



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

Patient Information

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 Healthcare Professional Information

Form with fields for Referring Healthcare Professional Name (Last, First), Phone, and Fax\* for both the professional and Genetic Counselor/Other Healthcare Professional.

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

Is this a postmortem specimen? Yes No If "Yes," attach autopsy report if available.

Reason for Testing Check all that apply.

Reason for testing options: Diagnosis, Family history\*\*, Sudden death. Includes a note: \*\*Genetic testing should be performed on an affected family member first, when possible. FMTT / Familial Variant, Targeted Testing should be ordered when there is a previous positive genetic test result in the family.

Clinical History

Diagnosis/Suspected Diagnosis options: Marfan Syndrome, Ehlers-Danlos Syndrome, Other, Loeys-Dietz Syndrome, Familial thoracic aortic aneurysm and dissection, Osteogenesis Imperfecta, Cerebrovascular disease/stroke.

Indicate whether the following are present: Aortic diameter at sinuses of Valsalva Z-score ≥ 2, Aortic dissection, Ectopia lentis, Systemic score ≥ 7 points, Aortic dilatation/aneurysm (Z-score < 2), Family history of independently diagnosed Marfan syndrome using the revised Ghent criteria, Talipes equinovarus, Hypertelorism, Craniosynostosis, Cleft palate, Bifid uvula, Blue sclerae, Arterial tortuosity, Patent ductus arteriosus, Velvety/translucent skin, Easy bruising, Widened atrophic scars, Spontaneous organ rupture, Aortic Dimensions, Fractures, Hearing loss, Stroke, Other aneurysm, Other.

Table for Ghent Systemic Score Calculation for Marfan Syndrome with columns for Feature, Value, and Enter Value if Present. Features include Wrist and thumb sign, Wrist or thumb sign, Pectus carinatum, Pectus excavatum or chest asymmetry, Hindfoot deformity, Plain flat foot (pes planus), Pneumothorax, Dural ectasia, Protrusio acetabulae, Reduced upper/lower segment and increased armspan/height, Scoliosis or thoracolumbar kyphosis, Reduced elbow extension, 3 of 5 facial features (dolichocephaly, enophthalmos, downslanting palpebral fissures, malar hypoplasia, retrognathia), Skin striae, Myopia > 3 diopters, Mitral valve prolapse, and a Total row.

List any additional features present:

# Connective Tissue/Cerebrovascular Disease Genetic Testing Patient Information (continued)

Patient Name (Last, First Middle)	Birth Date (mm-dd-yyyy)
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## Family History

Are there similarly affected relatives? <input type="checkbox"/> Yes <input type="checkbox"/> No If "Yes," indicate relationship and symptoms: _____
Have any family member had genetic testing? <input type="checkbox"/> Yes*** <input type="checkbox"/> No <input type="checkbox"/> Unknown <b>***FMTT / Familial Variant, Targeted Testing should be ordered when there is a previous positive genetic test result in the family. Contact the lab for ordering assistance.</b>
History of consanguinity: <input type="checkbox"/> No <input type="checkbox"/> Yes; relationship details: _____

## Ancestry

<input type="checkbox"/> African/African American	<input type="checkbox"/> East Asian	<input type="checkbox"/> Latinx/Latine	<input type="checkbox"/> South Asian	<input type="checkbox"/> Unknown
<input type="checkbox"/> Ashkenazi Jewish	<input type="checkbox"/> European	<input type="checkbox"/> Middle Eastern	<input type="checkbox"/> None of the above	<input type="checkbox"/> Choose not to disclose

**New York State Patients: Informed Consent for Genetic Testing is required.** See Informed Consent for Genetic Testing (T576)  
or Informed Consent for Genetic Testing – Spanish (T826).