

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		Ordering Physician Information
Patient Last Name		Full Name
Patient First Name		
Street Address		NPI OFF of 1/5 o
City, State, Zip		Office/Facility Name
Phone	DOB / /	Address
Fax	Male Fem	City, State, Zip
ID/MRN#		Phone Fax
Hospital In-Patient	Yes No	Account # Notes
Physician Signature	Date	
Send additional copic	es of test results to:	
Physician Name		Physician Phone Fax
Physician Name		Physician Phone Fax
Billing Information - 9	Select One Billing Method	Billing is done in accordance with the information provided below and OHSU policy. Appropriate areas must be completed or referring laboratory/physician will be billed.
	Bill Insurance Attach Copy of Insurance Card or Billing Face She	Bill Referring Provider or Institution
Primary Insurance Name		Secondary Insurance Name
Primary Policy #		Secondary Policy #
Primary Group #		Secondary Group #
Preauthorization #		Preauthorization #
Relation to Insured Self Child	Medicaid Medicare Spouse Other	Relation to Insured Medicaid Medicare Self Child Spouse Other
Sell Cillu	Spouse Other	Self Child Spouse Other
Clinical Information		
Amniocytes, Cultured Whole Blood Amniotic Fluid, Di Blood Spots DNA from* CVS, Direct CVS, Cultured Whole Blood		Family History / Pedigree Diagnosis
Saliva	African American	Caucasian/ NonHispanic Jewish (Other)
Tissue Other	———— Alaska Native ———— Asian	Hispanic American Native American Indian Jewish, Ashkenazi Other:
	Waldii	Jewish, Ashkenazi Other:

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^{*}DNA extraction must occur in a CLIA-certified lab or a lab meeting equivalent requirements.

Indication 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.
Symptomatic Possible Diagnosis Definite Diagnosis	Patient Status: Symptomatic Asymptomatic Name of Gene:
□ Carrier Testing □ Presymptomatic Testing □ Prenatal Testing □ Predispositional Testing □ Other (Please Specify) Pregnancy: LMP: GMP:	Variants to be tested:

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	1692	Inherited Colon Cancer Panel
6000	BRCA1 and BRCA2	1693	Inherited Ovarian Cancer Panel
1275	Cholestasis	1694	Inherited Pancreatic Cancer Panel
1277	Ciliopathies	1695	Inherited Prostate Cancer Panel
1165	Connective Tissue Disorders	4870	Lynch Syndrome
1245	Craniosynostosis	1895	Maturity-Onset Diabetes of the Young
1697	Disorders of Sex Development Panel	1135	Migraine and Strokes
1291	Epilepsy-Seizures	2142	Monogenic Hypertension Panel
1460	Fanconi Anemia	2101	NBIA Sequencing
1495	Hearing Loss	2135	Noonan and Other Related Disorders
1610	Heterotaxia	2240	Parkinson's Disease
1625	Holoprosencephaly	2250	Platelet Disorders
1645	Hypercholesterolemia	2405	Rett-Angelman Syndrome
1691	Inherited Breast Cancer Panel	2590	SOD and Schizencephaly
1696	Inherited Breast/Gyn Cancer panel	2810	Wilson's Disease

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Molecular Diagnostic Tests – Single Gene and Targeted Testing

Code	Test Name	Code	Test Name
Angelman Syn	drome/Prader-Willi	1420	Factor V Leiden, R506Q mutation
1020	SNRPN Methylation and Del/Dup	Hereditary He	mochromatosis (HH), HFE Mutation
1050 APOL	l Sequencing (exon 6)	1600	C282Y (heterozygous results are reflexed to H63D)
1160 CPT1A	Targeted Mutation, c.1436C → T (p.P479L)	1602	H63D
1150 Connexin 26, GJB2, Sequencing and Connexin 30, GJB6, Deletion			ngton Disease, HTT repeat expansion se-Specific Physician Statment Required
Cystic Fibrosis	, CFTR	Neurodegener	ation with Brain Iron Accumulation (NBIA)
1220	CFTR Screening Mutation Panel (60 mutations)	NBIA NextGer	n Panel
1224	CFTR Sequencing and Del/Dup	2101	Sequencing Only
1222	CFTR Sequencing	PANK2 (PKAN)
1226	CFTR Del/Dup	2230	Sequencing and Del/Dup
1280 Duche	enne/Becker Muscular Dystrophy Del/Dup	2231	Sequencing Only
FMR1-related disorders (including Fragile X), FMR1		2232	Del/Dup Only
	Fragile X Syndrome, FMR1, FMR1-related Primary	PLA2G6 (INAD)	
1480	1480 Ovarian Insufficiency (POI), Fragile X Tremor Ataxia Syndrome (FXTAS)		Sequencing and Del/Dup
Hereditary Hemochromatosis (HH), HFE Mutation		1682	Sequencing Only
1600	C282Y (heterozygous results are reflexed to H63D)	1681	Del/Dup Only
1602	H63D	1145 C19or	rf12 (MPAN), Sequencing
	ngton Disease, HTT repeat expansion	1400 FA2H	(FAHN), Sequencing
	se-Specific Physician Statement Required	1550 FTL, Sequencing	
Lynch Syndrom		1080 WDR45 (Beta-Propeller), Sequencing	
2027	MLH1 Promoter Hypermethylation	2290 Prothr	rombin-Related Mutation, G20210A
4870	Sequencing Panel (MLH1, MSH2, and MSH6)	Rett Syndrome	(RTT)
5020 MEN2,	RET, Sequencing	2402	MeCP2 Sequencing and Del/Dup
2050 Myoto	nic Dystrophy, DMPK Repeat Expansion	2400	MeCP2 Sequencing

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Other Lab	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) Detail:	
1240	Full Gene(s) Analysis Please Specify Gene(s):		Variant(s) Detail:	

NOTICE REGARDING MOLECULAR GENETIC TESTING

Maternal cell rule-out testing will be performed on all prenatal specimens received. Please provide maternal blood 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:	

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