NHS Genomic Medicine Service, WGS Test Request Rare Disease, August 2021, v1.2 to be used for WGS golive. This document is subject to version control and is regularly updated. Please confirm you are using the current version by contacting your local Genomic Laboratory Hub

## **Genomic Medicine Service**

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



Requesting orga	anisation:											
GLH laboratory:												
- 1 11 61 .												
Proband's first name				status			Ethnicity					
Drohand's last name						Decea	isea					
Proband's last name					nily test		Trio	Othor	Inrovido n	umbai	٠١.	
Date of birth (dd/mm/yyyy) Hospital number					Singleton evant clinic				(provide n	umbei	<u>)·</u>	
Date of birtir (ad/r	nm/yyyy) Tiospitai	Hullibel							cular testing wi	th date(s) and	any othe	r pertinent
Gender		Please state	in clinico	linformation		cal information				, ,		
	emale Oth		ypic and	or phenotypic								
Postcode												
NHS number		<u> </u>										
Reason NHS Nur	mher not availah	ıle.										
	ble for NHS number (e		tional)									
Other (please pr	rovide reason):											
Test request												
Clinically urgent	Test Directo	ory Clinical	Indi	cation & d	code	e (reason fo	or tes	ting)		Proband's	s age c	of onset
										у	ears	months
Additional papal	(s) (if relevant, m		fa [	200/	D:			C	-:f: - u - u u	ام ما شام ما سا	d:	
Additional panel (use panels with panels)		-	101 1	109)	Disease penetrance   Specific rare or inherited diseases that   are suspected or have been confirmed							
http://panelapp.genom					Complete							
						Incomplet	:e					
Family members	to be tested (no	ot required <sub>.</sub>			ly ref	ferrals)						T
First name	Last name	Date of birth		Number ostcode if			Relationship to proband					
			not	known)								
	CILL BAIA		1.1.7	. , .					,			
Samples being se	ent to GLH DNA (	extraction	iab (	only requi	red ij	f also using	this fo	orm fo	or sample co	llection)	l	
First name	Last name	Date of b	oirth	Sample II	D	Collection date / time		Sar	mple type	Sample volume	Co	mments
						uate / tilli	-			volulile		
Responsible clini	ician / consultar	nt			Ma	ain contac	<b>t</b> (if di	ifferer	nt from resp	onsible clini	cian/co	nsultant)
Name:				Name:								
Department address:				De	partment	addre	ess:					
Phone:					Phone:							
Email:					Email:							

I have attached a copy of the Record of Discussion form for all individuals

Patient conversation taken place; Record of Discussion form to follow

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Proband first name	Proband last name	Date of birth (dd/mm/yyyy)	NHS number

HPO terms are important for the analysis and interpretation of WGS data.

Please enter valid HPO terms present in the proband/family members being tested

HPO terms can be copied from the lists below

HPO Terms - Please ensure those given match those available at						
(https://hpo.jax.org/app/)	Present	Absent	Present	Absent	Present	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
Macrocephaly
Tall stature

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

Skeletal dysplasia
Disproportionate short stature
Proportionate short stature
Short stature
Skeletal dysplasia

Neonatal insulin-dependent diabetes mellitus
Transient neonatal diabetes mellitus

Diabetes

Renal
Multiple renal cysts
Nephronophthisis
Hepatic cysts
Enlarged kidney
Renal insufficiency

Neurology
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
Neurodegeneration
Dementia

Epilepsy
Seizures
Generalized seizures
Focal seizures
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

Cardiology	
Hypertrophic cardiomyopathy	
Dilated cardiomyopathy	
Cardiomyopathy	

Eye Disorders
Cataract
Retinal dystrophy
Macular dystrophy
Microphthalmia
Anophthalmia
Coloboma
Developmental glaucoma
Aniridia
Abnormal anterior eye segment morphology
Nystagmus

Immune Disorders
Immunodeficiency
Abnormal lymphocyte morphology
Abnormal lymphocyte physiology
Abnormal lymphocyte count
Abnormality of neutrophils
Abnormality of humoral immunity
Abnormal inflammatory response
Abnormality of complement system

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