

GENETIC CARRIER & PRENATAL SCREENING – EXON

FEMALE PATIENT MALE PATIENT

LAST NAME _____

FIRST NAME _____

PATIENT ID _____

DATE OF BIRTH (MM/DD/YYYY) _____ TELEPHONE NUMBER _____

Asian African American/Black Hispanic
 Caucasian/White Jewish, Ashkenazi Other/Mixed/Unknown _____

EMAIL _____

STREET NUMBER _____ STREET NAME _____ APT NUMBER _____

CITY _____ STATE _____ ZIP _____

ACKNOWLEDGEMENT: I authorize the laboratory to provide to my health plan the information on this form and other information provided by my healthcare provider if necessary for reimbursement. I understand that the laboratory may seek prior authorization for testing from my health plan on my behalf. I also authorize all benefits of the plan to be payable directly to the laboratory, and I agree to remit to the laboratory any payment for these services made directly to me. I understand that the laboratory may be an out-of-network provider for my health plan and that I am responsible for all amounts not reimbursed by my health plan. I hereby designate the laboratory as my Authorized Representative, as provided under ERISA, 29 C.F.R. § 2560.5031 (b)(4), and/or as my Attorney in Fact, for the purpose of pursuing administrative appeals to which I am entitled and, if the laboratory deems it appropriate, any legal and/or equitable claims that I could bring against my health plan, and/or its fiduciaries, and/or its administrators, with respect to their handling or resolution of my insurance claim. I authorize information to be shared with my partner if also undergoing testing.

I authorize the laboratory to retain and use my de-identified specimen and test data (where all information that could link me to the specimen or data has been removed) for research and/or help develop new products or services, in compliance with applicable laws.

I do not authorize the laboratory to retain and use my de-identified specimen and test data as described above.

All leftover specimens from New York State will be destroyed within 60 days.

REQUIRED **PATIENT SIGNATURE** _____ DATE (MM/DD/YY) _____

PREGNANCY INFO – REQUIRED FOR FEMALE PATIENTS

Is patient pregnant?
 NO Z31.430
 YES, singleton
 YES, twin
 YES, other: _____

First pregnancy?
 YES, primigravida
 NO, multigravida
Which trimester?
 First Z34.01/81
 Second Z34.02/82
 Third Z34.03/83

EST. DATE OF DELIVERY (MM/DD/YY) _____ DATING METHOD
 LMP Ultrasound

PATIENT HEIGHT _____ FT _____ IN PATIENT WEIGHT _____ LBS

Egg donor or surrogate pregnancy 009.811
 Age of donor _____

CARRIER TESTING

PREPARENT® EXON CARRIER TEST
 Full-exon sequencing and select copy number variant analysis.
 For disorders tested, visit: progenity.com/resources

5030 5040 **Trio** CF, SMA, and FX

5050 5060 **Standard**¹ 25 genes with ACOG/ACMG guidelines

5010 5020 **Exon**¹ 150+ genes

5064 5068 **Global**¹ 280+ genes

Other (All tests can be ordered individually.) _____

¹ In certain cases, orders for Standard, Exon 150, or Global may be adjusted to Trio based on health plan policy.

PRENATAL SCREENING

INNATAL® PRENATAL SCREEN Gestational age 10 weeks or later
 Test ordered will be processed and billed based upon health plan policy.

2800 **Singleton Pregnancy** noninvasive screening for chromosomes 21, 18, 13, X, and Y
 OPT OUT of screening for sex chromosome aneuploidies (no fetal sex information)

2800 **Twin Pregnancy** noninvasive screening for chromosomes 21, 18, 13, and Y
 OPT OUT of screening for Y chromosome (no fetal sex information)

MATERNAL SERUM SCREENING
 Patient weight is required for all maternal serum screening tests.

2901 **First Trimester Screen (PAPP-A, hCG)**
 Gestational age 10 – 13 weeks 6 days
Requires nuchal translucency ultrasound.
 For twins, attach separate form: progenity.com/ultrasound

2902 **Second Trimester Quad Screen (AFP, uE3, hCG, DIA)**
 Gestational age 15 – 22 weeks 6 days

2900 **Open Neural Tube Screen (AFP)** Gestational age 15 – 22 weeks 6 days

ULTRASOUND INFO

ULTRASOUND DATE _____

CROWN RUMP LENGTH _____ mm (34.0 – 85.0)

NUCHAL TRANSLUCENCY _____ mm

CLINICIAN INFO

ACKNOWLEDGEMENT: I hereby confirm that information has been provided to the patient about the test(s) to be performed and the patient has given consent as required under applicable laws and regulations for the test(s) to be performed. The patient has completed pre-testing genetic counseling. The test(s) to be performed are medically necessary and the results will be used for medical management and treatment decision purposes for this patient. I confirm that the person listed as the Ordering Clinician is authorized by law to order the test(s) requested herein.

REQUIRED **ORDERING CLINICIAN SIGNATURE** _____ DATE (MM/DD/YY) _____

ADDITIONAL REPORTS TO: CLINICIAN NAME _____

NPI# _____ FAX NUMBER _____

BILLING INFO

BILL INSURANCE Attach legible front and back copy of insurance cards.

INSURANCE COMPANY _____ IPA NAME _____

MEMBER ID _____ PRIOR AUTHORIZATION, IF NEEDED _____

BILL PATIENT Patient will be contacted to provide payment method.
 CLIENT BILL

SPECIMEN INFO – REQUIRED

Collected on: _____ Time: _____ AM PM

Collected by: _____

Collection Requirement Key: See Specimen Guide for additional specimen types

4 mL Lavender-top EDTA tube 4 mL serum separator tube (SST)
 10 mL Streck DNA tube

CLINICAL INFO

REQUIRED - Family history of genetic disorder?

NO, and patient is female Z31.430
 NO, and patient is male Z31.440
 YES, Hereditary genetic disorder Z84.81 (specify condition) _____
 YES, Musculoskeletal disorder Z82.69 (specify condition) _____

Patient had previous carrier testing (attach report)
 Patient had a blood transfusion (past 3 months) or a bone marrow/organ transplant

PARTNER INFO

PARTNER'S FIRST NAME _____
 PARTNER'S DATE OF BIRTH (MM/DD/YYYY) _____
 PARTNER'S LAST NAME _____

SPECIFY CONDITION _____
 RELATIONSHIP TO PATIENT OR PARTNER _____
 OTHER DIAGNOSIS (SPECIFY ICD-10) _____

CLINICAL INFO

REQUIRED - Is patient at increased risk for aneuploidy (i.e., Down syndrome)?

NO, patient is average risk. Z13.79
 YES, advanced maternal age (35+ at EDD) 009.511, 512, 513, 519, 521, 522, 523, 529
 YES, abnormal serum screening 028.1 (specify aneuploidy risk) _____
 YES, ultrasound indicating structural anomaly 028.3 (specify aneuploidy risk) _____

YES, prior pregnancy w/ aneuploidy Z82.79 (specify condition) _____
 YES, family history of chromosomal abnormality Z82.79 (specify condition) _____
 YES, patient history of chromosomal abnormality (specify condition) _____

Patient had a blood transfusion (past 3 months) or a bone marrow/organ transplant
 OTHER DIAGNOSIS (SPECIFY ICD-10) _____

CLINICAL INFO

Patient is an insulin-dependent diabetic
 Patient smokes cigarettes

OTHER DIAGNOSIS (SPECIFY ICD-10) _____

Credentialed by: NTQR FMF Other _____

NOTE: Carrier testing for X-linked disorders is not performed in males.

HEMOGLOBIN EVALUATION

| | | |
|----------------------|-----------------------------------|-----------------------------------|
| Red blood cell count | Mean corpuscular volume (MCV) | Red cell distribution width (RDW) |
| Hemoglobin | Mean corpuscular hemoglobin (MCH) | Hemoglobin electrophoresis |
| Hematocrit | | |

CLINICAL INDICATIONS

GENETIC CARRIER SCREENING

| | |
|--|---------|
| Screening for other metabolic disorders | Z13.228 |
| Nonprocreative screening for genetic disease carrier status | Z13.71 |
| Female for testing for genetic disease carrier status for procreative management | Z31.430 |
| Male for testing for genetic disease carrier status for procreative management..... | Z31.440 |
| Supervision of normal first pregnancy, first trimester | Z34.01 |
| Supervision of normal first pregnancy, second trimester | Z34.02 |
| Supervision of normal first pregnancy, third trimester | Z34.03 |
| Supervision of other normal pregnancy, first trimester | Z34.81 |
| Supervision of other normal pregnancy, second trimester..... | Z34.82 |
| Supervision of other normal pregnancy, third trimester..... | Z34.83 |
| Supervision of normal pregnancy, unspecified, first trimester | Z34.91 |
| Supervision of normal pregnancy, unspecified, second trimester..... | Z34.92 |
| Supervision of normal pregnancy, unspecified, third trimester..... | Z34.93 |
| Family history of intellectual disabilities | Z81.0 |
| Family history of other diseases of the musculoskeletal system and connective tissue | Z82.69 |
| Family history of carrier of genetic disease | Z84.81 |

PRENATAL ANEUPLOIDY SCREENING

| | |
|--|----------|
| Supervision of pregnancy with history of infertility, first trimester | 009.01 |
| Supervision of pregnancy with history of infertility, second trimester | 009.02 |
| Supervision of pregnancy with other poor reproductive or obstetric history, first trimester | 009.291 |
| Supervision of pregnancy with other poor reproductive or obstetric history, second trimester | 009.292 |
| Supervision of elderly primigravida, first trimester | 009.511 |
| Supervision of elderly primigravida, second trimester | 009.512 |
| Supervision of elderly primigravida, third trimester | 009.513 |
| Supervision of elderly multigravida, first trimester | 009.521 |
| Supervision of elderly multigravida, second trimester | 009.522 |
| Supervision of elderly multigravida, third trimester | 009.523 |
| Supervision of pregnancy resulting from assisted reproductive technology, first trimester | 009.811 |
| Supervision of pregnancy resulting from assisted reproductive technology, second trimester | 009.812 |
| Supervision of pregnancy resulting from assisted reproductive technology, third trimester | 009.813 |
| Supervision of other high risk pregnancies, first trimester | 009.891 |
| Supervision of other high risk pregnancies, second trimester | 009.892 |
| Supervision of other high risk pregnancies, third trimester | 009.893 |
| Abnormal hematological finding on antenatal screening of mother | 028.0 |
| Abnormal biochemical finding on antenatal screening of mother | 028.1 |
| Abnormal cytological finding on antenatal screening of mother | 028.2 |
| Abnormal ultrasonic finding on antenatal screening of mother | 028.3 |
| Abnormal radiological finding on antenatal screening of mother | 028.4 |
| Abnormal chromosomal and genetic finding on antenatal screening of mother | 028.5 |
| Other abnormal findings on antenatal screening of mother | 028.8 |
| Unspecified abnormal findings on antenatal screening of mother | 028.9 |
| Maternal care for (suspected) chromosomal abnormality in fetus, fetus 1 | O35.1XX1 |
| Maternal care for (suspected) hereditary disease in fetus, fetus 1 | O35.2XX1 |
| Other screening for genetic and chromosomal anomalies | Z13.79 |
| Family history of other disabilities and chronic diseases leading to disablement, not elsewhere classified | Z82.8 |