



**Instructions:** The accurate interpretation and reporting of the genetic results is contingent upon the reason for referral, clinical information, ethnic background, 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: Personalized Genomics Laboratory Genetic Counselors at 507-284-1759. Phone: 507-266-5700 / International clients: +1-507-266-5700 or email [mclglobal@mayo.edu](mailto:mclglobal@mayo.edu)**

**Patient Information**

Patient Name <i>(Last, First, Middle)</i>	Birth Date <i>(mm-dd-yyyy)</i>	Sex <input type="checkbox"/> Male <input type="checkbox"/> Female
Referring Provider Name <i>(Last, First)</i>	Phone	Fax*
Other Contact Name <i>(Last, First)</i>	Phone	Fax*

\*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.

**Clinical History**

Reason for Testing (Check all that apply.) <input type="checkbox"/> Diagnosis <input type="checkbox"/> Carrier testing <input type="checkbox"/> Presymptomatic diagnosis <input type="checkbox"/> Family history <input type="checkbox"/> Sudden death <b>Note:</b> Genetic testing should always be initiated on an affected family member first, if possible, in order to be most informative for at-risk relatives. See Ethnic Background and Family History section for more information.			
Diagnosis/Suspected Diagnosis <input type="checkbox"/> Marfan Syndrome <input type="checkbox"/> Ehlers-Danlos Syndrome <input type="checkbox"/> Loeyes-Dietz Syndrome <input type="checkbox"/> Familial thoracic aortic aneurysm and dissection <input type="checkbox"/> Other: _____			
Indicate whether the following are present: <input type="checkbox"/> Aortic diameter at sinuses of Valsalva Z-score $\geq 2$ <input type="checkbox"/> Aortic dissection <input type="checkbox"/> Ectopia lentis <input type="checkbox"/> Systemic score $\geq 7$ points (see table to the right for calculation) <input type="checkbox"/> Aortic dilatation/aneurysm (Z-score $< 2$ ) <input type="checkbox"/> Family history of independently diagnosed Marfan syndrome using the revised Ghent criteria	<b>Systemic Score Calculation</b>		
	<b>Feature</b>	<b>Value</b>	<b>Enter Value if Present</b>
	Wrist <b>and</b> thumb sign	3	
	Wrist <b>or</b> thumb sign	1	
	Pectus carinatum	2	
	Pectus excavatum or chest asymmetry	1	
	Hindfoot deformity	2	
	Plain flat foot (pes planus)	1	
	Pneumothorax	2	
	Dural ectasia	2	
	Protrusio acetabulae	2	
	Reduced upper/lower segment <b>and</b> increased armspan/height	1	
	Scoliosis or thoracolumbar kyphosis	1	
	Reduced elbow extension	1	
	3 of 5 facial features: <ul style="list-style-type: none"> <li>• dolichocephaly</li> <li>• enophthalmos</li> <li>• downslanting palpebral fissures</li> <li>• malar hypoplasia</li> <li>• retrognathia</li> </ul>	1	
	Skin striae	1	
	Myopia $> 3$ diopters	1	
	Mitral valve prolapse	1	
		<b>Total</b>	
List any additional features present:			

Patient Name (Last, First, Middle)

Birth Date (mm-dd-yyyy)

### Ethnic Background and Family History

European Caucasian    African American    Hispanic    Asian    Middle Eastern    Other (specify): \_\_\_\_\_

Are other relatives known to be affected?    Yes    No

If yes, indicate their diagnosis and relationship to the patient: \_\_\_\_\_

Have other relatives had molecular genetic testing?    Yes    No

**For known mutation test requests, order known variant analysis:**

KVAR1 / Known Variant Analysis-1 Variant

KVAR2 / Known Variant Analysis-2 Variants

KVAR3 / Known Variant Analysis-3+ Variants

**New York State Patients: Informed Consent for Genetic Testing is required.** See Informed Consent for Genetic Testing (T576), Informed Consent for Genetic Testing – Spanish (T826), or Informed Consent for Genetic Testing for Deceased Individuals (T782).