



# Knight Diagnostic Laboratories

Fax: (855) 535-1329  
 Email: KDLClientServices@ohsu.edu  
 Shipping: 2525 SW 3rd Ave, Ste 350, Portland, OR 97201  
 Questions? (855) 535-1522

## Molecular Genetics Test Requisition

### Patient Information

Patient Last Name

Patient First Name

Street Address

City, State, Zip

Phone  DOB  /  /

Fax  Male  Female

ID/MRN #

Hospital In-Patient Yes  No

### Ordering Healthcare Provider Information

Full Name

NPI

Office/Facility Name

Address

City, State, Zip

Phone  Fax

Account #

Notes

Physician Signature \_\_\_\_\_ Date \_\_\_\_\_

### Send additional copies of test results to:

Healthcare Provider Name

Healthcare Provider Name

Provider Phone  Fax

Provider Phone  Fax

### Billing Information - Select One Billing Method

Self Pay

Bill Insurance

Attach Copy of Insurance Card or Billing Face Sheet

Bill Referring Provider or Institution

Invoice will be sent to Client Account and Address Listed Above

Primary Insurance Name

Primary Policy #

Primary Group #

Preauthorization #

Secondary Insurance Name

Secondary Policy #

Secondary Group #

Preauthorization #

Relation to Insured  Self  Child  Spouse  Other \_\_\_\_\_

Medicaid  Medicare

Relation to Insured  Self  Child  Spouse  Other \_\_\_\_\_

Medicaid  Medicare

### Clinical Information

**Specimen Type**

Amniocytes, Cultured  Date of Specimen Collection

Amniotic Fluid, Direct  Time of Specimen Collection

Blood Spots  ICD-10 (required)

DNA from\* \_\_\_\_\_

CVS, Direct  African American

CVS, Cultured  Alaska Native

Whole Blood  Asian

Saliva

Tissue \_\_\_\_\_

Other \_\_\_\_\_

### Family History / Pedigree

### Diagnosis

Caucasian/NonHispanic  Jewish (Other)

Hispanic American  Native American Indian

Jewish, Ashkenazi  Other: \_\_\_\_\_

Indications for Testing	Known Familial Mutations
Family History, Mutation Known: Yes* No *If Yes, please complete KNOWN FAMILIAL MUTATIONS	Please call Client Services at 1-855-535-1522 and provide clinical report if proband testing was performed outside of OHSU.
	Patient Status: Symptomatic Asymptomatic
Symptomatic	Name of Gene:
Possible Diagnosis	Variant(s) to be tested:
Definite Diagnosis	Name of Proband:
Carrier Testing	Relationship to Proband:
Prenatal Testing	OHSU Sample # of Proband:
Other (Please Specify)	*If proband testing was performed outside of OHSU, please provide clinical report.

### Molecular Diagnostic Tests - Next Generation Sequencing Panels

Code	Test Name	Code	Test Name
2264	Autosomal Dominant Osteogenesis Imperfecta Panel	1690	Inherited Cancer Panel
1085	Bone Marrow Failure Panel	1692	Inherited Colon Cancer Panel
6000	BRCA1 and BRCA2	1693	Inherited Ovarian Cancer Panel
1275	Cholestasis Panel	1694	Inherited Pancreatic Cancer Panel
1277	Ciliopathies Panel	1695	Inherited Prostate Cancer Panel
1165	Connective Tissue Disorders Panel	1895	Maturity-Onset Diabetes of the Young Panel
1245	Craniosynostosis Panel	1135	Migraine and Strokes Panel
1697	Disorders of Sex Development Panel	2142	Monogenic Hypertension Panel
1291	Epilepsy-Seizures Panel	2101	NBIA sequencing Panel
1460	Fanconi Anemia Panel	2135	Noonan and Other Related Disorders Panel
1495	Hearing Loss Panel	2240	Parkinson's Disease Panel
1610	Heterotaxia Panel	2250	Platelet Disorders Panel
1625	Holoprosencephaly Panel	2405	Rett-Angelman Syndrome Panel
1645	Hypercholesterolemia Panel	2590	SOD and Schizencephaly Panel
1691	Inherited Breast Cancer Panel	2810	Wilson's Disease
1696	Inherited Breast/GYN Cancer Panel		

**Molecular Diagnostic Tests - Single Gene and Targeted Testing**

<b>Code</b>	<b>Test Name</b>	<b>Code</b>	<b>Test Name</b>
<b>Angelman Syndrome/Prader-Willi</b>		1420	Factor V Leiden, R506Q Mutation
1020	SNRPN Methylation and Del/Dup	<b>Neurodegeneration with Brain Iron Accumulation (NBIA)</b>	
1050	APOL1 Sequencing (exon 6)	2101	NBIA Panel
1160	CPT1A Targeted Mutation, c.1346C-> T(p.P479L)	1145	C19orf12 (MPAN), Sequencing
1150	Connexin 26, GJB2, Sequencing and Connexin 30, GJB6, Deletion	1400	FA2H (FAHN) Sequencing
<b>Cystic Fibrosis, CFTR</b>		1550	FTL (Neurferitinopathy)
1220	CFTR Screening Mutation Panel	2231	PANK2 (PKAN)
1224	CFTR Sequencing and Del/Dup	1682	PLA2G6 (INAD)
1222	CFTR Sequencing	1080	WDR42 (BPAN)
1226	CFTR Del/Dup	2290	Prothrombin-Related Mutation, G20210A
1280	Duchenne/Becker Muscular Dystrophy Del/Dup	<b>Rett Syndrome (RTT)</b>	
1480	Fragile X Syndrome, FMR1, FMR1-related Primary Ovarian Insufficiency (POI), Fragile X Tremor Ataxia Syndrome (FXTAS)	2402	MECP2 Sequencing and Del/Dup
1600	Hereditary Hemochromatosis (HH), HFE, Common Variants	2400	MECP2 Sequencing
1620	Huntington Disease, HTT repeat expansion Disease-Specific Healthcare Provider Statement Required	2403	MECP2 Del/Dup
<b>Lynch Syndrome (HNPCC)</b>			
2027	MLH1 Promoter Hypermethylation		
5020	MEN2, RET, Sequencing		
2050	Myotonic Dystrophy, DMPK Repeat Expansion		

**Other Laboratory Services**

Code	Test Name	Code	Test Name
1300	DNA Banking Services	2900	Zygosity Testing
1980	Maternal Cell Rule Out	1230	Known Mutation/Familial Variant Targeted Mutation Variant(s) Details
1240	Full Gene Analysis Please Specify Gene(s):		

**NOTICE REGARDING MOLECULAR GENETIC TESTING**

Maternal cell rule-out testing will be performed on all prenatal specimens received. Please provide maternal blood or saliva in addition to the fetal specimen sent for genetic testing. Additional charges apply for the maternal cell rule-out test.

**Result Release**

**Results will be immediately available to the patient unless you mark the box below.**

Do not release (I reasonably believe that an Information Blocking exception applies)

Comments / Requests: