

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. Testing may be performed at Synnovis, any other NHSE GLH or by other international laboratories where necessary. 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		SAMPLE TYPE:	
First name: <input type="text"/>		Blood EDTA <input type="checkbox"/> for DNA or gene tests	
Last name: <input type="text"/>		Lithium Heparin <input type="checkbox"/> for karyotype only	
DOB: <input type="text"/>	Biological sex: Male <input type="checkbox"/> Female <input type="checkbox"/> Other <input type="checkbox"/>	CVS <input type="checkbox"/> Amnio <input type="checkbox"/> Fetal blood <input type="checkbox"/> POC <input type="checkbox"/> RNA <input type="checkbox"/>	
NHS number: <input type="text"/>	<input type="text"/>	Histology sample <input type="checkbox"/> specimen number	
Hospital no: <input type="text"/>	Family ref no: <input type="text"/>	Tissue type	
Postcode: <input type="text"/>	Life status: Alive <input type="checkbox"/> Deceased <input type="checkbox"/>	Other (please state)	
Non-NHSE funded (please attach invoicing details): <input type="checkbox"/>		Date of collection: <input type="text"/>	Time of collection: <input type="text"/>
Ethnicity: <input type="text"/>		For Departmental Use Only:	

CLINICAL INFORMATION AND FAMILY HISTORY	
Please give as much clinical and genetic information as possible. Interpretation of results depends on the quality of clinical information provided. Please use HPO terms (https://hpo.jax.org/app/) when possible, see reverse.	
Affected <input type="checkbox"/> Unaffected <input type="checkbox"/>	
Age of onset:	
For familial cases, please include a pedigree with the patient clearly marked:	
Is patient pregnant? Y/N If yes how many weeks gestation? <input type="text"/>	
Have other members of this family had gene testing? Y/N	
Please provide details:	
Has this patient had a bone marrow transplant or a blood transfusion? Yes / No	

TEST REQUEST:	
URGENT <input type="checkbox"/>	Routine <input type="checkbox"/>
If you are a member of a Clinical Genetics team, is this test: For a consultant <input type="checkbox"/> For a proband <input type="checkbox"/> A diagnostic / predictive / carrier test	If you are not a member of a Clinical Genetics team, is this test diagnostic? Yes / No
Test Directory ID number: <input type="text"/> (R or M code) <i>This is a mandatory field.</i>	Array CGH <input type="checkbox"/>
<i>Please ensure that this referral matches the testing criteria:</i> https://www.england.nhs.uk/publication/national-genomic-test-directories/	QF-PCR (rapid aneuploidy) <input type="checkbox"/>
	Karyotype analysis <input type="checkbox"/>
	Storage only (no test activation) <input type="checkbox"/>
	Other (please specify)

CLINICIAN DETAILS	
Requesting clinician / consultant Name: <input type="text"/> Hospital & department: <input type="text"/>	Reporting clinician/ consultant (if different) Name: <input type="text"/> Hospital & department: <input type="text"/>
NHS email: <input type="text"/> Phone: <input type="text"/>	NHS email: <input type="text"/> Phone: <input type="text"/>

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

