

Pathology and Laboratory Medicine Clinic Building, K6, Core Lab, E-655 2799 W. Grand Bvd. Detroit, MI 48202 855.916.4DNA (4362)

## PRENATAL CYTOGENOMICS REQUISITION

CENTER FOR PRECISION DIAGNOSTICS

Required Patient Information	Ordering Physician Information			
Name: Gender: M F	Name:			
MRN:DDB:MM/DD/YYYY	Address:			
ICD10 Code(s):	City: State: Zip:			
ICD-10 Codes are required for billing. When ordering tests for which reimbursement will be sought, order only those tests that are medically necessary for the diagnosis and treatment of the patient.	Phone: Fax:			
Billing & Collection Information	NPI:			
Patient Demographic/Billing/Insurance Form is required to be submitted with this for	m. Most genetic testing requires insurance prior authorization.			
Due to high insurance deductibles and member policy benefits, patients may elect to	self-pay. Call for more information (855.916.4362)			
Bill Client or Institution Client Name:	Client Code/Number:			
Bill Insurance Prior authorization or reference number:				
Patient Self-Pay Call for pricing and payment options Toll Free: 8	355.916.4362			
Patient status at time of collection:  Inpatient  Outpatient				
roviders are responsible to obtain informed consent, as required by Michigan law, for predictive or pre-symptomati equisition.				
Specimen/Source				
Maternal peripheral blood (required for MCC studies and Toxoplasma Serology, 5m				
Amniotic fluid (15-20mL of fluid in 2-3 aliquots) Fluid color:				
Chorionic villus (CVS)				
Products of Conception (POC) (send in sterile media, Ringer's lactate or saline) Tiss	ue source:			
Extracted DNA – Source:	(provide CLIA certificate of lab that performed the DNA extraction)			
ndication for Testing				
Maternal Age:	☐ Family history (specify) :			
Abnormality on ultrasound (specify):	☐ Other:			
☐ NIPT positive for: ☐+21 ☐+18 ☐+13 ☐ Other				
Pregnancy History				
Gestational age (GA): weeks Last Menstrual Period (LMP)	:/			
Gravidity: Parity: Abortus:				
Biparietal Diameter or other: mm Date ultrasound performed:				
Prenatal Testing Options				
	Some testing includes pathologist interpretation at a separate, additional charge.  us (CMV) PCR			
(with reflex to AChE and Fetal Hemoglobin if AFP MoM ≥2.0) on amniotic flu	, , , , , , , , , , , , , , , , , , , ,			
1 Chromosome analysis (Amniotic Fluid/CVS: 88235, 88267, 88280, 88285, 88291; PO	C: 88233, 88262, 88291)			
FISH Prenatal Aneuploidy Screen (88271x5, 88274x2) Note: requires additional 5mL	of sample			
Microarray (SNP Array) (81229)				
☐ Direct ☐ Cultured cells if GA ≤17 weeks				
Maternal Cell Contamination (MCC) Studies (81265)				
Additional Testing	Send Additional Report To			
	Name:			

Address: Phone #:

Fax #:

## HENRY FORD HEALTH

## **Consent for Genetic Testing**

Place patient label here or fill out information below:	
Patient Name:	
Date of Birth:	
MRN:	

Office Use Only					
Ordering Provider Information (Last, First)	Genetic Testing Requested for:				
Name:					
Phone:	(name of medical condition)				
Sample Type:  Amniotic fluid  Blood  Cheek swab  Chorionic villus sample (CVS)  Skin  Tissue block  Other	The purpose is (check all that apply):  ☐ Carrier status ☐ Diagnostic ☐ Predictive ☐ Prenatal ☐ Pre-symptomatic ☐ Screening ☐ Other				

## I understand and agree to the following:

- This form goes with an information booklet that has more information on genetic tests. I can find the booklet online at <u>What Michigan Patients Need to Know Before Getting a Genetic Test</u> or a written copy can be provided.
- 2. This genetic test has been explained to me. I understand why I am having this test done.
- 3. I understand what this genetic test may or may not be able to find.
- 4. I was able to talk to my doctor or other health care provider about the benefits and the risks of this test. I know that some genetic tests can involve health, mental health, or insurance issues for me or my family.
- 5. I understand what the results may mean and how I will get them.
- 6. I understand that genetic tests can sometimes find other results that have nothing to do with the original reason for the test. These are called secondary findings. I talked to my doctor or health care provider and I understand that I can decide if I want secondary findings shared with me.
- 7. I was told who may access my sample. I understand that any leftover sample may be kept by the laboratory and used for quality checks.
- 8. I was told who may see my test results. These results will be part of my health record.
- 9. I was able to talk to my doctor and have my questions answered about this test.
- 10. I was given a copy of this form for my records.

I have read this form, or it was read to me. I understand and agree to what it says. I agree to have a sample taken for genetic testing for the condition(s) listed above. If the signer is not the patient, the signer confirms that they are the patient's legally authorized representative.

Person signing form (circle one):	Self	Parent(s)	Legal Guardian	Durable Power of Attorney for Health Care		
Signature of Patient or Authorized Designee				Time	Date	
Signature of Physician or person e	xplaini	ng informat	ion	Time	Date	