

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 | | Ordering Healthcare Provider Information |
|--|--|---|
| Patient Last Name | | Full Name |
| Patient First Name | | NPI |
| Street Address | | Office/Facility Name |
| City, State, Zip | | Address |
| Phone | DOB / / | City, State, Zip |
| Fax | Male Female | Phone Fax |
| ID/MRN # | | Account # |
| Hospital In-Patient Ye | es No | Notes |
| Physician Signature | Date | |
| Send additional copies of te | est results to: | |
| Healthcare Provider Name |) | Provider Phone Fax |
| Healthcare Provider Name | | Provider Phone Fax |
| Billing Information - 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. |
| Self Pay Bill Insur Attach Co | ance ppy 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 | | Secondary Insurance Name |
| Primary Policy # | | Secondary Policy # |
| Primary Group # | | Secondary Group # |
| Preauthorization # | | Preauthorization # |
| Relation to Insured | Medicaid Medicare | Relation to Insured Medicaid Medicare |
| Self Child Spous | se Other | Self Child Spouse Other |
| Clinical Information Amniocytes, Cultured | Date of Specimen Collection | Family History / Pedigree |
| 2 Amniotic Fluid, Direct | | |
| Blood Spots | Time of Specimen Collection | |
| | | Diagnosis |
| DNA from* CVS, Direct | ICD-10 (required) | Diagnosis |
| CVS, Cultured | | |
| Whole Blood | | |
| Saliva | African American | Caucasian/NonHispanic Jewish (Other) |
| Tissue | Alaska Native | Hispanic American Native American Indian |
| Other | ——— 🗌 Asian | Jewish, Ashkenazi Other: |

| ndications 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 Specifiy) | *If proband testing was performed outside of OHSU, please provide clini- cal 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

| Code | Test Name | Code | Test Name |
|---------------|---|-------------|---|
| Angelman S | Angelman Syndrome/Prader-Willi | | Factor V Leiden, R506Q Mutation |
| 1020 | SNRPN Methylation and Del/Dup | Neurodegene | eration with Brain Iron Accumulation (NBIA) |
| 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 |
| Cystic Fibros | sis, CFTR | 1550 | FTL (Neurferritinopathy) |
| 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 | | |
| 1480 | Fragile X Syndrome, FMR1, FMR1-related Primary Ovarian Insufficincy (POI), Fragile X Tremor Ataxia Syndrome (FXTAS) | Rett Syndro | ome (RTT) |
| 1600 | Hereditary Hemochromatosis (HH), HFE, Common Variants | 2402 | MECP2 Sequencing and Del/Dup |
| 1620 | Huntington Disease, HTT repeat expansion Disease-Specific Healthcare Provider Statement Required | 2400 | MECP2 Sequencing |
| Lynch Syndr | ome (HNPCC) | 2403 | MECP2 Del/Dup |
| 2027 | MLH1 Promoter Hypermethylation | | |
| 5020 | MEN2, RET, Sequencing | | |
| 2050 | Myotonic Dystrophy, DMPK Repeat Expansion | | |

| ther Labo | ner 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): | | | |
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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: