## **Genomic Medicine Service**

Whole Genome Sequencing (WGS) Test Request PLEASE DO NOT USE FOR NON-WGS TESTS

RARE AND INHERITED DISEASES
FAMILY MEMBER



Requesting organisation:						
GLH laboratory to receive	e sample:					
First name		Life status Alive Deceased	Ethnicity (Please tick	c on Page 2)		
Last name		HPO terms ( <a href="https://hpo.jax.org/app/">https://hpo.jax.org/app/</a> ) phenotypes & presence in this individual (if relevant, please list below or				
Date of birth (dd/mm/yyyy)	Hospital number	tick on page 2)				
Gender  Male  Female	Other	Specific rare or inherited have been confirmed (ON		•		
NHS Number (or postcode			,	o (		
THIS TRAINSET (OF POSICOGO						
Patient not eligible for NHS number not of the Other (please provide reason)	umber (e.g. foreign national)					
Additional clinical information Please include any previous molecu	ation (if relevant) ular testing with date(s) and any other	r pertinent clinical information				
Relationship to proband	For the condition being to disease status  Affected Unaffect	ested, please describe the in	dividual's nown	Age of onset (if relevant) State in years and months		
Proband details						
Proband first name		Proband NHS nu	ımber (or po	stcode if not known)		
Proband last name			Proband	date of birth (dd/mm/yyyy)		
Test request						
Test required Whole Genome Sequence	•	al Indication & code (reason	for testing)			
Samples (being sent to GL	LH DNA extraction lab)					
_ `	Amniotic fluid Fetal b	_	Fresh	Tissue (not tumour)		
Sample ID	Collection date / time	Sample volume if applicat	ole Com	ments		
·						
Clinician details						
Responsible clinician / cor	nsultant	Main contact (if different from responsible clinician/consultant)				
Name:		Name:				
Department address:		Department address:				
Phone:		Phone:				
Email:		Email:				

Version 1.15 Page 1 of 2

I have attached a copy of the Record of Discussion form

Patient conversation taken place; Record of Discussion form to follow

First name	Last name	Date of birth (dd/mm/yyyy)	NHS number (or postcode if not		t kn	known)									
															i

## 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  Intellectual disability - mild  Intellectual disability - moderate  Intellectual disability - profound  Intellectual disability - severe  Autistic behaviour  Global developmental delay  Delayed fine motor development  Delayed gross motor development  Delayed speech and language development  Generalized hypotonia  Feeding difficulties  Failure to thrive  Abnormal facial shape  Abnormality of metabolism/homeostasis  Microcephaly  Tall stature	r icase committee in o term	is that ha	
Intellectual disability - mild Intellectual disability - moderate Intellectual disability - profound Intellectual disability - severe Autistic behaviour Global developmental delay Delayed fine motor development Delayed gross motor development Delayed speech and language development Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly	**	Present	Absent
Intellectual disability - moderate Intellectual disability - profound Intellectual disability - severe Autistic behaviour Global developmental delay Delayed fine motor development Delayed gross motor development Delayed speech and language development Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	metabolic		
Intellectual disability - profound Intellectual disability - severe Autistic behaviour Global developmental delay Delayed fine motor development Delayed gross motor development Delayed speech and language development Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly	Intellectual disability - mild		
Intellectual disability - severe Autistic behaviour Global developmental delay Delayed fine motor development Delayed gross motor development Delayed speech and language development Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Intellectual disability - moderate		
Autistic behaviour Global developmental delay Delayed fine motor development Delayed gross motor development Delayed speech and language development Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Intellectual disability - profound		
Global developmental delay  Delayed fine motor development  Delayed gross motor development  Delayed speech and language development  Generalized hypotonia  Feeding difficulties  Failure to thrive  Abnormal facial shape  Abnormality of metabolism/homeostasis  Microcephaly  Macrocephaly	Intellectual disability - severe		
Delayed fine motor development Delayed gross motor development Delayed speech and language development Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Autistic behaviour		
Delayed gross motor development  Delayed speech and language development  Generalized hypotonia  Feeding difficulties  Failure to thrive  Abnormal facial shape  Abnormality of metabolism/homeostasis  Microcephaly  Macrocephaly	Global developmental delay		
Delayed speech and language development Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Delayed fine motor development		
Generalized hypotonia Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Delayed gross motor development		
Feeding difficulties Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Delayed speech and language development		
Failure to thrive Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Generalized hypotonia		
Abnormal facial shape Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Feeding difficulties		
Abnormality of metabolism/homeostasis Microcephaly Macrocephaly	Failure to thrive		
Microcephaly Macrocephaly	Abnormal facial shape		
Macrocephaly	Abnormality of metabolism/homeostasis		
	Microcephaly		
Tall stature	Macrocephaly		
	Tall stature		

Craniosynostosis	Present	Absent
Bicoronal synostosis		
Unicoronal synostosis		
Metopic synostosis		
Sagittal craniosynostosis		
Lambdoidal craniosynostosis		
Multiple suture craniosynostosis		

Skeletal dysplasia	Present	Absent
Disproportionate short stature		
Proportionate short stature		
Short stature		
Skeletal dysplasia		

Epilepsy	Present	Absent
Seizures		
Generalized-onset seizure		
Focal-onset seizure		
Epileptic spasms		
Infantile encephalopathy		
Atonic seizures		
Generalized myoclonic seizures		
Generalized tonic seizures		
Generalized tonic-clonic seizures		
EEG with focal epileptiform discharges		
EEG with generalized epileptiform discharges		
Multifocal epileptiform discharges		

Neurology	Present	Absent
Muscular dystrophy		
Myopathy		
Myotonia		
Fatigable weakness		
Peripheral neuropathy		
Distal arthrogryposis		
Arthrogryposis multiplex congenita		
Cognitive impairment		
Parkinsonism		
Spasticity		
Chorea		
Dystonia		
Ataxia		
Cerebellar atrophy		
Cerebellar hypoplasia		
Dandy-Walker malformation		
Olivopontocerebellar hypoplasia		
Diffuse white matter abnormalities		
Focal White matter lesions		
Leukoencephalopathy		
Cortical dysplasia		
Heterotopia		
Lissencephaly		
Pachygyria		
Polymicrogyria		
Schizencephaly		
Holoprosencephaly		
Hydrocephalus		

Diabetes	Present	Absent
Neonatal insulin-dependent diabetes mellitus		
Transient neonatal diabetes mellitus		

Renal	Present	Absent
Multiple renal cysts		
Nephronophthisis		
Hepatic cysts		
Enlarged kidney		
Other (please specify)	Present	Absent

## Ethnicity – Please tick the relevant Self Defined Ethnicity code below

	•		•	
White	Mixed	Asian or Asian British	Black or Black British	Other Ethnic Groups
A British	D White and	H Indian	M Caribbean	R Chinese
	Black Caribbean			
B Irish	E White and	J Pakistani	N African	S Any other
	Black African		_	ethnic group
C Any other	F White	K Bangladeshi	P Any other	Z Not stated
White background	and Asian		Black background	
	G Any other	L Any other Asian background		99 Not known

## Additional local identifiers – please use the table below, these will also be displayed in the interpretation portal

Type – O (Test order/Case ID), P (local patient ID), S (local sample identifier)						
Organisation	Туре	Identifier				

Version 1.15 Page 2 of 2