

# TEST REQUEST AND PATIENT CONSENT FORM

## PATIENT DETAILS

Name \_\_\_\_\_ Surname \_\_\_\_\_  
 Date of birth \_\_\_\_\_ Place of birth \_\_\_\_\_  
 Address \_\_\_\_\_  
 \_\_\_\_\_  
 Date of blood withdraw \_\_\_\_\_  
 Gynecologist name \_\_\_\_\_  
 Address \_\_\_\_\_  
 \_\_\_\_\_  
 Phone No. \_\_\_\_\_  
 E-mail \_\_\_\_\_

Protocol no. (internal use only): \_\_\_\_\_

## ORDERING LABORATORY / CLINICIAN

Clinic Name \_\_\_\_\_  
 Address \_\_\_\_\_  
 \_\_\_\_\_  
 Contact No. \_\_\_\_\_

## PATIENT MEDICAL HISTORY

\_\_\_\_\_  
 \_\_\_\_\_

## PREGNANCY HISTORY

Patient current weight \_\_\_\_\_ Kg. Patient height \_\_\_\_\_  
 Gestational age at draw \_\_\_\_\_ weeks + days \_\_\_\_\_  
 Gestational age calculated by:  
 Ultrasound  last menstrual period  IVF treatment  
**Twin pregnancy?**  Yes  NO Monochorial  Dichorial   
**IVF Pregnancy?**  Yes  NO  
 Homologous pregnancy  Heterologous Pregnancy  
 Embryo donation  Eggs donation  Sperm donation

## INDICATION FOR TESTING

Advanced maternal age  Advanced paternal age  
 Partner carrier of a genetic disorder:  Male  Female  
 Specify disorder \_\_\_\_\_  
 Gene and mutation \_\_\_\_\_  
 Parental anxiety (low-risk) \_\_\_\_\_  
 Abnormal ultrasound (describe) \_\_\_\_\_  
 Other indication \_\_\_\_\_  None

## TEST REQUESTED, please tick all that apply

- PrenatalSAFE® 3 test (for chromosomes 21, 18,13 only)  
 PrenatalSAFE® 5 test (for chromosomes 21, 18, 13, X, Y)  
 PrenatalSAFE® Plus test (for chromosome 21, 18, 13, X, Y) +  Panel 6 Microdeletion  Trisomies 9 and 16 option  
 PrenatalSAFE® Karyo test (genome-wide NIPT that provides karyotype-level insight)  
 PrenatalSAFE® Karyo Plus test (genome-wide NIPT that provides karyotype-level insight + Panel 9 Microdeletions)  
 GeneSAFE™ Inherited  GeneSAFE™ *de novo*  GeneSAFE™ Complete (Inherited + *de novo*)  
 PrenatalSAFE® COMPLETE (PrenatalSAFE® Karyo + GeneSAFE™ Complete)  
 PrenatalSAFE® COMPLETE Plus (PrenatalSAFE® Karyo Plus + GeneSAFE™ Complete)  
 Do you wish to know the fetal gender?  Yes  NO Is it a redraw?  Yes  NO

## PATIENT INFORMED CONSENT

I consent to the test(s) I have chosen and confirm that I have been informed about the purpose, scope, and limitations of the test by my healthcare provider. I understand that the test is a screen for selected abnormalities; that the result should be reviewed by my healthcare provider. I have had the opportunity to ask questions and understand that I can request further information or appropriate counselling. I consent to the use of the leftover specimen and health information as described in the Test Information Document.

\_\_\_\_\_  
 Patient Name

\_\_\_\_\_  
 Signature

\_\_\_\_\_  
 Date

## REPORTING PREFERENCES

PHYSICIAN / LABORATORY  
 E-mail \_\_\_\_\_  
 On-Line  Post  
 In order to activate the on-line reporting option, you need to provide us  
 Username \_\_\_\_\_ Password \_\_\_\_\_  
 \_\_\_\_\_  
 Signature Date