



Hereditary Hemorrhagic Telangiectasia and Vascular Malformations Gene Panel

Patient Information

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

Patient Name (Last, First Middle)		Birth Date (mm-dd-yyyy)
Sex Assigned at Birth <input type="checkbox"/> Male <input type="checkbox"/> Female <input type="checkbox"/> Unknown <input type="checkbox"/> Choose not to disclose		Legal/Administrative Sex <input type="checkbox"/> Male <input type="checkbox"/> Female <input type="checkbox"/> Nonbinary

Referring Provider Information

Referring Provider Name (Last, First)	Phone	Fax*
Other Contact Name (Last, First)	Phone	Fax*

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

Reason for Testing

Diagnosis Family History** Other, specify: _____

**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.

Indications

Indications

Hereditary hemorrhagic telangiectasia (HHT) Multiple cutaneous and mucosal venous malformations (VMCM)

Hereditary glomuvenous malformations Capillary malformation-arteriovenous malformation syndrome (CM-AVM)

Familial cerebral cavernous malformation (CCM) Other, specify: _____

Clinical History

<input type="checkbox"/> Telangiectasia Location and number: _____	<input type="checkbox"/> Cerebral cavernous malformation Number: _____
<input type="checkbox"/> Epistaxis (nosebleeds) Frequency: _____	<input type="checkbox"/> Retinal vascular malformation
<input type="checkbox"/> Visceral arteriovenous malformations (AVM) Location and number: _____	<input type="checkbox"/> Parkes-Weber syndrome
<input type="checkbox"/> Arteriovenous (AV) fistula Location and number: _____	
<input type="checkbox"/> Capillary malformations Location and number: _____	

Other Relevant Clinical History

Hereditary Hemorrhagic Telangiectasia and Vascular Malformations Gene Panel 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 ***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.
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"/> Choose not to disclose
<input type="checkbox"/> Ashkenazi Jewish	<input type="checkbox"/> European	<input type="checkbox"/> Middle Eastern	<input type="checkbox"/> None of the above	<input type="checkbox"/> Unknown

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).