



Instructions: The accurate interpretation and reporting of genetic results is contingent upon the reason for testing and clinical information. To help provide the best possible service, supply the information requested below and send this paperwork with the specimen or return by fax to Mayo Clinic Laboratories, Attn: Genetic Counselors at 507-284-1759. Phone: 800-533-1710 / International clients: +1-507-266-5700 or email MLINT@mayo.edu

Patient Information (required)

Form with fields for Patient Name (Last, First, Middle), Birth Date (mm-dd-yyyy), Sex Assigned at Birth (Male, Female, Unknown, Choose not to disclose), and Legal/Administrative Sex (Male, Female, Nonbinary).

Referring Provider Information

Form with fields for Requesting Provider Name (Last, First), Phone, Fax*, and Genetic Counselor Name (Last, First), Phone, Fax*.

*Fax number given must be from a fax machine that complies with applicable HIPAA regulations.

Reason for Testing

Form with checkboxes for Abnormal ultrasound, Abnormal testing, Family history, and Other, each followed by a blank line for details.

Clinical Information

Large form section containing Donor Egg, Fetal Specimen Source, Fetal Sex, Multiple Gestation Pregnancy, Fetal Sample Collection Date, Maternal Blood Collection Date, and Previous Testing sections.

** See Lab Test Catalog for available cytogenetic tests and ordering guidance.

Molecular Genetics: Prenatal Patient Information (continued)

Test Selection

Maternal Sample

- CYPZ / 21-Hydroxylase Gene, CYP21A2, Full Gene Analysis, Varies (sent as positive control)
 - FMTT / Familial Mutation, Targeted Testing, Varies (sent as positive control)
 - MATCC / Maternal Cell Contamination, Molecular Analysis, Varies[§]
 - UNIPD / Uniparental Disomy, Varies[§]
 - Other:
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-

Fetal Sample

- MATCC / Maternal Cell Contamination, Molecular Analysis, Varies[§]
 - BWRS / Beckwith-Wiedemann Syndrome/Russell-Silver Syndrome, Molecular Analysis, Varies
 - CHDGG / Congenital Heart Disease Gene Panel, Varies[†]
 - CKDGP / Cystic Kidney Disease Gene Panel, Varies[†]
 - CYPZ / 21-Hydroxylase Gene, CYP21A2, Full Gene Analysis, Varies[¶]
 - DBMD / Duchenne/Becker Muscular Dystrophy, DMD Gene, Large Deletion/Duplication Analysis, Varies
 - F81P / Hemophilia A F8 Gene, Intron 1 Inversion Known Mutation Analysis, Prenatal[†]
 - F822P / Hemophilia A F8 Gene, Intron 22 Inversion Mutation Analysis, Prenatal[†]
 - FMTT / Familial Mutation, Targeted Testing Varies^{§§}
 - FXS / Fragile X Syndrome, Molecular Analysis, Varies
 - NSRGG / Noonan Syndrome and Related Conditions Gene Panel, Varies[†]
 - OIBFG / Osteogenesis Imperfecta and Bone Fragility Gene Panel, Varies[†]
 - PWAS / Prader-Willi/Angelman Syndrome, Molecular Analysis, Varies
 - SMNDX / Spinal Muscular Atrophy Diagnostic Assay, Deletion/Duplication Analysis, Varies
 - UNIPD / Uniparental Disomy, Varies[§] (chromosome[s] to be tested:)
 - Other:
-
-

Paternal Sample (if applicable)

- Paternal sample unavailable for testing
- CYPZ / 21-Hydroxylase Gene, CYP21A2, Full Gene Analysis, Varies (sent as positive control)
- FMTT / Familial Mutation, Targeted Testing, Varies (sent as positive control)
- UNIPD / Uniparental Disomy, Varies[§]

Father's Name (*Last, First, Middle*): _____

Father's Birth Date (*mm-dd-yyyy*): _____

[§] If ordering MATCC or UNIPD, an order for MATCC or UNIPD must be placed on both the maternal and the fetal sample. Fetal and maternal samples must be sent under separate order numbers.

[¶] The CYP21A2 Gene Testing for Congenital Adrenal Hyperplasia Patient Information Form (T663) is required for prenatal CYPZ orders; CYPZ testing will not be performed unless this form is also completed and sent with the sample.

^{§§} Also complete Familial Mutations section below.

[†] Also complete the test-specific patient information sheet located in the Lab Test Catalog.

Molecular Genetics: Prenatal Patient Information (continued)

Familial Mutation Testing (required patient information)

Familial Mutations

FMTT / Familial Mutation, Targeted Testing, Varies cannot be performed without the information below.‡

Is the familial mutation a nucleotide substitution or small insertion/deletion of nucleotides? Yes No

If "Yes," provide the familial mutations here:

Mutation 1: Gene _____ Exon/Intron _____ Nucleotide _____ Amino Acid _____

Mutation 2: Gene _____ Exon/Intron _____ Nucleotide _____ Amino Acid _____

Mutation 3: Gene _____ Exon/Intron _____ Nucleotide _____ Amino Acid _____

Is the familial mutation a large deletion or duplication involving one or more exons? Yes No

If "Yes," provide the familial deletion/duplication here:

Deletion Duplication

Gene: _____ Exons: _____

Familial History

Include the name(s) and birth date(s) of the family member(s) who have had genetic testing (ie, proband):

Indicate the family member's relationship to the patient: _____

Important: Attach a copy of the proband's genetic test result and a detailed pedigree, if available.

‡Note: Analysis of regions surrounding the familial variant may be required and may result in the identification of additional sequence variants.