

Eurofins Biomnis Genetic Test Request, Information & Consent Form

PATIENT DETAILS

REQUESTING HOSPITAL/ CLINIC DETAILS Surname: _____ Hospital / Clinic Name: Forename: ____ Department: Date of Birth: / / Male Female Sex: Hospital/Clinic No.: Laboratory No.: Ward: Phone: Fax: Physician: ___ PLEASE REMEMBER ALWAYS TO COMPLETE THE INFORMED CONSENT SECTION **TESTS REQUIRED** MOLECULAR GENETICS **TEST NAME SELECT TEST NAME** CODE **SELECT** CODE **Huntingtons Disease** HUNT П Array CGH Analysis CGH MTHFR Mutation C677T **MTHFR** Chromosome YMicrodeletions ΥQ Muscular Dystrophy DUCH (Duchenne's) Cystic Fibrosis Screen CF36 (mostcommon mutations) **PTMUT** Prothrombin Factor V Leiden PCR FAC5 (Factor II) Mutation Rett's Syndrome **RETT** Fragile X Chromosome **FRAGX** PAI-1 Mutation PAI1M Haemochromatosis HFE Other Please Specify: _ **CYTOGENETICS TEST NAME** CODE **SELECT TEST NAME** CODE **SELECT** Chromosome Analysis / Prader Willi Syndrome **PRADW** KARY Karyotyping - Whole Blood (15q11-13Methylation) Chromosome Analysis -William's Syndrome WILL Products of Conception KARPP Other Please Specify: **ONCOGENETICS SELECT SELECT TEST NAME** CODE **TEST NAME** CODE Chromosome Analysis/ Bone Marrow Philadelphia Chromosome (Bone PHIL KARYB (Cytogenetic Bone Marrow) Marrow) Philadelphia Chromosome **PHILB** Other Please Specify: (Whole Blood) Other Tests Required **SAMPLE DETAILS**

Form RQF36 Issue No.: 2.03 Active Date: 16/12/2019 Page 1 of 2

Specimen Type: _____

Specimen Collection Date: ____/___/



Eurofins Biomnis Genetic Test Request, Information & Consent Form

CLINICAL INFORMATION Please note full clinical information is essential: Ple	ease sp	pecify the condition/syndro	me sı	uspected clinically, if	
known:					
HAEMATOLOGICAL KARYOTYPE		CONSTITUTIONAL KARYOTYPE			
Indication (necessary for conclusive interpretat	ion)	In Infants			
		Small Birth Weight		Sexual Ambiguity	
Acute Leukaemia (AL):		Hypotonia		Dysmorphic Syndrome	
Acute Lymphoid Leukaemia (ALL)		Malformation Syndrome			
Acute Myeloid Leukaemia (AML)		lm abildran	_		
Chronic Myeloid Leukaemia		In children Developmental Delay		Psycho-Motor Delay	
Chronic Lymphoblastic Leukaemia				r syone moter Bolay	
Lymphoma		In adolescents			
Myelodycplastic syndrome (MDS)		Girls: Delayed Puberty		Boys: Gynecomastia	
Myelodysplastic syndrome (MDS) Myeloproliferative syndrome		Boys: delayed puberty			
Fanconi anemia		In adults:			
		Multiple miscarriages:		Number:	
Recent bone marrow transplant		Sterility or hypofecundity			
		Male infertility / abnormal			
Other (specify):		sperm Primary or accordant			
		Primary or secondary amenorrhea			
Immuno:		Pre IVF			
FAB Type:		Pre ICSI			
IMPOPTANT: Please note that in accordance with a	ood clir	nical practice we will automati	cally r	porform additional tosts fo	r on
<u>IMPORTANT</u> : Please note that in accordance with good clinical practice we will automatically perform additional tests for an accurate diagnosis where required. This will incur further charges and, where applicable, please ensure your patient is aware of					
this. We recommend that you obtain signed consent from the patient that they will accept such charges.					
INFORMED CONSENT SECTION					
Patient or Guardian:	fully in	formed by the Dector/Dath	ologi	at/ Canaticiat	
I/we the undersigned confirm that I/we have been fully informed by the Doctor/Pathologist/ Geneticist regarding cytogenetic and/or molecular genetic tests that will be performed on cells					
and/or DNA extracted from my/our child's blood and/or tissue to:					
o confirm or exclude the diagnosis					
o determine heterozygote status wit o examine gene locus/loci.	h a vie	ew to obtaining genetic cou	ınsell	ing.	
o examine gene locus/loci. I/we give my/our consent to such testing and confi	rm tha	t I/we have received all the	e nec	essary information	
according to the law.					
D // //O // O		D 4			
Patient/Guardian Signature:		Date: _		//	
Doctor/ Pathologist/Genetic Consul	tant				
The Cytogenetic and/or molecular genetic test information is to be given by the Clinical Pathologist prescribing the					
test, or by the Physician collecting the sample. All	releva	nt issues regarding the inv	olved	d pathology etiology,	
development, prognosis and potential treatment m					
and clearly understood by the patient. All informati Biomnis. The result will be reported to the Physicia			e WIII	be retained by Eurofins	;
Biominio. The result will be reported to the Fiftysion	ari Orny	•			
Doctor/Pathologist Signature:		Date: _			