

- Sample date:

Weeks of amenorrhea:

- Sample type

☐ Amniotic fluid (CLA ou CHN)

☐ Chronic villi sampling (CVS)

☐ Products of conception (PoC)

Tube 1 ▶ Volume: mL

▶ Appearance: ☐ clear ☐ brown ☐ bloody

☐ Foetal blood (FB)

Tube 2 ▶ Volume: mL

▶ Appearance: ☐ clear ☐ brown ☐ bloody

☐ DNA extracted from:
(specify the nature of the sample)

In cases of twin pregnancy, please indicate the number of foetus sampled:

TEST REQUEST

☐ FETAL STANDARD KARYOTYPE

☐ FETAL DNA MICROARRAY (SNP ARRAY) (SNPRE) - Please supply 5 mL EDTA maternal blood sample

☐ Risk of trisomy 21 by MSM greater than or equal to 1/50 - The report must be supplied

☐ 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 karyotype - The report must be supplied

☐ Maternal age > 38 years [without trisomy 21 screening (MSM or NIPT)] after 18 weeks' gestation

☐ Personal request:

☐ Other:

☐ IN SITU HYBRIDISATION (FISH)

- Rapid diagnosis on uncultivated nuclei: ☐ Chromosome 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 tube according to the term of pregnancy.

☐ FETAL BIOCHEMISTRY

☐ Alpha fetoprotein (αFP) (AFPLA) ☐ Acetylcholinesterase (AChE) (ACOLA) ☐ Digestive enzymes

☐ Other:

FETAL INFECTION PROFILE

• Indication:

Ultrasound signs : ☐ YES ☐ NO If yes, please specify:

Seroconversion: ☐ YES ☐ NO If yes, date of pregnancy at the time of seroconversion:

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• Desired pathogens:

☐ Cytomegalovirus (CMVLA) ☐ Parvovirus B19 (PARLA) ☐ Toxoplasmosis (TOXLA) ☐ HSV1/HSV2 (HSVLA)

☐ Rubella (RUBLA) ☐ Varicella Zoster (VZVLA) ☐ Other:

☐ Zika (ZIKLA) ▶ Stay in endemic areas: ☐ YES ☐ NO If yes, return date:

▶ Spouse presenting a positive sample for Zika: ☐ YES ☐ NO

Please supply the maternal serology results and the date of seroconversion 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)
(attach the R66-INTGB information form)

☐ Cystic fibrosis (CFTR, screening for most common mutations) (MUCOL): (see specific file available on www.eurofins-biomnis.com)

☐ Fragile X syndrome (XFRAP)

☐ Prader-Willi syndrome (SNRPL)

☐ Exome (specific request form required, please see www.eurofins-biomnis.com)

☐ Other:

Documents

- Test request form indicating which tests to be performed