

Biomnis

Test request form

Antenatal diagnostics on invasive sampling

INTERNATIONAL DIVISION

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Request Identification Request Form Number (Numéro du patient)

Your reference (Référence)

Customer Identification

Compulsory Stick
your laboratory identification sticker here

PATIENT					
First name(s):					
Surname:					
Maiden name:					
Address:					
Post code: City:					
Country:					
Tel.:					
Date of birth:					
Date of last period:					
Expected due date:					
Name of spouse :					
Date of birth:					
CLINICAL DETAILS, ME	DICAL BACKGROUND				
	three miscarriages: OUI NON				
	liseases in the family: OUI NON				
_					
PRESCRIBER					
First name(s):					
Surname:					
Address:					
Post code:	City:				
Country:					
Tel.:					
Fax:					
PRACTITIONER					
Client number:	1				
First name(s):					
Surname:	ser stamp				
Surname: Address:	Practition				
Post code:	City:				
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IVI					

INFORMATION AND CONSENT FOR PREGNANT WOMEN

for the collection of a sample and the performance of one or more tests for the purposes of prenatal diagnostics in utero (as per Article R. 2131-1 of the French Public Health Code)

I, the undersigned, ▶ confirm that the doctor (Surname, First Name) :

as part of a medical consultation on (date):

I received the following:

1. Has given me information about:

- the risk of the unborn child being born with a particularly serious condition:
- · the characteristics of this condition;
- · the means of diagnosing it;
- the possible opportunities offered by foetal medicine for the treatment or care of the child after birth;
- 2. Has given me information on the laboratory tests that could establish a prenatal diagnosis in utero; these tests have been offered to me and I wish to receive them:
- this (these) test(s) require(s) a sample of amniotic fluid, chorionic villi (placenta), foetal blood, or other foetal sample;
- the methods, risks, constraints and possible consequences of each sampling technique needed to perform this (these) test(s) have been explained to me;
- I have been informed that a second sample may be needed in the event of technical failure; in this event, I will need to sign a new written consent form;
- I have been informed about possible conditions other than those initially investigated that could be revealed by the examination;
- I have been informed that the result of the test will be given to me and explained by the doctor who prescribed it to me.

I give my consent for the sampling of (required for the conduct of the test(s)) (*):

- · amniotic fluid
- · chorionic villi
- · foetal blood
- other foetal sample (please specify)

I also give my consent for the test or tests (*) for which the sample is being collected:

- cytogenetic tests, including molecular tests applicable for cytogenetics
- molecular genetics tests
- · foetal biochemistry tests for diagnostic purposes
- · laboratory tests for the diagnosis of infectious diseases

This (or these) test(s) will be carried out in a medical laboratory that is duly authorized by the regional health agency to conduct them.

The original copy of this document will be stored in my medical records.

A copy of this document will be given to me and to the person responsible for conducting the tests.

The medical laboratory in which the person responsible for conducting the tests is employed shall keep this document under the same conditions as the report of the examination.

(*) Delete as applicable

Date						_	_
	Sia	natu	re of	fthe	clini	cian	

Signature of the interested party

- Sample date:	Weeks of amenorrhea:				
- Sample type	_				
Amniotic fluid (CLA ou CHN)	Chronic villi sampling (CVS)				
Tube 1 ▶ Volume: mL	Products of conception (PoC)				
▶ Appearance: □ clear □ brown □ bloody	Feetal blood (FB)				
Tube 2 ▶ Volume: mL	DNA extracted from: (specify the nature of the sample)				
▶ Appearance: □ clear □ brown □ bloody	In cases of twin pregnancy, please indicate the number of fœtus sampled:				
TEST REQUEST					
☐ FETAL STANDARD KARYOTYPE					
☐ FETAL DNA MICROARRAY (SNP ARRAY) (SNPRE) - Please ☐ Risk of trisomy 21 by MSM greater than or equal to 1/50 - The ☐ Positive NIPT - The report DPNI-ADNIc must be supplied ☐ Ultrasound abnormalities - The ultrasound report must be supplied ☐ Parental chromosomal anomalies - The report must be supplied ☐ The couple has a previous history of pregnancy with an abnormal age > 38 years [without trisomy 21 screening (MSN☐ Personal request: ☐ Other:	e report must be supplied ed from the report must be supplied from the report must be supplied from the report must be supplied from the report must be supplied				
☐ IN SITU HYBRIDISATION (FISH)					
- Rapid diagnosis on uncultivated nuclei:	ome 21 (NC21) Chromosomes 13, 18, 21, X, Y (NCKIT)				
- Microdeletion screening - Please specify:					
For any additional request, it is highly recommended to take a third tub					
FŒTAL BIOCHEMISTRY	e decoraing to the term of pregnancy.				
Alpha fetoprotein (αFP) (AFPLA) Acetylcholinesterase Other:					
Seroconversion: YES NO If yes, o	lease specify:late of pregnancy at the time of seroconversion:				
• Desired pathogens: ☐ Cytomegalovirus (CMVLA) ☐ Rubella (RUBLA) ☐ Varicella Zoster (VZVLA)	☐ Toxoplasmosis (TOXLA) ☐ HSV1/HSV2 (HSVLA) ☐ Other:				
	NO If yes, return date: VES NO				
Please supply the maternal serology results and the date of seroconversion	n as well as the latest ultrasound report				
☐ MOLECULAR GENETICS					
For all test requests, please supply 5 mL EDTA maternal blood sample and 5 mL EDTA paternal blood sample.					
Uniparental disomy (DUPRE) Which chromosomes 7, 14, 15:	qPCR: contact us (QPRE) (attach the R66-INTGB information form)				
☐ Mono-/di-zygotic (twins) (zygo)	Sanger: contact us (SEPRE)				
☐ Cystic fibrosis (CFTR, screening for most common	(attach the R66-INTGB information form)				
mutations) (MUCOL): (see specific file available on	Fragile X syndrome (XFRAP)				
www.eurofins-biomnis.com)	Exome (specific request form required, please see				
☐ Prader-Willi syndrome (SNRPL)	www.eurofins-biomnis.com)				

Documents