\GENOM. **PATIENT DETAILS** Protocol no. (internal use only): _____ PrenatalSat Name Surname_____ Date of birth_____ Place of birth____ ORDERING LABORATORY / CLINICIAN VAT no. ______ Address:_____ asiakasneuvonta@vita.fi Phone no.: __ hanna.sipi@vita.fi Date of blood withdraw ____ Gynecologist name:_____ PATIENT MEDICAL HISTORY Address:_____ Phone no.: _____ E-mail: _____ **PREGNANCY HISTORY** INDICATION FOR TESTING ☐ Advanced maternal age; ☐ Advanced paternal age; Patient current weight Kg _____ Patient height_ ☐ Parental anxiety (low-risk) Gestational age at draw _____ + days ____ ☐ Abnormal ultrasound (describe): _ Gestational age calculated by: ☐ Previous pregnancy with aneuploidy; ☐ Ultrasound; ☐ last menstrual period; ☐ IVF treatment ☐ Abnormal maternal serum screening test; Twin pregnancy? ☐ Yes; ☐ NO Monochorial ☐ Bichorial ☐ ☐ Partner carrier of a genetic disorder: ☐ Male ☐ Female IVF Pregnancy? ☐ Yes; ☐ NO ☐ Specify disorder: ______ ☐ Homologous pregnancy; ☐ Heterologous Pregnancy ☐ Specify gene and mutation: ☐ Embryo donation; ☐ Eggs donation; ☐ Sperm donation ☐ Other indication ___ ☐ None TYPE OF TEST ✓PrenatalSAFE® Plus test (for chr. 21, 18, 13, X, Y) * (can not be requested for bichorial pregnancy) + Panel 6 Microdeletion* *This option includes the following syndromes: 22q11 deletion (DiGeorge); 15q11 deletion (Angelman/ Prader-Willi); 1p36 deletion, 4p- (Wolf-Hirschhorn); 5p- (Cri-du-chat) Do you wish to know the fetal gender? 🔲 Yes; 🔲 NO Is it a redraw? ☐ Yes; ☐ NO REPORTING PREFERENCES **BILLING** PHYSICIAN / LABORATORY Sending facility

Signature

Mod.PR 11.H3/ENG Rev. 14



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= ISO 9001 =

PATIENT CONSENT STATEMENT:

By signing this form, I, the patient having the testing performed, acknowledge that:

- (i) I have received and read or have had read to me the above informed consent information about the PrenatalSafe® Non-Invasive Prenatal Test (NIPT) in its entirety and realize I may retain a copy for my records;.
- (ii) I have had the opportunity to ask questions of my health care provider regarding this test, including the reliability of test results, the risks, and the alternatives prior to my informed consent.;
- (iii) I have discussed with the healthcare provider ordering this test the reliability of positive or negative test results and the level of certainty that a positive test result for a given disease or condition serves as a predictor of that disease or condition;
- (iv) I have been informed about the availability and importance of genetic counseling and have been provided with information identifying an appropriate healthcare provider from whom I might obtain such counseling;
- (v) I consent to the use of the leftover specimen and health information as described in the Patient Informed Consent;
- (vi) I consent to having this test performed and I will discuss the results and appropriate medical management with my healthcare provider.

Date:	
Signature of Patient	
Printed Name	
The physician who has collected the consent (first a	and last name)
Tel. E-Mail	
Signature and stamp of the physician who has colle	cted the consent:

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