

NIPT REFERRAL FORM FOR FETAL ANOMALIES (PILOT STUDY)

PATIENT DETAILS:		
SURNAME	/	ADDRESS:
FIRST NAME (S)		
		DOCTOODE
DATE OF BIRTH		POSTCODE:
NHS NUMBER		
HOSPITAL NUMBER		
CLINICIAN DETAILS:		
Hospital:	Leac	d Professional:
Requesters signature:		Date: / /
Email address for return of results:		
PREGNANCY INFORMAT	ION:	
Singleton pregnancy YE	es/No	The following have been excluded (please tick box
Twin pregnancy YES/NO		to confirm)
Multiple fetal anomalies strongly suggestive of trisomy 13,		my 13, Blood transfusion within 4 months
18 or 21		
Please list		Transplant surgery
Duodenal atresia with additional anomalies		Immunotherapy/ stem cell therapy
Plea <mark>se l</mark> ist additional anomalies		Maternal malignancy
		Multiple pregnancy >2 fetuses
Isolated AVSD		
Isolated exomphalos		Pregnancy with vanishing twin
Isolated cystic hygroma*		
isolated cystic hygronia		Failure to complete this section will lead to a delay in
* The NIPT test does not c	heck for Turner syndrome whic	ch is an the testing of this sample
important cause of cystic h		
Gestation by scan:		
Specimen taken by: (Please print) Sign:		
Date of collection: / / Time:		
Sample requirements:		
10ml of maternal blood in Streck tube. Please send to laboratory as soon as possible following sample collection. Please inform the laboratory that a sample is on its way; by tel: 029 218 44072.		
Laboratory contact details:		
Address: All Wales Genomics Laboratory, Institute of Medical Genetics, University Hospital of		
	Wales, Heath Park, Cardiff. CF14 4XW	
Laboratory working hours: Monday - Friday 08.30-17.00		
For further enquiries please email <u>lab.genetics@wales.nhs.uk</u> or ring 029 218 44072.		