

MOLECULAR HAEMOSTASIS & THROMBOSIS TEST REQUEST FORM

Molecular Haemostasis & Thrombosis, Viapath Analytics LLP 4th Floor North Wing St. Thomas' Hospital. London, SE1 7EH *T*: 020 7188 6817/2798 viapath.mol.haem@nhs.net

All fields are mandatory. Illegible, unclear or incomplete forms will result in delays or rejection.

CONSENT STATEMENT: It is the referring clinician's responsibility to ensure that the patient/carer knows the purpose of the test and that the sample may be stored for future diagnostic testing. In signing this form the clinician has obtained consent for testing, storage and for the use of this sample and the information gathered from it to be shared with members of the donor's family through their health professionals (if appropriate). The patient should be advised that the sample may be used anonymously for quality assurance and training purposes. If the patient does not wish information to be shared please write this clearly in the clinical summary box.

PATIENT DEMOGRAPHICS					PATIENT ETHNICITY									
First name:									White:	sh 🗆 Irish 🗆 Any Other White Background 🗆				
Last name:									Mixed:	Whi	te And Black Caribbean 🗆 te And Black African 🗆 White And Asian 🗆			
DOB:	Gender:	Male 🗆	🗌 Fe	male 🗆	Oth	ner 🗆		_			Other Mixed Background			
NHS number:									Asian or Asian British:		an 🗆 Pakistani 🗆 Bangladeshi 🗆 Other Asian Background 🗆			
								-	Black or		bbean African Any Other Black			
Hospital no:	Family r	ef no:	10:						Black British:	Background 🗆				
Postcode:	Life stat	us: Alive	Alive Deceased						Other Ethnic	Chir	ese 🗆 Any Other Ethnic Group 🗆			
Non-NHSE funded (please attach invoicir	ng details):							_	Groups:	(please specify:				
		_							Not stated 🗆	NOT	Known 🗆			
CLINICAL INFORMATION AND FAMILY HI	STORY													
Please give as much clinical and genetic ir clinical information provided. Please use										of	Have other members of this family had gene testing? Y/N Please provide details:			
											For familial cases, please include a pedigree with the patient clearly marked:			
Is patient pregnant? Y/N If yes ho	ow many w	eeks ges	station?											
Affected Unaffected Age of onset: Pa				Patients to be tested: Patient only \Box Patient and both parent						nts 🗆 Other 🗆				
CLINICALLY URGENT?														
CLINICIAN DETAILS									- alialate - 1 -		+ (if different)			
					e spons ame:	sponsible clinician / consultant (if different) me:								
Hospital & department:			Hospital & department:											
NHS email:						NHS email:								
Phone:	Phone: Phone:													
SAMPLE TYPE:		TE	ST REQ	UEST:										
Blood EDTA														
(if not EDTA please state)		Gei	ne test:											

Test directory clinical indication and ID (R number):

information provided.

DNA storage only

Other (please specify)

https://www.england.nhs.uk/publication/national-genomic-test-directories/

If the clinical indication and code are not provided, a test or panel will be applied based on the clinical

Extracted DNA

Date of collection:

Time of collection:

For Departmental Use Only:

Amnio 🗆

Other (please state e.g. buccal swab)

РОС 🗆

Fetal blood

cvs 🗆



MOLECULAR HAEMOSTASIS & THROMBOSIS TEST REQUEST FORM

All fields are mandatory. Illegible, unclear or incomplete forms will result in delays or rejection.

CONSENT STATEMENT: It is the referring clinician's responsibility to ensure that the patient/carer knows the purpose of the test and that the sample may be stored for future diagnostic testing. In signing this form the clinician has obtained consent for testing, storage and for the use of this sample and the information gathered from it to be shared with members of the donor's family through their health professionals (if appropriate). The patient should be advised that the sample may be used anonymously for quality assurance and training purposes. If the patient does not wish information to be shared please write this clearly in the clinical summary box.

Patient first name:	Patient last name:	DOB:	NH	IHS no:									

HPO terms phenotypes and presence in this individual – please tick

Please confirm the HPO terms that have been assessed, and select whether they are present or absent

Bleeding disorder	Present	Absent	Thrombocytopenia / Platelet disorder	Present	Absent
Bruising susceptibility <u>HP:0000978</u>			Thrombocytopenia <u>HP:0001873</u>		
Epistaxis <u>HP:0000421</u>			Congenital thrombocytopenia <u>HP:0001905</u>		
Prolonged bleeding after surgery <u>HP:0004846</u>			Macrothrombocytopenia <u>HP:0040185</u>		
Prolonged bleeding after dental extraction <u>HP:0006298</u>			Abnormal platelet function <u>HP:0011869</u>		
Gingival bleeding <u>HP:0000225</u>			Abnormal platelet granules <u>HP:0011883</u>		
Petechiae <u>HP:0000967</u>			Glanzmann Thrombasthenia <u>ORPHA:849</u>		
Menorrhagia <u>HP:0000132</u>			Bernard Soulier syndrome ORPHA:274		
Post partum haemorrhage <u>HP:0011891</u>			Gray Platelet syndrome ORPHA:721		
Intracranial haemorrhage HP:0002170			Thrombotic Thrombocytopenic Purpura <u>ORPHA:54057</u>		
Cerebral haemorrhage HP:0001342					
Gastrointestinal haemorrhage HP:0002239					

Thrombosis / Thrombophilia	Present	Absent
Venous thrombosis <u>HP:0004936</u>		
Deep venous thrombosis <u>HP:0002625</u>		
Pulmonary embolism <u>HP:0002204</u>		
Arterial thrombosis <u>HP:0004420</u>		
Stroke <u>HP:0001297</u>		
Cerebral venous thrombosis HP:0005305		

Other (please specify)	Present	Absent
Sensorineural hearing		
impairment <u>HP:0000407</u>		
Presenile cataracts <u>HP:0007819</u>		
Renal insufficiency <u>HP:0000083</u>		
Ocular albinism <u>HP:0001107</u>		
Neutrophil inclusion bodies		
<u>HP:0008264</u>		

Note: Please ensure the latest version of this request form is used, found on our website: www.southeastgenomics.nhs.uk