



GENETIC DIAGNOSTIC LABORATORY
UNIVERSITY OF PENNSYLVANIA SCHOOL OF MEDICINE
DEPARTMENT OF GENETICS
560 Clinical Research Building • 415 Curie Boulevard • Philadelphia, PA 19104
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CLIA ID: 39D0893887

REQUEST FOR HEMOPHILIA A (FACTOR VIII DEFICIENCY)

Please provide the following information. We cannot perform your test without ALL of this information. PLEASE PRINT ALL ANSWERS

PATIENT INFORMATION*		Sample Collection Date & Time: _____		
FIRST NAME	MI	LAST NAME	BIRTH DATE (MM/DD/YYYY)	GENDER
ANCESTRY <input type="checkbox"/> Western/Northern European <input type="checkbox"/> Central/Eastern European <input type="checkbox"/> Latin American/Caribbean <input type="checkbox"/> African				
<input type="checkbox"/> Asian <input type="checkbox"/> Jewish (Ashkenazi) <input type="checkbox"/> American Indian <input type="checkbox"/> Near East/Middle Eastern <input type="checkbox"/> Native Hawaiian or Pacific Islander				
Specify countries: _____ <input type="checkbox"/> Other: _____				
CLINICAL INFORMATION				
<input type="checkbox"/> Patient is affected: <input type="checkbox"/> Mild <input type="checkbox"/> Moderate <input type="checkbox"/> Severe <input type="checkbox"/> Unknown				
Patient's Factor VIII level: _____				
<input type="checkbox"/> Patient is a female and a possible carrier. Please list the affected relative(s)*: _____				
<i>*Please include a family pedigree indicating affected individuals and relationship of at-risk female carriers to affected individuals.</i>				
Has anyone in the patient's family ever had DNA testing for Hemophilia A? <input type="checkbox"/> No <input type="checkbox"/> Yes. If yes:				
Name of person previously tested and relationship: _____				
Was the previous testing performed at the Genetic Diagnostic Laboratory? <input type="checkbox"/> Yes <input type="checkbox"/> No				
Result (Please include a copy of the result): _____				
ICD-10 CODE(S):* <input type="checkbox"/> D66 Hemophilia A <input type="checkbox"/> Z31.43 Testing for genetic disease carrier status				
<input type="checkbox"/> Z83.2 Family history of hemophilia A <input type="checkbox"/> Z14.01 Asymptomatic hemophilia A carrier				
<input type="checkbox"/> Z84.81 Family history of genetic disease <input type="checkbox"/> Other: _____				
TEST REQUESTED*				
Approximately 50-53% of severe hemophilia A cases will have an inversion in either intron 22 or intron 1:				
<input type="checkbox"/> Inversion analysis of intron 22 and 1 of F8 gene				
<input type="checkbox"/> Inversion analysis of introns 22 and 1; if negative, reflex to sequence analysis of F8 gene				
<input type="checkbox"/> Inversion analysis of intron 22 and intron 1; if negative, reflex to sequence analysis; if negative, reflex to deletion/duplication analysis of F8 gene				
Approximately 98% of mild to moderate cases of hemophilia A will have a sequencing mutation:				
<input type="checkbox"/> Sequence analysis; if negative, reflex to deletion/duplication analysis of F8 gene				
<input type="checkbox"/> Sequence analysis; if negative, reflex to deletion/duplication analysis; if negative, reflex to inversion analysis of intron 22 and intron 1 of F8 gene				
<input type="checkbox"/> Sequence analysis of F8 gene				
<input type="checkbox"/> Deletion/duplication analysis of F8 gene				
Other testing options:				
<input type="checkbox"/> Linkage Analysis (please call the laboratory before ordering)				
<input type="checkbox"/> Site specific analysis (familial): Inversion (Intron 22) _____ Inversion (Intron 1) _____ Sequencing _____ Del/Dup _____				
<input type="checkbox"/> PRENATAL site specific analysis (familial):** Inversion (Intron 22) _____ Inversion (Intron 1) _____ Sequencing _____ Del/Dup _____				

*Required information

**Please call the laboratory prior to sending a prenatal sample. Please refer to the special requirements for prenatal samples on the Instructions for Sample Submission page.



Perelman
School of Medicine
UNIVERSITY of PENNSYLVANIA

**GENETIC DIAGNOSTIC LABORATORY
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PATIENT REGISTRATION FORM

Please provide the following information. We cannot perform your test without ALL of this information. PLEASE PRINT ALL ANSWERS

PATIENT INFORMATION

FIRST NAME	MI	LAST NAME	BIRTH DATE	GENDER
STREET ADDRESS				
CITY	STATE	ZIP	PHONE	

PHYSICIAN INFORMATION*

REFERRING PHYSICIAN	PHONE	FAX	
GENETIC COUNSELOR	PHONE	FAX	
EMAIL ADDRESS FOR COUNSELOR	EMAIL ADDRESS FOR PHYSICIAN		
INSTITUTION AND DEPARTMENT			
STREET ADDRESS	CITY	STATE	ZIP

PAYMENT OPTIONS* (must choose one) [a receipt will be mailed to the patient for self-pay options]

- I have enclosed a check payable to the "Genetic Diagnostic Laboratory" for \$ _____

Please charge my credit card for the amount of \$ _____
 VISA Master Card Discover American Express

Card Number: _____ Exp date: _____

Name of cardholder as it appears on card: _____

I have Pennsylvania Medicaid. A copy of my Medicaid card is attached.

INSTITUTIONAL BILLING: The Institution where my testing originated has agreed to pay all charges for the testing.
 INCLUDE Billing Address, Person Authorizing Payment, Telephone, and Fax below:

BILLING ADDRESS

BILLING ADDRESS

INFORMED CONSENT: GENETIC TESTING FOR HEMOPHILIA

Background: Hemophilia A is the most common severe bleeding disorder occurring in approximately 1:4,000 births and inherited as an X-linked recessive disease. Hemophilia B is also an X-linked recessive disease with a birth prevalence of approximately 1:20,000 births. Both diseases have varied clinical presentations from mild to severe disease, and the severity of the disease can be related to the type of molecular alteration.

Purpose: The diagnostic samples will be used for the purpose of attempting to determine if I (or my child/fetus) am/is a carrier of an altered gene known to cause hemophilia. This information may help establish appropriate medical management.

Results: I understand that there are five possible results to this testing:

PATHOGENIC VARIANT: A clinically significant variant is detected in gene(s) analyzed. This may explain my personal or family history of hemophilia. My or my child's healthcare provider will make medical management recommendations based on this information.

LIKELY PATHOGENIC VARIANT: A variant is detected in the gene(s) analyzed which is the likely deleterious. This may explain my personal or family history of hemophilia. My or my child's healthcare provider will make medical management recommendations based on this information.

VARIANT OF UNCERTAIN SIGNIFICANCE: The laboratory may detect an alteration in the gene(s) analyzed which is currently of unknown significance, called a "variant of unknown significance (VUS)". The laboratory will work with my physician to help determine if the VUS can be further classified as to whether it is associated with hemophilia.

LIKELY BENIGN VARIANT: A variant is detected in the gene(s) analyzed which is not likely to be clinically significant. This result reduces the likelihood that I, or my child, have a clinically significant variant in the gene(s) tested.

NEGATIVE: No clinically significant variants were identified in the gene(s) analyzed. This result reduces the likelihood that I, or my child, have a clinically significant variant in the gene(s) tested. Methods currently in use are unable to detect all variants and therefore I may still carry a variant that was not detected by the current technology.

Disclosure Policy: The Genetic Diagnostic Laboratory will release my test results to the ordering healthcare provider or genetic counselor, and otherwise only as permitted by law. The results will be kept confidential to the extent allowed by law. If I provide separate written consent, the lab will release my test results to other medical professionals or third persons I want to receive my results.

Limitations: While genetic testing is highly accurate for detection of the majority of disease causing mutations, a small fraction of mutations may be missed by the current technology. Due to the nature of the testing, there is a small possibility that the test will not work properly or that an error will occur. Occasionally, testing may reveal a variant of unknown significance that is unable to be definitively interpreted as positive or negative for disease-association based on the current knowledge of the variant. The DNA analysis performed at the University of Pennsylvania Genetic Diagnostic Laboratory is specific only for gene(s) analyzed and in no way guarantees my health.

There are federal laws in place that prohibit health insurers and employers from discriminating based on genetic information, such as test results. There currently are no federal laws prohibiting discrimination based on genetic information by life insurance, long term care, or disability insurance companies, but state laws may restrict this. I understand I can ask my ordering provider or genetic counselor for more information about how insurers might use genetic information.

Initials _____

Use of Specimens After Clinical Test Performed: I understand my blood or tissue specimen will not be returned to me or the ordering healthcare provider, and becomes the property of the lab upon receipt. The laboratory is not a DNA banking facility; therefore this is no guarantee that samples will be available or usable for additional or future testing. Samples from New York residents will be disposed of 60 days after clinical testing is complete.

After the laboratory completes the ordered clinical test, the lab may retain and preserve the specimen to validate the development of future genetic tests or for future research or education purposes. The laboratory is committed to continuous improvement and therefore I understand my coded sample may be used to validate a new assay. If testing reveals a clinically significant result during the validation process of a new assay related to the original indication for testing, my health provider may be contacted. If the lab uses the specimen for future research or education purposes, the specimen will be de-identified by removing my personally identifying information. My name, address and other personal identifying information will not be linked to the samples, or the results of the research, and I will not be identified in any research results or publications. I will not receive a copy of the research results. I can decline for my sample to be retained at the lab by filling out "Research Opt Out" form found on the following website: <http://www.med.upenn.edu/genetics/gdl/>.

I understand the lab may wish to contact me, or my ordering healthcare provider, for additional information. The additional information may include, but would not be limited to, information about health and family history that might be relevant to the research. I understand I can decline future contact from the lab by filling out "Research Opt Out" form found on the following website: <http://www.med.upenn.edu/genetics/gdl/>.

Genetic Counseling provided by a qualified specialist (i.e. genetic counselor/ medical geneticist) is a recommendation for individuals proceeding with genetic testing. This service is available before and after genetic testing. Additionally, other testing or further physician consults may be warranted.

The Genetic Diagnostic Laboratory is also an available resource to ask more questions about this testing. The laboratory genetic counselor can be reached at 215-573-9161 and Arupa Ganguly, PhD, FACMG can be reached at 215-898-3122. I will be given a copy of this consent form to keep.

HEALTHCARE PROVIDER STATEMENT:

I have explained to _____ the purpose of this genetic testing, the procedures required and the possible risks and benefits to the best of my ability.

Printed Name of Professional Obtaining Consent

Signature of Professional Obtaining Consent

Date

CONSENT OF PATIENT:

I have read and received a copy of this consent form. I agree to have genetic testing performed for myself, child or my fetus, and accept the risks. I understand the information provided in this document and I have had the opportunity to ask questions I have about the testing, the procedure, the associated risks and the alternatives.

Patient's Printed Name: _____

DOB: _____

Patient's Signature: _____
(or Parent/Guardian if patient is a minor)

Date: _____

Name and Relationship: _____
(Parent/Guardian if patient is a minor)

Initials _____