



## VANADIS® NIPT and PRE-ECLAMPSIA (PIGF) Requisition Form

Please complete every field clearly. Missing information may result in a delay of sample processing.

PATIENT INFORMATION		PATIENT SAMPLE AND CL	INICAL INDIC	CATION		
	MM/ DD /YYYY	Collection Date: MM/DD/	Y Was th	is sample collec	ted in NY St	ate: ○ Yes ○ No
First Name	Date of Birth	ICD-10 code(s) required:  AMA Primigravida:				
Last Name	Patient ID Number					
Patient's Street Address		<ul><li>Abnormal ultrasonic find</li><li>Abnormal serum screeni</li></ul>	•			
	<ul><li>Abnormal finding unspect</li><li>History of recurrent pregr</li></ul>		6 20			
City / Town	Maternal care for (suspe	-		ity in fetus - (	035.1XX0	
	Other ICD-10:					
Country Patient's Preferred Phone	TEST MENU (check all tha	t apply)				
	<b>NIPT Testing Options: P</b>	ease Fill Out	Section 1 Belo	w if selecte	d	
Patient's Email (Used for communication of insurance coverage	O VAN110 Vanadis NIPT (Trisomies 21, 18, 13) + Fetal Sex Determination					
ORDERING ACCOUNT	○ VAN100 Vanadis NIPT Excluding Fetal Sex Determination  Vanadis - Whole blood in two 10ml Speckled Top Cell Free BCT Tubes required					
		•		•		
Provider's First and Last Name		PIGF PRE-ECLAMPSIA To ○ PGF100 PIGF for pre-			it Section 2	Below if selected
PerkinElmer Account Number NPI	PIGF - Pre-Eclampsia W			p tube requir	red	
		Section 1: Required for NI	PT Testing O	nly (select all t	hat apply)	
Clinic/Hospital/Laboratory Name				•	Yes	No
		Abnormal ultrasound finding			0	0
Primary Contact Email Primary C	Contact Phone	Positive serum screen in cu Patient or father of pregnanc		•	0	0
		chromosome abnormality	y with a fairing i	listory or	0	0
Account Street Address		Patient or father of pregnar	cy with history	of recurrent		
		pregnancy loss			0	0
City / Town	State Postal Code	Previous Pregnancy – Triso Previous Pregnancy – Triso	•		0	0
		Previous Pregnancy – Trisc	-		0	0
Country Account Fax		Previous pregnancy with ar		e abnormality	0	0
PREGNANCY DETAILS	If yes to any items, please	specify:				
EDC: MM/ DD / YYYY Dating by: OLMP O Ultra Maternal Weight: Olbs Okg Maternal Heigh	isound nt: O ft/in O cm	Section 2: Required for Pl	GF and Pre-E	clampsia Testi	ng Only	
	IL © 10111 © CITI	Mean Arterial Pressure (M	AP) Details	l Increas	ses detectio	n rate by > 19%
Fetus Details	O.D. L. d. d.	Blood Pressure Measure Da	•	•	300 40100110	
○ Singleton ○ Twins If Twins: ○ Monochorionic	O Dicnorionic	blood Flessule Measule Da		Arm	R	ght Arm
Conception Details	-/N-t 0 IV/F 0 1001	Blood Pressure (mm/HG)	Systolic	Diastolic	Systolic	Diastolic
Conception Method (select all that apply): Spontaneous  PGS/PGT-A Ovulation drugs Other assiste	ed reproduction	First Reading			,	
If IVF pregnancy, egg source: O Self O Donor	,a reproduction	Second Reading				
Age of donor/self at time of extraction:						
Maternal History	Scan Details		! Increa	ses detectio	on rate by > 49%	
Yes Insulin Dependent Diabetes Mellitus (IDDM)	No ○	Date of Ultrasound: MM/ DCRL (mm) - for twins, use la		measurements		
Insulin Dependent Diabetes Mellitus (IDDM)  Diabetes Type II	0			Left	Right	Mean
If yes, is she using insulin?	0	Uterine Artery Pulsatility Ind [Acceptable Range: 0.4-4]	ex (UTPI)	Leit	rtigitt	Weari
Smoking  Is this patient's first pregnancy?  If no pumber of pregnancy?	0	Maternal History		1		'
If no, number of pregnancies after 24 weeks:  ETHNICITY (check all that apply):   Caucasian	n - Non Hispanic			Yes	No	
O African American O French Ca	· ·	Chronic Hypertension				
Asian O Hispanic		Anti-phospholipid antibody syndrome or Systemic lupus erythematosus				
Olndian subcontinent O Jewish - A		Mother of Patient had Pre-eclampsia				
<ul><li>○ Chinese</li><li>○ Filipino</li><li>○ Middle East</li></ul>		Patient or father of pregnancy of recurrent pregnancy loss		0	0	
<ul><li>○ Filipino</li><li>○ Middle East</li><li>○ Japanese</li><li>○ Native Am</li></ul>		Previous Pregnancy - Pre-e		0	0	
O Korean O Pacific Isla		If yes: Delivery date: MM/				
Other Asian descent Other (specify): Gestational age: wks. Baby's bir			birth weight:	Ibs/oz	S.	
CONFIRMATION OF INFORMED CONSENT AND MEDICAL NECESSITY						
The undersigned person (or representative thereof) ensures her	she is a licensed medical profes	sional authorized to order genetic	testing and cor	nfirms that the pa	tient has give	n appropriate

The undersigned person (or representative thereof) ensures he/she is a licensed medical professional authorized to order genetic testing and confirms that the patient has given appropriate informed consent for the testing ordered, including a discussion of the benefits and limitations. I confirm that testing is medically necessary and that test results may impact medical management for the patient. Furthermore, all information on this TRF is true to the best of my knowledge. My signature applies to the informed consent and/or attached letter of medical necessity.

for the patient. I distribute, an information on the first	, and to the beet of my fallowledge. My digitation	applied to the informed deficent analysis attached let	
Signature	Date	FOR	INTER
authorized medical professional		Date Rec'd	

FOR INTERNAL USE ONLY

Date Rec'd Rec'd

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## VANADIS® NIPT and PRE-ECLAMPSIA (PIGF) Requisition Form (page 2)

PAYMENT INFORMATION							
■ INSTITUTIONAL BILLING (Please ensure all fields in Ordering Account section on page 1 are complete)							
Billing Account Name	Billing Account Number						
Contact Name	Contact Number						
□ PATIENT BILLING  ○ Credit Card (Please fill out all information):  ○ Check: \$	Amount Enclosed (Please make checks payable to: PerkinElmer Genetics, Inc.)						
Credit Card Number	CVV Card Exp. Date Billing Zip Code						
Cardholder Printed Name as Appears on Card	Cardholder Signature						
■ INSURANCE BILLING (Include a copy of both sides of insurance card)							
Insurance Carrier and ID	Prior Authorization # (attach confirmation) Policy Holder Relationship to Patient:						
Policy Holder Name	Policy Holder DOB Self O Parent O Spouse O Other:						
ADDITIONAL INFORMATION AND INFORMED CONSENT							
SAMPLE AND DATA RETENTION							
Pursuant to laboratory best practices, your DNA sample will be retained by PerkinElmer for a minimum of two years and then destroyed. Additionally, your PHI, the data from the Tests (including those performed before any withdrawal of consent) and the related reports will be retained by PerkinElmer indefinitely, unless otherwise noted. In some instances, it may be beneficial to you for PerkinElmer to retain your sample for a longer period of time in order to conduct additional testing, and PerkinElmer will do so with appropriate documentation from you or your HCP.							
PerkinElmer is requesting consent to keep your anonymized sample and data indefinitely for ongoing test development, scientific research, and/or other activities. This consent is optional, and the Test will be performed whether or not you provide consent to the following:							
If consent is given for Sample retention, PerkinElmer will anonymize and retain your Sample indefinitely for internal quality control, test validation, assay development and improvement. By allowing PerkinElmer to retain your Sample, you understand and agree that you give up any property rights you may have in the Sample and are donating it to PerkinElmer Genetics, Inc. If you withdraw your consent, no additional tests or anonymization will be carried out on your Sample; no results will be reported and your sample, reports and data that have not been anonymized will be destroyed.							
If consent is given for data retention, PerkinElmer will anonymize your data and retain the anonymized data and related anonymized reports from your Tests indefinitely for internal statistical, quality analysis, research, scientific and technical development, and market research.							
PerkinElmer is requesting consent to keep your anonymized sample indefinitely for ongoing test development, scientific research, and/or other activities.  O Check here if you would like to opt out of anonymized sample retention. Note, if not checked, this is interpreted as "consent given".							
PerkinElmer is requesting consent to keep your anonymized data indefinitely for ongoing test development, scientific research, and/or other activities.  O Check here if you would like to opt out of anonymized sample retention. Note, if not checked, this is interpreted as "consent given".							
PATIENT INFORMED CONSENT							
☐ By checking this box, I am requesting that PerkinElmer contact me if my estimate PIGF - Pre-Eclampsia. I acknowledge that PerkinElmer will make three attempts t							
I certify that the insurance information that I have provided is accurate, complete and current and that no other coverage or insurance exists. I hereby authorize PerkinElmer Genetics, Inc. ("PerkinElmer") to bill my designated insurance carrier(s) and share health information as needed for the purposes of billing and reimbursement, and I request that payment of authorized benefits be made on my behalf to PerkinElmer for any services furnished the patient listed above by PerkinElmer. If any insurance benefits are remitted to me for services performed by PerkinElmer for the patient, I will forward said benefits to PerkinElmer. I authorize PerkinElmer to file an appeal on my behalf for any denial of payment and/or adverse benefit determination related to services and care provided. I agree to pay all charges for services provided by PerkinElmer to the patient which are not covered by my health insurance plan or which I am responsible for payment under my health insurance plan. Furthermore, I grant PerkinElmer permission to share health information with my insurance as needed for the purposes of billing and reimbursement.							
By signing below I attest that I have read and understood the information in this Patient Informed Consent and I have had my questions about the testing to be performed answered by my healthcare provider. I attest that all information I have provided is accurate, and I consent to PerkinElmer performing the testing ordered by my healthcare provider.							
Patient Signature Patient Na	me Date						