

APPLY LABELS TO PATIENT SPECIMENS ONLY.

To find the nearest patient service center, visit Labcorp.com or call 888-Labcorp (888-522-2677).

Patient's Legal Name (Last, First, MI)		Sex	Date of Birth MO DAY YR	Collection Time AM PM	Fasting <input type="checkbox"/> Yes <input type="checkbox"/> No	Collection Date MO DAY YR	Urine hrs/vol hrs ____ vol ____
NPI	Physician's ID #	Patient's ID #		Hospital Patient Status: <input type="checkbox"/> In-Patient <input type="checkbox"/> Out-Patient <input type="checkbox"/> Non-Patient			
Physician's Name (Last, First)		Physician/Authorized Signature X _____		Genetic Counselor (Last, First)			
Diagnosis/Signs/Symptoms in ICD-CM format in effect at Date of Service <b>Highest Specificity REQUIRED</b>				Patient's Address		Phone	
				City		State ZIP	
				Email Address			
				Name of Policy Holder (if different from patient)			
				Address of Policy Holder		APT #	
				City		State ZIP	
I hereby authorize the release of medical information related to the service described herein and authorize payment directly to Labcorp. I agree to assume responsibility for payment of charges for laboratory services that are not covered by my healthcare insurer. X Patient's Signature _____ Date _____							
<b>MEDICARE ADVANCE BENEFICIARY NOTICE OF NON-COVERAGE (ABN)</b> Refer to policies published by your Medicare Administrative Contractor (MAC), CMS, or Labcorp.com/MedicareMedicalNecessity when ordering tests that are subject to ABN guidelines.							
PRIMARY BILLING PARTY				SECONDARY BILLING PARTY			
Insurance Carrier *		Insurance Carrier *		Insurance Carrier *		Insurance Carrier *	
ID #		ID #		ID #		ID #	
Group #		Group #		Group #		Group #	
Insurance Address		Insurance Address		Insurance Address		Insurance Address	
Name of Insured Person		Name of Insured Person		Name of Insured Person		Name of Insured Person	
Relationship to Patient		Relationship to Patient		Relationship to Patient		Relationship to Patient	
Employer Name		Employer Name		Employer Name		Employer Name	
*If Medicaid State		Physician's Provider #		Workers Comp <input type="checkbox"/> Yes <input type="checkbox"/> No			
TEST #				OTHER TESTS / INDIVIDUAL PROFILE COMPONENTS TEST NAMES			

### INFORMED CONSENT

I have obtained informed consent for the above ordered genetic test(s). (Required)

Physician's Signature \_\_\_\_\_

Please indicate the diagnostic center to which you want screen positive results reported (NY State only)

Additional tests available. Call Genetics Services for info. 1-800-345-GENE

**REQUIRED INFORMATION**

GA \_\_\_\_ wks \_\_\_\_ days on date \_\_\_\_/\_\_\_\_/\_\_\_\_

Patient Weight \_\_\_\_ lbs

# of Fetuses  1  2  Other \_\_\_\_

Patient Race  Cauc  Hispanic  Black  
 Asian  Amer Indian  Other

Yes  No Is patient an insulin dependent diabetic?  
 Yes  No Egg donor:  Self  Non-self  
Age of donor at egg retrieval: \_\_\_\_ years

**CLINICAL HISTORY**

Yes  No Prior Down Syndrome/ONTD Screen in Current Pregnancy? If yes, prior test was:  
 in 1st Tri  in 2nd Tri  elevated msAFP

Yes  No Family history of NTD?

Yes  No Previous pregnancy with Down Syndrome?

Yes  No Parental cytogenetics following abnormal prenatal results.

Yes  No Parental balanced Robertsonian Translocation with increased risk of Trisomy.

Yes  No Other Indications: \_\_\_\_\_

### CYTOGENETICS

**Amniotic Fluid (specify GA below)**

511580  Chromosome & AFP/AChE/HbF

052040  Chromosome Analysis

002428  AFP, Amniotic fluid

510305  AFP, AChE with reflex to HbF

511894  FISH, InSight Aneuploidy Evaluation

052104  Chromosome rfx SNP Microarray (Reveal®)

511966  FISH, InSight rfx to Chromosome or Microarray

511590  Chrom. 5 Count + Reveal® SNP Microarray

510100  SNP Microarray - Prenatal (Reveal®)

510200  SNP Microarray (Direct) - Prenatal (Reveal®)

**Chorionic Villi (specify GA below)**

510988  Chromosome Analysis

510960  FISH, InSight Aneuploidy Evaluation

511033  Chromosome rfx SNP Microarray (Reveal®)

511625  FISH, InSight rfx to Chromosome or Microarray

511555  Chrom. 5 cell + Reveal® SNP Microarray

510100  SNP Microarray - Prenatal (Reveal®)

510200  SNP Microarray (Direct) - Prenatal (Reveal®)

**POC / Tissue / Other (specify GA below)**

052052  Chromosome Analysis

510110  SNP Microarray - POC Tissue (Reveal®)

511997  SNP Microarray - POC FFPE (Reveal®)

052065  Chromosome rfx Reveal® SNP Microarray

**Chromosome, Blood**

511035  Chromosome, Blood

510770  FISH Microdeletion (Specify)

**Fetal Sex:**  Male  Female  Unknown

**By:**  U/S  NIPS  PGT

**Gestational age:** \_\_\_\_\_

Hold cells

511402  Maternal cell contamination (mat blood required)

Clin Info, Fam Hx, Other \_\_\_\_\_

Abnl NIPS: Specify \_\_\_\_\_

### PREGNANCY COMPLICATIONS

005199  Prothrombin Time (PT)

005207  PTT, Activated (APTT)

161802  Anticardiolipin Ab, IgG, IgM

015594  Antithrombin Deficiency Profile

001610  Fibrinogen Activity

511162  Factor II (Prothrombin)

511154  Factor V Leiden

706994  Homocysteine

117892  Lupus Anticoagulant w Reflex

511238  MTHFR

283655  Protein C Deficiency Profile

117754  Protein S Deficiency Profile

504295  ReproSURE™ (Ovarian Reserve Profile)  
See Reverse GEL

502051  Recurrent Miscarriage

504808  Von Willebrand Reflexive

**MATERNAL SERUM SCREENING W/O NT**

017200  Serum Integrated 1 (10w0d-14w0d)

017270  Serum Integrated 2 (15w0d-21w6d)

017319  AFP Tetra (15w0d-21w6d)

010801  msAFP (15w0d-23w6d) (optimal 16w-18w6d)

GA \_\_\_\_ wks \_\_\_\_ days on date \_\_\_\_/\_\_\_\_/\_\_\_\_

By  LMP  EDC/EDD  U/S

**MATERNAL SERUM SCREENING WITH NT**

017500  1st Trimester Screen (10w0d-14w0d)

017700  Sequential Part 1 (10w0d-14w0d)

017750\*  Sequential Part 2 (15w0d-21w6d)

017100  Integrated Part 1 (10w0d-14w0d)

017170\*  Integrated Part 2 (15w0d-21w6d)

CRL date \_\_\_\_/\_\_\_\_/\_\_\_\_ CRL \_\_\_\_ mm (45.0-84.0)

Twin B, if applicable CRL \_\_\_\_ mm (45.0-84.0)

NT \_\_\_\_ mm Chorionicity:  Mono  DI  Unknown

NT \_\_\_\_ mm

Sonographer Name\*: Last \_\_\_\_\_ First \_\_\_\_\_

Sonographer ID #: \_\_\_\_\_

Credentialed by:  NTQR  FMF  Other

Reading MD ID #: \_\_\_\_\_

Site ID#: \_\_\_\_\_

Nasal Bone:  Not Evaluated  Present  Absent

NB Twin B:  Not Evaluated  Present  Absent

**MATERNAL PLASMA SCREENING: NONINVASIVE PRENATAL SCREENING (NIPS)**

451927  MaternIT® 21 PLUS (9w+)

451931  MaternIT® 21 PLUS w/ ESS\* (9w+)

451934  MaternIT® 21 PLUS w/ SCA\*\* (9w+)

451937  MaternIT® 21 PLUS w/ ESS & SCA\*\* (9w+)

451941  MaternIT® Genome (9w+)

451951  MaternIT® 21 PLUS No Gender (9w+)

452104  GENOME-Flex\*\* (Add On)

452114  GENOME-Flex\*\* (Add On) Redraw

\* ESS = chr 16, chr 22, and select microdeletions

\*\* SCA = sex chromosome aneuploidies; singleton only

GA \_\_\_\_ wks \_\_\_\_ days on date \_\_\_\_/\_\_\_\_/\_\_\_\_

Indication for Noninvasive Prenatal Screening (NIPS)

AMA  Positive maternal serum screening test

Previous pregnancy with aneuploidy  Family history of chromosomal abnormality

Ultrasound findings (Check all that apply):

cystic hygroma  IUGR  heart defect  CNS abnormality

Other: \_\_\_\_\_

**CARRIER SCREENING**

481758  Inheritest® CF/SMA panel

481776  Inheritest® Core Panel

481797  Inheritest® 14-gene Panel

481816  Inheritest® High Frequency Panel

481855  Inheritest® 100 PLUS Panel

481874  Inheritest® 300 PLUS Panel

481893  Inheritest® 500 PLUS Panel

482370  GeneSeq PLUS  VUS opt out

481025  Cystic Fibrosis (CF), 97 Variants

482632  Cystic Fibrosis (CF), Full-gene Carrier Screen

481684  Fragile X Syndrome, Carrier

481630  Spinal Muscular Atrophy (SMA)

511172  α - Thalassemia, DNA Analysis

252823  β - Thalassemia: HBB (Full Gene Sequencing)

121690  Hgb Fractionation Cascade

By providing the reproductive partner's information, you, the ordering provider, confirm that you have obtained from the patient and reproductive partner all required consents and/or authorizations necessary for the use and disclosure of protected health information, including test results, between the patient and reproductive partner. A separate requisition is required for each partner.

**Required: Gene(s)** \_\_\_\_\_

Partner's name: \_\_\_\_\_ Partner's DOB: \_\_\_\_\_

482595  Partner Reflex to GeneSeq (male partner only)

## TEST COMBINATION/PANEL POLICY

Labcorp's policy is to provide physicians, in each instance, with the flexibility to choose appropriate tests to assure that the convenience of ordering test combinations/panels does not distance physicians who wish to order a test combination/profile from making deliberate decisions regarding which tests are truly medically necessary. All the tests offered in test combinations/panels may be ordered individually using the Labcorp® request form. Labcorp encourages clients to contact their local Labcorp representative or Labcorp location if the testing configurations shown here do not meet individual needs for any reason, or if some other combination of procedures is desired.

In an effort to keep our clients fully informed of the content, charges and coding of its test combinations/panels when billed to Medicare, we periodically send notices concerning customized test combinations/panels, as well as information regarding patient fees for all Labcorp services. We also welcome the opportunity to provide, on request, additional information in connection with our testing services and the manner in which they are billed to physicians, health care plans, and patients.

For CPT codes please contact the CPT coding department at telephone number 800-222-7566 ext 6-8400 or [www.labcorp.com](http://www.labcorp.com). Please note, correct coding often varies from one carrier to another. Consequently, the codes provided by Labcorp are intended as general guidelines and should not be used without confirming with the appropriate payor that their use is appropriate in each case. All laboratory procedures will be billed to third-party carriers (including Medicare and Medicaid) at fees billed to patients and in accordance with the specific CPT coding required by the carrier. Microbiology CPT code(s) for additional procedures such as susceptibility testing, identification, serotyping, etc. will be billed in addition to the primary codes when appropriate. Labcorp will process the specimen for a microbiology test based on source.