NHS South East Genomic Laboratory Hub

TEST REQUEST FORM

Genetics Laboratories, 5th Floor, Tower Wing, Guy's Hospital, Great Maze Pond, London, SE1 9RT

> T: 020 7188 1696/1709 gst-tr.viapathgeneticsadmin@nhs.net gst-tr.londonsouthglh@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				•		Γ	PATIENT ETHNI				
First name:				Ē	White:	British 🗆 Irish 🗆 Any Other White Background 🗆					
Last name:									d Black Caribbean 🗆		
Last name:							Mixed:		d Black African □ White And Asian □ r Mixed Background □		
DOB: G	ender: Male 🗆	Fema	le 🗆 O	ther 🗆		-	Asian or		Pakistani 🗆 Bangladeshi 🗆		
NHS number:							Asian British:	,	r Asian Background		
Hospital no: Fa	amily ref no:						Black or Black British:	Caribbea Backgrou	n 🗆 African 🗆 Any Other Black nd 🗆		
Postcode: Li	ife status: Alive 🗆] Decea	ised 🗆				Other Ethnic Groups:	Chinese [Any Other Ethnic Group (please specify:)		
Non-NHSE funded (please attach invoicing d	etails): 🗆					Ľ	Not stated	Not Knov			
CLINICAL INFORMATION AND FAMILY HISTO	RY										
Please give as much clinical and genetic inform information provided. Please use HPO terms		prg/app/)) when pos					or clinical	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:		
Affected Unaffected Age of ons	set: Pati	ients to b	oe tested: F	Patient c	onlv □		Patient and both	parents 🗆	Other 🗆		
CLINICALLY URGENT?					,						
CLINICIAN DETAILS Requesting clinician / consultant				Res	nonsi	ihl	e clinician / cons	ultant <i>(if di</i>	ifferent)		
Name: Hospital & department:				Na	Name: Hospital & department:						
NHS email: Phone:					S ema one:	ail					
SAMPLE TYPE:	TEST R	EQUEST:									
Blood EDTA 🛛 for DNA or gene tests	Gene to	est:									
Lithium Heparin 📮 for karyotype only											
CVS 🕀 Amnio 🕀 Fetal blood 🕀 PG	DC 🗔 Test dir	rectory c	linical indi	cation a	nd ID	(R	number):				
Other (please state)	https://	/www.en	igland.nhs.	uk/publ	icatio	n/	national-genomi	c-test-direc	tories/		
	If the cl	inical ind	i cation and	code are	: not p	pre	wided, a panel w	ill be applied	d based on the clinical information provided.		
Date of collection:	Array 0	сөн 🗆									
	QF-PCF	R (rapid a	aneuploidy	') 🗆							
Time of collection: For Departmental Use Only:	Karyot	ype anal	ysis 🗆			_					
	DNA st	orage or	nly 🗆								
						_					

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

Other (please specify)

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Patient first name:	Patient last name:	DOB:	NHS 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

Intellectual disability, developmental and metabolic	Present	Absent	Neurology	Present	Absen	
Intellectual disability - mild			Muscular dystrophy			
Intellectual disability - moderate			Myopathy			
Intellectual disability - profound			Myotonia			
Intellectual disability - severe			Fatigable weakness			
Autistic behaviour			Peripheral neuropathy			
Global developmental delay			Distal arthrogryposis			
Delayed fine motor development			Arthrogryposis multiplex congenita			
Delayed gross motor development			Cognitive impairment			
Delayed speech and language development			Parkinsonism			
Generalized hypotonia			Spasticity			
Feeding difficulties			Chorea			
Failure to thrive			Dystonia			
Abnormal facial shape			Ataxia			
Abnormality of metabolism/homeostasis			Cerebellar atrophy			
Microcephaly			Cerebellar hypoplasia			
Macrocephaly			Dandy-Walker malformation			
Tall stature			Olivopontocerebellar hypoplasia			
	1		Diffuse white matter abnormalities		1	
			Focal White matter lesions			
		1	Leukoencephalopathy			
Craniosynostosis	Present	Absent	Cortical dysplasia			
Bicoronal synostosis			Heterotopia			
Unicoronal synostosis			Lissencephaly			
Metopic synostosis			Pachygyria			
Sagittal craniosynostosis			Polymicrogyria			
Lambdoidal craniosynostosis			Schizencephaly			
Multiple suture craniosynostosis			Holoprosencephaly			
· · · · · ·			Hydrocephalus			
Skeletal dysplasia	Present	Absent				
Disproportionate short stature	1 resent	Abbent	Diabetes	Present	Absen	
Proportionate short stature			Neonatal insulin-dependent diabetes	1 resent	7105011	
Short stature			mellitus			
Skeletal dysplasia			Transient neonatal diabetes mellitus			
			Renal	Present	Absen	
Epilepsy	Present	Absent	Multiple renal cysts			
Seizures		ļ	Nephronophthisis	_		
Generalized-onset seizure		ļ	Hepatic cysts	_		
Focal-onset seizure		ļ	Enlarged kidney	_		
Epileptic spasms		ļ		_		
Infantile encephalopathy						
Atonic seizures						
Generalized myoclonic seizures		ļ	Other (please specify)	Present	Absen	
	1			_		
Generalized tonic seizures					1	
Generalized tonic-clonic seizures						
Generalized tonic-clonic seizures EEG with focal epileptiform discharges						
Generalized tonic-clonic seizures EEG with focal epileptiform discharges EEG with generalized epileptiform						
Generalized tonic-clonic seizures EEG with focal epileptiform discharges EEG with generalized epileptiform discharges						
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