

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

Patient's Legal Name (Last, First, MI)				Sex M <input type="checkbox"/> F <input type="checkbox"/>	Date of Birth MO DAY YR	Collection Time AM <input type="checkbox"/> PM <input type="checkbox"/>	Fasting Yes <input type="checkbox"/> No <input type="checkbox"/>	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 _____							
Diagnosis/Signs/Symptoms in ICD-CM format in effect at Date of Service Highest Specificity REQUIRED									
PRIMARY BILLING PARTY					SECONDARY BILLING PARTY				
Insurance Carrier *					Insurance Carrier *				
ID #					ID #				
Group #					Group #				
Insurance Address					Insurance Address				
Name of Insured Person					Name of Insured Person				
Relationship to Patient					Relationship to Patient				
Employer Name					Employer Name				
*If Medicaid State					Physician's Provider #		Workers Comp <input type="checkbox"/> Yes <input type="checkbox"/> No		

PATIENT	Patient's Address		Phone
	City	State	ZIP
	Name of Policy Holder (if different from patient)		
	Address of Policy Holder		APT #
RESP. PARTY	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 _____			

MEDICARE ADVANCE BENEFICIARY NOTICE OF NONCOVERAGE (ABN)
Refer to policies published by your Medicare Administrative Contractor (MAC), CMS, or www.LabCorp.com/MedicareMedicalNecessity when ordering tests that are subject to ABN guidelines.

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

MATERNAL PLASMA SCREENING: NON INVASIVE PRENATAL TESTING (NIPT)

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 chromosomes aneuploidies

GA ____ wks ____ days on date ____/____/____

By LMP EDC/EDD U/S

Indication for Non Invasive Prenatal Testing (NIPT)

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: _____

MATERNAL SERUM SCREENING W/O NT

017200	<input type="checkbox"/>	Serum Integrated 1 (10w0d-13w6d)
017270	<input type="checkbox"/>	Serum Integrated 2 (15w0d-21w6d)
017319	<input type="checkbox"/>	AFP Tetra (15w0d-21w6d)
010801	<input type="checkbox"/>	msAFP (15w0d-23w6d)

MATERNAL SERUM SCREENING W NT

017500	<input type="checkbox"/>	1st Trimester Screen (10w0d-13w6d)
017700	<input type="checkbox"/>	Sequential Part 1 (10w0d-13w6d)
017750*	<input type="checkbox"/>	Sequential Part 2 (15w0d-21w6d)
017100	<input type="checkbox"/>	Integrated Part 1 (10w0d-13w6d)
017170*	<input type="checkbox"/>	Integrated Part 2 (15w0d-21w6d)

CARRIER SCREENING

480533 Cystic Fibrosis (32)

450020 CFplus[®] (97 mutation test)

511919 Frag X, PCR w/ rfx to Southern blot analysis**

450010 Spinal Muscular Atrophy

121679 Hemoglobinopathy Profile

511172 α- Thalassemia, DNA Analysis

252823 β- Thalassemia: HBB (Full Gene Sequencing)

Inherited Carrier Screen

451910 Gene-specific Sequencing**

Required: Gene (s) _____

451950 Inheritest[®] Comprehensive Panel (144 genes)

451920 Inheritest[®] Ashkenazi Jewish Panel (48 genes)

451960 Inheritest[®] Society-guided Panel (14 genes)

451964 Inheritest[®] Core (CF97, SMA, FragX)

452172 Inheritest[®] CF/SMA Panel

NT MEASUREMENTS

CRL date ____/____/____ CRL ____ mm (45.0-84.0) NT ____ mm Chorionicity: Mono DI

Twin B, if applicable CRL ____ mm (45.0-84.0) NT ____ mm Unknown

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 Present Absent

Please also check YES under "Other Indications" in Clinical History section if NB data provided.

* Gestational age will be based on CRL data provided for Part 1. Integrated & Sequential Testing options require 2 specimens within a specified period. Part 2 follow-up information will be listed on the Part 1 report.

+ The NT and nasal bone must be performed by a sonographer credentialed by the FMF, NTQR or equivalent entity.

REQUIRED INFORMATION	Patient Weight _____ lbs	CLINICAL HISTORY	<input type="checkbox"/> Yes <input type="checkbox"/> No	Prior Down Syndrome/ONTD Screen in Current Pregnancy? If yes, prior test was:
	# of Fetuses <input type="checkbox"/> 1 <input type="checkbox"/> 2 <input type="checkbox"/> Other _____		<input type="checkbox"/> Yes <input type="checkbox"/> No	<input type="checkbox"/> in 1st Tri <input type="checkbox"/> in 2nd Tri <input type="checkbox"/> elevated msAFP
	Patient Race <input type="checkbox"/> Cauc <input type="checkbox"/> Hispanic <input type="checkbox"/> Black		<input type="checkbox"/> Yes <input type="checkbox"/> No	Family history of NTD?
	<input type="checkbox"/> Asian <input type="checkbox"/> Amer Indian <input type="checkbox"/> Other		<input type="checkbox"/> Yes <input type="checkbox"/> No	Previous pregnancy with Down Syndrome?
<input type="checkbox"/> Yes <input type="checkbox"/> No	Is patient an insulin dependent diabetic?	<input type="checkbox"/> Yes <input type="checkbox"/> No	Parental cytogenetics following abnormal prenatal results.	
<input type="checkbox"/> Yes <input type="checkbox"/> No	Egg donor: <input type="checkbox"/> Self <input type="checkbox"/> Non-self	<input type="checkbox"/> Yes <input type="checkbox"/> No	Parental balanced Robertsonian Translocation with increased risk of Trisomy.	
	Age of donor at egg retrieval: ____ years	<input type="checkbox"/> Yes <input type="checkbox"/> No	Other Indications: _____	

PREGNANCY COMPLICATIONS

005199 Prothrombin Time (PT)

005207 PTT, Activated (APTT)

365200 IUFD Profile

365300 IUFD Extended Profile

161802 Anticardiolipin Ab, IgG, IgM

015594 Antithrombin Deficiency Profile

001610 Fibrinogen Activity

511162 Factor II (Prothrombin)

511154 Factor V Leiden

706994 Homocysteine

365500 Inherited Thrombophilias of Preg

117892 Lupus Anticoagulant w Reflex

512358 MTHFR

283655 Protein C Deficiency Profile

117754 Protein S Deficiency Profile

504295 ReproSURE[™] (Ovarian Reserve Profile) See Reverse (GEL)

CYTOGENETICS

Amniotic Fluid (specify GA above)			Chorionic Villi (specify GA above)			POC / Tissue (specify GA above)		
511580	<input type="checkbox"/>	Chromosome & AFP/ACH/HbF	510988	<input type="checkbox"/>	Chromosome Analysis	052052	<input type="checkbox"/>	Chromosome Analysis
052040	<input type="checkbox"/>	Chromosome Analysis	510960	<input type="checkbox"/>	InSight [®] Prenatal CVS Aneuploid FISH	510110	<input type="checkbox"/>	Reveal [®] SNP Microarray
002428	<input type="checkbox"/>	AFP, Amniotic fluid	511033	<input type="checkbox"/>	Chromosome rfx Reveal [®] SNP Microarray	052065	<input type="checkbox"/>	Chromosome rfx Reveal [®] SNP Microarray
510305	<input type="checkbox"/>	AFP, AChE with reflex to HbF	511625	<input type="checkbox"/>	FISH, rfx Chrom. or Reveal [®] SNP Microarray	511035	<input type="checkbox"/>	Chromosome, Blood
511894	<input type="checkbox"/>	InSight [®] -Prenatal Amnio Aneuploid FISH	511555	<input type="checkbox"/>	Chrom. 5 cell + Reveal [®] SNP Microarray	510770	<input type="checkbox"/>	FISH Microdeletion (Specify)
052104	<input type="checkbox"/>	Chromosome rfx Reveal [®] SNP Microarray	510100	<input type="checkbox"/>	SNP Microarray - Prenatal (Reveal [®])			
511966	<input type="checkbox"/>	FISH, rfx chrom. or Reveal [®] SNP Microarray	510200	<input type="checkbox"/>	SNP Microarray (Direct) - Prenatal (Reveal [®])			
511590	<input type="checkbox"/>	Chrom. 5 Count + Reveal [®] SNP Microarray			Clinical Info, Family History: _____			Other: _____
510100	<input type="checkbox"/>	SNP Microarray-Prenatal (Reveal [®])			Abnl NIPT: Specify _____			Fetal Gender: <input type="checkbox"/> Male <input type="checkbox"/> Female <input type="checkbox"/> Unknown
510200	<input type="checkbox"/>	SNP Microarray (Direct)-Prenatal (Reveal [®])						

* Reflex testing will cause additional CPT codes to be billed.

1-FORWARD TO LABCORP WITH SPECIMEN-RETAIN LAST COPY

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Form containing patient information: Patient's Legal Name, Sex, Date of Birth, Collection Time, Fasting, Collection Date, Urine hrs/vol, NPI, Physician's ID #, Patient's ID #, Hospital Patient Status, Physician's Name, Physician/Authorized Signature, Patient's Address, Phone, City, State, ZIP, Name of Policy Holder, Address of Policy Holder, APT #, City, State, ZIP, Insurance Carrier, ID #, Group #, Insurance Address, Name of Insured Person, Relationship to Patient, Employer Name, and Medicare Advance Beneficiary Notice of Noncoverage (ABN).

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

MATERNAL PLASMA SCREENING: NON INVASIVE PRENATAL TESTING (NIPT) section with test options like MaterniT 21 PLUS, MaterniT 21 PLUS w/ ESS, MaterniT 21 PLUS w/ SCA, MaterniT Genome, MaterniT 21 PLUS No Gender, GENOME-Flex, and GENOME-Flex**.

GA ___ wks ___ days on date ___/___/___, By ___ LMP ___ EDC/EDD ___ U/S, Indication for Non Invasive Prenatal Testing (NIPT) with checkboxes for AMA, Previous pregnancy with aneuploidy, Family history of chromosomal abnormality, Ultrasound findings, cystic hygroma, IUGR, heart defect, CNS abnormality, and Other.

MATERNAL SERUM SCREENING W/O NT section with test options: 017200 Serum Integrated 1 (10w0d-13w6d), 017270 Serum Integrated 2 (15w0d-21w6d), 017319 AFP Tetra (15w0d-21w6d), 010801 msAFP (15w0d-23w6d).

MATERNAL SERUM SCREENING W NT section with test options: 017500 1st Trimester Screen (10w0d-13w6d), 017700 Sequential Part 1 (10w0d-13w6d), 017750* Sequential Part 2 (15w0d-21w6d), 017100 Integrated Part 1 (10w0d-13w6d), 017170* Integrated Part 2 (15w0d-21w6d).

CARRIER SCREENING section with test options: 480533 Cystic Fibrosis (32), 450020 CFplus (97 mutation test), 511919 Frag X, PCR w/ rfx to Southern blot analysis**, 450010 Spinal Muscular Atrophy, 121679 Hemoglobinopathy Profile, 511172 alpha-Thalassemia, DNA Analysis, 252823 beta-Thalassemia: HBB (Full Gene Sequencing), Inheritest Carrier Screen, 451910 Gene-specific Sequencing**.

PREGNANCY COMPLICATIONS section with test options: 005199 Prothrombin Time (PT), 005207 PTT, Activated (APTT), 365200 IUFD Profile, 365300 IUFD Extended Profile, 161802 Anticardiolipin Ab, IgG, IgM, 015594 Antithrombin Deficiency Profile, 001610 Fibrinogen Activity, 511162 Factor II (Prothrombin), 511154 Factor V Leiden, 706994 Homocysteine, 365500 Inherited Thrombophilias of Preg, 117892 Lupus Anticoagulant w Reflex, 512358 MTHFR, 283655 Protein C Deficiency Profile, 117754 Protein S Deficiency Profile, 504295 ReproSURE (Ovarian Reserve Profile).

NT MEASUREMENTS section with CRL date, CRL measurements, NT measurements, Chorionicity, Sonographer Name, Sonographer ID, Reading MD ID, Nasal Bone, and NB Twin B checkboxes.

REQUIRED INFORMATION section with Patient Weight, # of Fetuses, Patient Race, Is patient an insulin dependent diabetic?, Egg donor, Age of donor at egg retrieval, and CLINICAL HISTORY checkboxes for Down Syndrome, Family history of NTD, Previous pregnancy with Down Syndrome, Parental cytogenetics, Parental balanced Robertsonian Translocation, and Other Indications.

CYTOGENETICS section with Amniotic Fluid, Chorionic Villi, and POC / Tissue test options, including Chromosome & AFP/ACHE/HbF, Chromosome Analysis, InSight Prenatal CVS Aneuploid FISH, Chromosome rfx Reveal SNP Microarray, FISH, rfx Chrom. or Reveal SNP Microarray, Chrom. 5 cell + Reveal SNP Microarray, SNP Microarray - Prenatal (Reveal), SNP Microarray (Direct) - Prenatal (Reveal), Chromosome Analysis, Reveal SNP Microarray, Chromosome rfx Reveal SNP Microarray, Chromosome, Blood FISH Microdeletion (Specify), and Clinical Info, Family History, Other, and Fetal Gender checkboxes.

Test No.	Description	Specimen	CPTs	Components
Genetic Disorders				
480533	Cystic Fibrosis (32)	7mL LAV *	81220	Includes more than 110 disorders including SMA, Cystic Fibrosis and Fragile X Includes more than 13 disorders included in ACMG and ACOG guidelines Includes more than 40 disorders specific to individuals of Ashkenazi descent
450020	CF ^{plus} ® (97 mutation test)	10mL YEL	81220	
511919	Frag X, PCR w/ rfx to Southern blot analysis	10mL LAV	81243	
450010	Spinal Muscular Atrophy	10mL YEL	81329	
121679	Hemoglobinopathy Profile	1mL LAV	83021; 85660	
451950	Inheritest® Comprehensive Panel		81443, 81329, 81243	
451960	Inheritest® Society-guided Panel		81220, 81243, 81329, 81200, 81209, 81242, 81251, 81361, 81260, 81290, 81330, 81255, 81257	
451920	Inheritest® Ashkenazi Jewish Panel		81329, 81243, 81257, 81412	
451382	Mutation Sequence Analysis (call before sending)			
451385	Prenatal Mutation Specific Sequencing (call before sending)			
511172	α-Thalassemia, DNA Analysis	7mL LAV *	81257	
252823	β-thalassemia	2mL LAV	81364	
Pregnancy Complications				
005199	Prothrombin Time (PT)	5mL BLU	85610	CBC; TSH; Human Parvovirus B19 IgG & IgM; Lupus Anticoagulant; Anticardiolipin Ab IgG, IgM; RPR 365200 plus Factor V Leiden; Factor II (Prothrombin); Antithrombin Activity; Homocysteine; Protein S Antigen, free; Protein C Activity Antithrombin Activity, Antithrombin Antigen
005207	PTT, Activated (APTT)	see DOS	85730	
365200	IUFD Panel	see DOS	84443, 86592, 86747 x2, 85732, 85613, 86147 x2, 85025	
365300	IUFD Extended Panel	see DOS	81240, 81241, 83090, 84443, 86592, 86747(x2), 85300, 85303, 85306, 85732, 85613, 86147(x2), 85025	
161802	Anticardiolipin Ab, IgG, IgM	1mL GEL	86147(x2)	
015594	Antithrombin Deficiency Profile	3mL BLU +	85300; 85301	
001610	Fibrinogen Activity	see DOS	85384	
511162	Factor II (prothrombin)	7mL LAV *	81240	
511154	Factor V Leiden	7mL LAV *	81241	
706994	Homocysteine	2mL LAV	83090	
365500	Inherited Thrombophilias of Pregnancy	see DOS	81240, 85300, 85303, 85306, 85307	
117892	Lupus Anticoagulant w Reflex	6mL BLU	85613; 85732	Lupus sensitive APTT & Dilute Russel Viper Venom Time; if prolonged, confirmation performed
511238	MTHFR	7mL LAV *	81291	
283655	Protein C Deficiency Profile	3mL BLU +	85302; 85303	Protein C, Functional, Protein C, Antigen
117754	Protein S Deficiency Profile	3mL BLU	85305; 85306(x2)	Protein S Antigen, Total, Protein S Antigen, Free, Protein S, Functional
Maternal Plasma Testing: Non Invasive Prenatal Testing (NIPT)				
451927	MaterniT21® PLUS (9w+)	MaterniT Collection Kit	Call Client Services	Please go to www.labcorp.com
451931	MaterniT21® PLUS w/ ESS (9w+)	MaterniT Collection Kit	Call Client Services	Please go to www.labcorp.com
451934	MaterniT21® PLUS w/ SCA (9w+)	MaterniT Collection Kit	Call Client Services	Please go to www.labcorp.com
451937	MaterniT21® PLUS w/ ESS & SCA (9w+)	MaterniT Collection Kit	Call Client Services	Please go to www.labcorp.com
451941	MaterniT® Genome (9w+)	MaterniT Collection Kit	Call Client Services	Please go to www.labcorp.com
451951	MaterniT21® PLUS No Gender (9w+)	MaterniT Collection Kit	Call Client Services	Please go to www.labcorp.com
452104	GENOME-Flex (Add On)	No collection required	Call Client Services	Please go to www.labcorp.com
452114	GENOME-Flex (Add On) Redraw	MaterniT Collection Kit	Call Client Services	Please go to www.labcorp.com
Maternal Serum Testing				
017500	1st Trimester Screen	3mL GEL	84163, 84702, 86336	PAPP-A, hCG, DIA
017700	Sequential Part 1	3mL GEL	84163, 84702	PAPP-A, hCG
017750	Sequential Part 2	5mL GEL	82105, 82677, 84702, 86336	AFP, uE3, hCG, DIA
017100	Integrated Part 1	3mL GEL	84163	PAPP-A
017170	Integrated Part 2	5mL GEL	82105, 82677, 84702, 86336	AFP, uE3, hCG, DIA
017200	Serum Integrated 1	3mL GEL	84163	PAPP-A
017270	Serum Integrated 2	5mL GEL	82105, 82677, 84702, 86336	AFP, uE3, hCG, DIA
017319	AFP Tetra	5mL GEL	82105, 82677, 84702, 86336	AFP, uE3, hCG, DIA
010801	msAFP	3mL GEL	82105	AFP
Cytogenetics				
511580	Chromosome & AFP/AChE/HbF	Amnio (20-30mL)	Call CPT Coding 800-222-7566 Ext. 68400	Amniotic fluid, cultured cells, or chorionic villus sample (CVS). Please submit maternal blood (sodium heparin or EDTA) for maternal cell contamination (MCC) studies. Volume 10 to 20 mL amniotic fluid, 2 T-25 flasks, or 10 to 20 mg CVS ReproSURE™ (Ovarian Reserve Profile) Test No. 504295 When ordered as a profile CPT Codes used: 82397, 82670, 83001 ReproSURE is a blood test comprised of AMH, FSH and Estradiol hormones designed to provide information about ovarian reserve.
052040	Chromosome Analysis	Amnio (20-30mL)	Call CPT Coding 800-222-7566 Ext. 68400	
002428	AFP, Amniotic fluid	Amnio (2mL)	82106	
510305	AFP, AChE with reflex to HbF	Amnio (2mL)	82013, 82106	
511894	FISH, Prenatal Aneuploidy	Amnio (3-5mL)	Call CPT Coding 800-222-7566 Ext. 68400	
052104	Chromosome rfx Reveal® SNP Microarray	Amnio (20-30mL)	Call CPT Coding 800-222-7566 Ext. 68400	
511966	FISH, reflex chromosomes or Reveal® SNP Microarray	Amnio (25mL)	Call CPT Coding 800-222-7566 Ext. 68400	
511590	Chromosome Five-cell Count Plus Reveal® SNP Microarray	Amnio (25mL)	Call CPT Coding 800-222-7566 Ext. 68400	
510100	SNP Microarray-Prenatal (Reveal®)	Amnio (25mL)/CVS (20-30 mg)	Call CPT Coding 800-222-7566 Ext. 68400	
510200	SNP Microarray (Direct)-Prenatal (Reveal®)	Amnio (25mL)/CVS (20-30 mg)	Call CPT Coding 800-222-7566 Ext. 68400	
510988	Chromosome Analysis	CVS (20-30mg)	Call CPT Coding 800-222-7566 Ext. 68400	
510960	FISH, Prenatal Aneuploidy	CVS (5mg)	Call CPT Coding 800-222-7566 Ext. 68400	
511033	Chromosome rfx Reveal® SNP Microarray	CVS (20-30mg)	Call CPT Coding 800-222-7566 Ext. 68400	
511625	FISH, reflex chromosomes or Reveal® SNP Microarray	CVS (20-30mg)	Call CPT Coding 800-222-7566 Ext. 68400	
511555	Chromosome Five-cell Count Plus Reveal® SNP Microarray	CVS (20-30mg)	Call CPT Coding 800-222-7566 Ext. 68400	
052052	Chromosome, Biopsies	POC/Skin	Call CPT Coding 800-222-7566 Ext. 68400	
510110	Reveal® SNP Microarray	POC/Tissue	Call CPT Coding 800-222-7566 Ext. 68400	
052065	Chromosome rfx Reveal® SNP Microarray	POC/Tissue	Call CPT Coding 800-222-7566 Ext. 68400	
511035	Chromosome, Blood	5mL GRN		
510770	FISH Microdeletion (specify)	see DOS		

* Buccal swab also acceptable + Two tubes required (1/2 volume in each)

The lab only accepts isolated or extracted nucleic acids for which extraction or isolation is performed in an appropriately qualified CLIA or CAP/CMS equivalent laboratory