\_\_\_\_

Phone: ( \_\_\_\_\_ ) \_\_\_\_\_ Fax: ( \_\_\_\_\_ ) \_\_\_\_\_

Email: \_\_\_\_\_

Genetic Counselor/Lab Contact Name: \_\_\_\_\_

Phone: ( \_\_\_\_\_ ) \_\_\_\_\_ Fax: ( \_\_\_\_\_ ) \_\_\_\_\_

Email: \_\_\_\_\_

\_\_\_\_\_ Date: \_\_\_\_/\_\_\_\_/\_\_\_\_

Referring Physician Signature (REQUIRED)

Contact Information for Results/Questions (if different than ordering provider) :

Name & Title: \_\_\_\_\_

Phone: ( \_\_\_\_\_ ) \_\_\_\_\_ Fax: ( \_\_\_\_\_ ) \_\_\_\_\_

Email: \_\_\_\_\_

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-10 CODE

Reason for Testing:

- Diagnosis in symptomatic patient
- Asymptomatic infant with abnormal newborn screen
- Carrier (Heterozygote) testing
- Presymptomatic diagnosis of at-risk sibling
- Prenatal testing (by previous arrangement only)
- Family history of disease

### SAMPLE/SPECIMEN INFORMATION

**Has patient received a bone marrow transplant?**

- Yes
- No

If yes, date of bone marrow transplant: \_\_\_\_\_

Percent engraftment: \_\_\_\_\_

### PEDIGREE OR FAMILY HISTORY

Parental Consanguinity  Y  N

### TEST(S) REQUESTED

#### Custom Gene Testing

Sanger sequencing of clinically relevant rare disease genes is available by prior arrangement. Custom single gene requisition available at: [www.cincinnatichildrens.com/customtesting](http://www.cincinnatichildrens.com/customtesting)

#### Deletion/Duplication Assay

Custom deletion/duplication analysis available for many genes. List of available genes and test requisition available at: [www.cincinnatichildrens.com/deldup](http://www.cincinnatichildrens.com/deldup)

#### Whole Exome Sequencing

Exome sequencing test requisition available at: [www.cincinnatichildrens.org/exome](http://www.cincinnatichildrens.org/exome)

#### Epidermolysis Bullosa

- EBSeq Epidermolysis Bullosa Panel  
(*CD151, CDSN, CHST8, COL17A1, COL7A1, DSP, DST, EXPH5, FERMT1 (KIND1), ITGA3, ITGA6, ITGB4, JUP, KLHL24, KRT5, KRT14, LAMA3, LAMB3, LAMC2, PKP1, PLEC1, TGM5*)

#### Fatty Acid Oxidation Disorders

- ACADM* (K329E) genotyping only
- ACADM* (K329E) genotyping, with reflex to full sequencing if indicated
- ACADM* full gene sequence analysis
- ACADM* full gene sequence analysis, with reflex to MetaboSeq<sup>®</sup> if indicated
- ACADVL* full gene sequence analysis
- ACADVL* full gene sequence analysis, with reflex to MetaboSeq<sup>®</sup> if indicated
- CPT2* full gene sequence analysis
- CPT2* full gene sequence analysis, with reflex to MetaboSeq<sup>®</sup> if indicated
- HADHA* (E510Q) genotype analysis
- HADHA* (E510Q) genotype analysis, with reflex to MetaboSeq<sup>®</sup> if indicated
- MetaboSeq<sup>®</sup> (19 gene next-generation sequencing panel)
- SLC22A5* full gene sequence analysis
- SLC22A5* full gene sequence analysis, with reflex to MetaboSeq<sup>®</sup> if indicated

#### Genetic Pharmacology Services\*

- Psychiatry Pharmacogenetics Expanded Panel
- TPMT Genotype Analysis
- Opioid CYP2D6 Pharmacogenetics Panel
- CYP2C19
- CYP2C9
- CYP2D6
- CYP2D6/CYP2C19
- CYP3A5 (if liver transplant, must test DONOR sample)
- Tamoxifen
- Warfarin

\*DONOR sample required for liver transplant patients

#### Hearing Loss Testing

##### Hearing Loss Panels

(Provide audiogram and MRI/CT report)

- Hearing Loss Panel Tier I (*GJB2* sequencing, *GJB6* deletion analysis and 8 mitochondrial mutations)
- OtoSeq<sup>®</sup> Hearing Loss Panel (next-generation sequencing of 23 genes)
- Hearing Loss Panel Tier I with reflex to OtoSeq<sup>®</sup> Hearing Loss Panel, if indicated
- Branchiootorenal Spectrum Disorder (BOR/BOS) Panel (sequencing of *EYA1, SIX1, SIX5*)
- Branchiootorenal Spectrum Disorder (BOR/BOS) Panel with reflex to OtoSeq<sup>®</sup> reanalysis, if indicated
- Hearing loss mtDNA panel (961, 1555, 1494, 3242, 3271, 7445, 7511, 8344)
- Pendred Syndrome Panel (*SLC26A4, FOX11, KCNJ10*)
- Pendred Syndrome Panel with reflex to OtoSeq<sup>®</sup> reanalysis, if indicated
- Usher Syndrome Panel (sequencing of *CDH23, CLRN1, GPR98, MYO7A, PCDH15, USH1C, USH1G, USH2A, WHRN*)
- Usher Syndrome Panel with reflex to OtoSeq<sup>®</sup> reanalysis, if indicated

##### Single Gene Tests

- CDH23* (*USH1D* and *DFNB12*)
- EYA1* (Branchiootorenal spectrum disorder type 1)
- GJB2* (connexin 26)
- GJB6* (connexin 30) deletion analysis
- MYO7A* (*USH1B*, *DFNB2*, *DFNA11*)
- OTOF* (*AUNB1*, *DFNB9*)
- SLC26A4* (Pendred syndrome, *DFNB4*)

**TEST(S) REQUESTED CONTINUED**

**Hematology Testing**

- Dense Deposit Disease/C3 Glomerulonephritis Sequencing Panel**  
(Includes C3, CD46 (MCP), CFB, CFD, CFH, CFHR2, CFHR5 and CFI)

**Hemoglobin Testing**

- Alpha (HBA1/2) and Beta (HBB) Globin Gene Locus Analysis
  - HBA1 and HBA2 (α-globin) sequence analysis
  - HBA1 and HBA2 (α-globin) locus del/dup analysis (HBA1/2 & HBZ)
  - HBB (β-globin) sequence analysis
  - HBB (β-globin) locus del/dup analysis (HBB, HBD, HBG1/2, & HBE)
- Hemoglobin D (HBD) Delta (HBD) Globin Locus Sequence Analysis
- Gamma (HBG1/2) Globin Gene Analysis by Custom Gene Sequencing

**Hemolytic Anemia**

- Hemolytic Anemia Panel (includes sequence analysis of ABCG5, ABCG8, AK1, ALAS2, ALDOA, ANK1, ATP11C, C15orf41, CDAN1, COL4A1, EPB41, EPB42, G6PD, GATA1, GCLC, GPI, GPX1, GSR, GSS, GYPC, HK1, KCNN4, KIF23, KLF1, LPIN2, NT5C3A, PFKM, PGK1, PIEZO1, PKLR, RHAG, SEC23B, SLC2A1 (GLUT1), SLC4A1, SPTA1, SPTB, TPI1, XK)
- Congenital Dyserythropoietic Anemia (CDA) Panel (includes sequence analysis of ALAS2, C15orf41, CDAN1, GATA1, KIF23, KLF1, LPIN2, SEC23B)
- CDA Panel with reflex to Hemolytic Anemia Panel reanalysis, if indicated
- RBC Membrane Disorders Panel (includes sequence analysis of ABCG5, ABCG8, ANK1, ATP11C, COL4A1, EPB41, EPB42, GYPC, KCNN4, PIEZO1, RHAG, SLC2A1, SLC4A1, SPTA1, SPTB, XK)
- RBC Membrane Disorders Panel with reflex to Hemolytic Anemia Panel reanalysis, if indicated
- RBC Enzymopathy Panel (includes sequence analysis of AK1, ALDOA, G6PD, GCLC, GPI, GPX1, GSR, GSS, HK1, NT5C3A, PFKM, PGK1, PKLR, TPI1)
- RBC Enzymopathy Panel with reflex to Hemolytic Anemia Panel reanalysis, if indicated

**Thrombophilia Testing**

- Factor V (Leiden)
- MTHFR (677 C>T and 1298 A>C) genotype
- Prothrombin (Factor II) G20210A genotype
- Thrombophilic polymorphism panel (Factor V- Leiden and Factor II-Prothrombin G20210A)

**Thrombotic Microangiopathy (aHUS and TTP)**

- aHUS Genetic Susceptibility Panel (Includes sequence analysis of ADAMTS13, C3, C4BPA, CD46 (MCP), CD59, CFB, CFH, CFHR1, CFHR2, CFHR3, CFHR4, CFHR5, CFI, DGKE, MMACHC, PLG, THBD and deletion/duplication analysis of CFHR1 and CFHR3 via MLPA. Also includes analysis of variants c.2653C>T and c.2654G>A in the C5 gene, which are associated with poor response to eculizumab)
- ADAMTS13 gene sequencing

**Hereditary Immunodeficiency Testing**

**Autoimmune Lymphoproliferative Syndrome**

- Autoimmune Lymphoproliferative Syndrome (ALPS) Panel by next generation sequencing (NGS)  
(ADA2 (CECR1), CASP10, CASP8, CTLA4, FADD, FAS, FASLG, ITK, KRAS, LRBA, MAGT1, NRAS, PRKCD, RASGRP1, STAT3)
- FAS (TNFRSF6)
- FASLG (TNFSF6)
- CASP10

Somatic FAS sequence analysis of sorted double-negative T cell (DN2C)  
**You MUST call 513-636-2731 in advance for specimen requirements and to schedule this test!**

**Bone Marrow Failure Syndromes Panel by next-generation sequencing (NGS)**

(ABCB7, ACD, ADA2 (CECR1), AK2, AP3B1, ATM, ATR, BLM, BRCA1, BRCA2, BRIP1, CD40LG, CLPB, CSF3R, CTC1, CXCR2, CXCR4, DKC1, DNAJC21, EFL1, EIF2AK3, ELANE, EPO, ERCC4, ERCC6L2, FANCA, FANCB, FANCC, FANCD2, FANCE, FANCF, FANCG, FANCI, FANCL, FANCM, G6PC3, GATA1, GATA2, GF11, HAX1, HYOU1, JAGN1, LAMTOR2, LIG4, LYST, MAD2L2, MPL, MRTFA (MKL1), MYSM1, NAF1, NBN, NHEJ1, NHP2, NOP10, NSMCE3, PALB2, PARN, POT1, RAB27A, RAC2, RAD51, RAD51C, RBM8A, RFWD3, RMRP, RNF168, RPL11, RPL15, RPL18, RPL26, RPL27, RPL31, RPL35, RPL35A, RPL5, RPL9, RPS10, RPS15, RPS15A, RPS17, RPS19, RPS24, RPS26, RPS27, RPS27a, RPS28, RPS29, RPS7, RTEL1, RUNX1, SBDS, SLC37A4, SLX4, SMARCD2 (except exon1), SRP54, SRP72, STK4, STN1, TAZ, TCIRG1, TCN2, TERC, TERF2IP, TERT, TINF2, TP53, TSR2, UBE2T, USB1, VPS13B, VPS45, WAS, WDR1, WIPF1, WRAP53, XRCC2)

**Chromosome Breakage Syndrome Panel**

(ATM, BLM, BRCA1, BRCA2, BRIP1, ERCC4, FANCA, FANCB, FANCC, FANCD2, FANCE, FANCF, FANCG, FANCI, FANCL, FANCM, LIG4, MAD2L2, MYSM1, NBN, NHEJ1, NSMCE3, PALB2, RAD51, RAD51C, RFWD3, SLX4, UBE2T, XRCC2)

**Diamond-Blackfan anemia panel**

(EPO, GATA1, RPL11, RPL15, RPL18, RPL26, RPL27, RPL31, RPL35, RPL35A, RPL5, RPL9, RPS10, RPS15, RPS15A, RPS17, RPS19, RPS24, RPS26, RPS27, RPS27a, RPS28, RPS29, RPS7, TSR2)

**Dyskeratosis congenita Panel**

(ACD, CTC1, DKC1, NAF1, NHP2, NOP10, PARN, POT1, RTEL1, STN1, TERC, TERF2IP, TERT, TINF2, WRAP53)

**Familial hemophagocytic lymphohistiocytosis**

- Hemophagocytic Lymphohistiocytosis (HLH) Panel  
(AP3B1, AP3D1 (except exon 23), CD27, CD70, CTPS1, GATA2, ITK, LYST, MAGT1, NLRC4, PRF1, RAB27A, SH2D1A, SLC7A7, STX11, STXBP2, UNC13D, XIAP)

If inadequate DNA is present, we will prioritize testing according to our FHL testing algorithm, unless you indicate a different order of prioritization below.

- \_\_\_ UNC13D (MUNC13-4)
- \_\_\_ PRF1
- \_\_\_ RAB27A (Griscelli syndrome)
- \_\_\_ STXBP2
- \_\_\_ STX11

**Fanconi anemia**

- Fanconi Anemia Panel by next-generation sequencing (NGS)  
(BRCA1, BRCA2, BRIP1, ERCC4, FANCA, FANCB, FANCC, FANCD2, FANCE, FANCF, FANCG, FANCI, FANCL, FANCM, MAD2L2, PALB2, RAD51, RAD51C, RFWD3, SLX4, UBE2T, XRCC2)
- FANCA

**Lymphoproliferative disorders (Including EBV-Related)**

- SH2D1A
- XIAP (BIRC4)
- ITK
- MAGT1

## TEST(S) REQUESTED CONTINUED

### Severe Combined Immunodeficiencies

- Severe Combined Immunodeficiency Panel by next-generation sequencing (NGS)  
*(ADA, AK2, ATM, BCL11B, CD247, CD3D, CD3E, CDH17, CHD7, CIITA, CORO1A, DCLRE1C, DOCK8, FOXP1, IL2RG, IL7R, JAK3, LAT, LCK, LIG4, MSN, NHEJ1, ORAI1, PNP, PRKDC, PTPRC, RAC2, RAG1, RAG2, RFX5, RFXANK, RFXAP, RMRP, STAT5B, STIM1, STK4, TAP1, TAP2, TBX1, TTC7A, ZAP70)*
- X-linked severe combined immunodeficiency (*IL2RG*)

### Severe congenital neutropenia

- Inherited Neutropenia Panel by next-generation sequencing (NGS)  
*(AK2, AP3B1, CD40LG, CLPB, CSF3R, CXCR2, CXCR4, DNAJC21, EFL1, EIF2AK3, ELANE, G6PC3, GATA1, GATA2, GFI1, HAX1, HYOU1, JAGN1, LAMTOR2, LYST, MRTFA (MKL1), RAB27A, RAC2, RMRP, RUNX1, SBDS, SLC37A4, SMARCD2 (except exon1), SRP54, STK4, TAZ, TCIRG1, TCN2, TP53, USB1, VPS13B, VPS45, WAS, WDR1, WIPF1)*
- ELANE (ELA2)*
- HAX1*
- WAS (males only)*

### Other Primary Immunodeficiencies

- IPEX syndrome (*FOXP3*)
- Shwachman Diamond syndrome (*SBDS*)
- Wiskott-Aldrich syndrome (*WAS*)
- X-linked hyper IgM immunodeficiency (*CD40LG*)

### Hereditary Liver Disease Testing

- Liver Diseases Panel (*ABCB11, ABCB4, 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*)
- Jaundice Panel (*ABCB4, ABCB11, ATP8B1, JAG1, TJP2*)
- Cystic Diseases of the Liver/Kidney Panel (*ALG8, DNAJB11, DZIP1L, GANAB, LRP5, PKD2, PKHD1, PRKCSH, SEC63*)
- Bile Acid Defects Panel (*ABCD3, AKR1D1, AMACR, CYP7B1, HSD3B7*)

### Heritable Pancreatic Disease Testing

- Pancreas Panel (*CASR, CEL, CFTR, CLDN2, CPA1, CTRC, PRSS1, SBDS, SPINK1, UBR1*)
- Pancreatic Insufficiency Panel (*CEL, CFTR, SBDS, UBR1*)
- SBDS* full sequence analysis

### Single Gene Tests

- ABCB4* (PFIC3/MDR3 deficiency)
- ABCB11* (PFIC2/BSEP deficiency)
- ATP7B* Gene Sequencing (Wilson Disease)
- ATP8B1* (PFIC1/FIC1 deficiency)
- BAAT* (Familial Hypercholanemia)
- JAG1* (Alagille syndrome)
- SERPINA1* genotyping assay for PI\*Z and PI\*S alleles
- SERPINA1* (alpha-1-antitrypsin deficiency)
- TJP2* (PFIC4/Familial Hypercholanemia)
- UGT1A1* Gene Sequencing (Gilbert, Crigler-Najjar Syndromes)

### Intellectual Disabilities Testing

#### Creatine deficiency syndromes

- SLC6A8* (creatine transporter defect)
- GATM* (AGAT deficiency)
- GAMT*
- Fragile X syndrome (*FMR1* – By PCR and mPCR)
- MECP2* sequencing (Rett syndrome)
- MECP2* deletion/duplication analysis by MLPA
- Prader-Willi/Angelman syndrome (by MLPA)

#### Lysosomal Storage Disease Testing

##### Cystinosis

- CTNS* full gene sequencing only
- CTNS* 57 kb deletion analysis by MLPA only
- CTNS* 57 kb deletion analysis with reflex to full gene sequencing

##### Fabry Disease

- GLA* full gene sequencing

##### Gaucher Disease

- GBA* full gene sequencing
- Ashkenazi Panel (TESTS ONLY: N370S, L444P, 84GG, IVS2+1G>A)

##### MPS II – Hunter syndrome

- IDS* full gene sequencing

##### Pompe Disease

- GAA* full gene sequencing for Infantile Pompe (by prior arrangement only)
- GAA* full gene sequencing

### Mitochondrial Disorders

#### POLG-related disorders: AD-PEO, SANDO, MIRAS

- POLG* full gene sequencing

#### IBMPFD

- VCP* full gene sequencing

#### Oncology Molecular Testing *Samples must be received within 24 hours of drawing.*

- BCR/ABL t(9;22) Qualitative
- BCR/ABL t(9;22) Quantitative (p210)
- BCR/ABL t(9;22) Quantitative (p190)
- PML-RARα t(15;17)—(qualitative only)

#### Oncology Molecular Testing *Samples must be received within 48 hours of drawing.*

- JAK2* Quantitative (V617F)
- MAP2K1* full gene sequencing analysis (% Tumor: \_\_\_\_\_)
- PTEN* sequencing

#### Targeted (family specific) mutation analysis

- Gene of interest: \_\_\_\_\_
- Proband's name: \_\_\_\_\_
- Proband's DOB (mm/dd/yyyy): \_\_\_\_\_
- Proband's variant: \_\_\_\_\_
- Relationship to proband: \_\_\_\_\_

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

- Bone marrow engraftment by STR - same sex donor and recipient Please specify:  Donor  Host (pre-transplant)  Host (post-transplant)
- Sorted cell engraftment study (WBC sub-population)  
You must call 513-636-4474 to schedule this test prior to sample submission.  
Please specify:  STR  FISH
- Hereditary hemochromatosis (HFE- C282Y and H63D)
- DNA/RNA processing and storage (Call (513) 636-4474 to arrange)
- Maternal Cell Contamination (by STR; required for all prenatal diagnostic testing)
- Maternal Engraftment
- Twin zygosity (by STR)