

KRCC MOLECULAR PATHOLOGY TEST REQUEST FORM

Please send all samples to:

Synnovis Analytics Molecular Pathology Laboratory c/o Central Specimen Reception

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

Blood Sciences Laboratory Ground Floor Bessemer Wing King's College Hospital, Denmark Hill

1-5µg genomic DNA

London SE5 9RS Tel: 020 3299 2265 Email: kch-tr.PND@nhs.net

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:	British 🗆 Irish 🗆 Any Other White Background 🗆	
Last name: DOB: Gender: Male Female Other					Mixed:	White And Black Caribbean □ White And Black African □ White And Asian □ Any Other Mixed Background □	
NHS number:				Asian or Asian British:	Indian □ Pakistani □ Bangladeshi □ Any Other Asian Background □		
Hospital no:		Family ref no:		Black or Black British:	Caribbean African Any Other Black Background		
Postcode:		Antenatal: Yes 🗆 No 🗆		Other Ethnic Groups:	Chinese Any Other Ethnic Group (please specify:		
Non-NHSE funded (please attach invoicing details): \Box					Not stated	Not Known 🗆	
PATHOLOGY RESULTS Haematology indices					SAMPLE REQUIREMENTS		
Iron / liver parameters	Hb		HbF %		For haemoglob	inopathy investigation:	2 x 4 ml EDTA blood
Serum Iron	RBC		HbA2 %		Children and ad	lults (all other tests):	4 ml EDTA blood
Serum TSat	MCV		Hb variant %		Infants:		1 ml EDTA blood
Serum Bilirubin	МСН		Absolute Reticulocyte		As DNA for Nex	t Generation Sequencing:	3-5μg genomic DNA

Reason for referral / family details: Is the patient affected? \Box

Platelets

(Please ensure sufficient information is included to demonstrate that the patient meets NHS England eligibility criteria for desired test, where applicable)

Reticulocyte %

As DNA for all other tests:

Date sample collected:

Molecular Tests - For Haemoglobinopathy referrals FBC and HPLC results MUST be provided							
R93 Hb variant identification	R176 Gilbert's genotyping (TAs	5/6/7/8 repeat)					
□ R93 Haemoglobinopathy investigations	R95 Hereditary haemochromatosis (HFE - C282Y and H63D variants)						
R93 Alpha thalassaemia	R191 Alpha-1-antitrypsin genotype (S and Z alleles)						
R93 Beta thalassaemia	Thrombophilia genetic screen	(please tick all that apply):					
□ Other (please state gene):	FVL 🗆 PT 🗆 M						
	 R240/242/244 Familial variant testing (please state familial variant): 						
For further details of each test please refer to the <u>Synnovis website</u> .							
Next Generation Sequencing - Please select which panel(s) are required							
🗆 R91 Cytopenia (NOT Fanconi anaemia)	R96 Iron metabolism disorders	🗆 R92 Rare Anaemia					
Neutropenia	R168 Porphyria	Membranopathy					
Diamond-Blackfan anaemia	R169 Acute intermittent porphyria	Enzymopathy					
Inherited bone marrow failure	R170 Variegate porphyria	Haemoglobinopathy					
□ R313 Neutropenia consistent with ELANE mutations	R347 Inherited predisposition to AML	Congenital dyserythropoietic anaemia					
Пнін	□ R366 Inherited predisposition to ALL	 Diamond-Blackfan anaemia Sideroblastic anaemia 					
□ Single gene analysis:(name of gene) 🛛 R405 Hereditary erythrocytosis						
	R406 Thrombocythaemia	Haemolytic anaemia					
	□ R323 Sitosterolaemia						
For details of genes in each subpanel, please refer to the Synnovis website							
CLINICIAN DETAILS							

Requesting clinician / consultant	Responsible clinician / consultant (if different)		
Name:	Name:		
Hospital & department:	Hospital & department:		
NHS email:	NHS email:		
Phone:	Phone:		

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

SE GLH KRCC FORM v2

Serum

Ferritin

Blood Film comments: