

1 SELECT TEST PANEL

Samples without a test panel selected will not be processed.

UNITY Carrier Screen™

☐ ACOG Guideline Panel
CFTR • SMN1 • HBB • HBA1 • HBA2☐ add Fragile X Syndrome[†]
FMR1☐ add Plus Panel[†]
ACADM • ASPA • DHCR7 • DMD
• ELP1 • HEXA • PAH • PMM2A separate cfDNA fetal risk assay will be run
for carrier positive singleton pregnancies.
Check here to opt out: ☐UNITY Aneuploidy Screen™
and 22q11.2 Microdeletion
singleton pregnancy☐ chromosomes:
13 • 18 • 21 • X • Y • 22q11.2

OR

☐ order for twin pregnancy
NIPT analysis[†]
13, 18, 21, and 22q11.2 only. Zygosity included.☐ add Fetal RhD NIPT for RhD-negative mothers[†]☐ opt out fetal sex[†][†] These test options include their respective primary panel (highlighted yellow checkbox)UNITY Aneuploidy Screen™
singleton pregnancy☐ chromosomes:
13 • 18 • 21 • X • Y☐ order for twin pregnancy
NIPT analysis[†]
13, 18, and 21 only. Zygosity included.

2 SAMPLE COLLECTION DATE & BARCODE

MM-DD-YYYY

PLACE PROVIDED BARCODE HERE

3 CLINICAL INDICATION

Required to select at least one for each test panel*.
The following codes are not exhaustive.**UNITY Carrier Screen** If the patient with singleton pregnancy is found to be a carrier for tested disorders, fetal risk assessment will be performed and O28.5 ICD-10 code will be applied, unless opted out.

- | | | | |
|--|---------|---------|--------|
| <input type="checkbox"/> Family history of carrier genetic disease** | Z84.81 | Z84.89 | |
| <input type="checkbox"/> Testing for genetic disease carrier status | Z31.430 | Z31.440 | |
| <input type="checkbox"/> Supervision of other normal pregnancy | Z34.81 | Z34.82 | Z34.83 |
| <input type="checkbox"/> Family history of intellectual disabilities | Z81.0 | | |
| <input type="checkbox"/> Other: | | | |

UNITY Aneuploidy Screen Code determined by trimester of pregnancy.

- | | | | |
|---|---------|---------|---------|
| <input type="checkbox"/> Supervision of elderly primigravida | O09.511 | O09.512 | O09.513 |
| <input type="checkbox"/> Supervision of elderly multigravida | O09.521 | O09.522 | O09.523 |
| <input type="checkbox"/> Supervision of other high risk pregnancies | O09.891 | O09.892 | O09.893 |
| <input type="checkbox"/> Supervision other normal pregnancy | Z34.81 | Z34.82 | Z34.83 |
| <input type="checkbox"/> Abnormal ultrasound findings | O28.3 | | |
| <input type="checkbox"/> Abnormal chromosomal & genetic findings | O28.5 | | |
| <input type="checkbox"/> Other: | | | |

UNITY Fetal RhD/Fetal Antigen Code determined by trimester of pregnancy.

- | | | | |
|---|----------|----------|----------|
| <input type="checkbox"/> Maternal care for anti-D [Rh] antibodies | O36.0110 | O36.0120 | O36.0130 |
| <input type="checkbox"/> Encounter for Rh incompatibility status | Z31.82 | | |
| <input type="checkbox"/> Maternal care for other isoimmunization | O36.1910 | O36.1920 | O36.1930 |
| <input type="checkbox"/> Other: | | | |

*ICD-10 code selected under one test panel might be used in other test panels' billing.
**Requires Additional Code

Ordering Healthcare Provider(s) *

PROVIDER AUTHORIZATION

By submission of this requisition and accompanying sample, I hereby authorize and direct BillionToOne to (1) utilize the above information to process the indicated test for this patient and (2) release the results and patient information to the patient's third-party payer, as needed. I certify (1) all information provided herein is true and accurate, (2) I am authorized by law to request the test, (3) the test is reasonable and medically necessary for the treatment and management of this patient, (4) the patient has been counseled on the potential results, benefits and limitations of the test, and (5) I have obtained informed consent to the extent required under applicable law. I agree to provide the necessary information and medical records to BillionToOne needed to submit and process claims to payers.

MM-DD-YYYY

Provider Signature *

Date of Authorization

5 CLINIC INFORMATION

Clinic Name *

Clinic Phone

Clinic Fax

Clinic Account Number

Additional Notes

4 PATIENT INFORMATION

First Name *	MI	Last Name *
	lbs	MM-DD-YYYY
Sex assigned at birth *	Maternal Weight	Date of Birth *
		Estimated Due Date *
		<input type="checkbox"/> not pregnant

By providing the information below, I agree I or my provider may be contacted for test status, billing/ collection, marketing, quality assurance, or research purposes.

Cell Phone *	Email
Street Address	Apt / Unit / Suite
City	State
	Zip Code

Pregnancy Details Select if applicable

- ☐
- twins
- ☐
- triplets or more
- ☐
- vanishing twin
-
- ☐
- egg donor/gestational carrier age of egg donor

Abnormal Ultrasound Findings

Ethnicity or Race ICD-10 code Z15.89 will be applied for high risk ethnicities.

- ☐
- Asian
- ☐
- African/African American
- ☐
- Ashkenazi Jewish
- ☐
- Middle Eastern
-
- ☐
- French Canadian/Cajun
- ☐
- Hispanic
- ☐
- White
- ☐
- other
- ☐
- unknown

Reported Carrier History Include condition and variant when possible

- ☐
- Known Maternal Carrier
-
- ☐
- Known Paternal Carrier

Link to Reproductive Partner (if applicable) Include first, last name, and DOB

- ☐
- Maternal
- ☐
- Paternal

PATIENT ACKNOWLEDGEMENT I acknowledge I have read and agreed to the Patient Acknowledgement for testing on the back page.

MM-DD-YYYY

Patient Signature

Date of Acknowledgement

6 BILLING INFORMATION

- ☐
- Bill to Insurance
- ☐
- Bill to Patient
- ☐
- Bill to Client

☐ Ordering provider or facility is out-of-network with the patient's insurance plan

Attach copy of insurance card

Insurance Company Name

Member ID

Group ID

TEST PANEL	TEST DETAILS	SAMPLE REQUIREMENT
UNITY Complete®	UNITY Carrier Screen + UNITY Aneuploidy Screen See conditions below	3 × 10 mL Streck cell-free DNA BCT® blood tube Fill to the top (≥ 8mL)
UNITY Carrier Screen <i>Recessive Conditions Screened</i> <i>cfDNA fetal risk assessment will be provided for positive carrier test results unless opted out</i>	ACOG Guideline Panel <ul style="list-style-type: none"> Cystic Fibrosis <i>CFTR</i> Hemoglobinopathies (Sickle Cell Disease, Alpha / Beta Thalassemias) <i>HBA1, HBA2, HBB</i> Spinal Muscular Atrophy <i>SMN1</i> Fragile X Syndrome <i>FMR1*</i> Plus Panel <ul style="list-style-type: none"> Canavan Disease <i>ASPA</i> DMD-Associated Dystrophinopathies <i>DMD*</i> Familial Dysautonomia <i>ELP1</i> Medium-Chain Acyl-CoA Dehydrogenase Deficiency <i>ACADM</i> Phenylalanine Hydroxylase Deficiency <i>PAH</i> PMM2-Congenital Disorder of Glycosylation <i>PMM2</i> Smith-Lemli-Opitz Syndrome <i>DHCR7</i> Tay Sachs Disease <i>HEXA</i> 	1 × 10 mL Streck cell-free DNA BCT® blood tube Fill to the top (≥ 8mL)
UNITY Aneuploidy Screen <i>Chromosomal Conditions Screened</i>	<ul style="list-style-type: none"> Down Syndrome <i>T21</i> Edwards Syndrome <i>T18</i> Patau Syndrome <i>T13</i> Sex Chromosome Aneuploidies <i>Monosomy X, XXY, XXX, XYY</i> Zygosity for Twin Pregnancies Optional: Fetal Sex Optional: Fetal RhD for RhD-negative Pregnant Patients Optional: 22q11.2 Microdeletion Syndrome 	2 × 10 mL Streck cell-free DNA BCT® blood tube Fill to the top (≥ 8mL)

UNITY Fetal Risk Screen: cfDNA fetal risk assessment for recessive conditions can be performed at ≥9 weeks gestation, can only be performed for singleton pregnancies, and cannot be performed for egg donors or gestational carriers. *Carrier screening for X-linked conditions is not performed for male patients, and cfDNA fetal risk assessment is provided via fetal sex, unless fetal-sex has been opted out. **UNITY Aneuploidy:** can be performed at ≥9 weeks gestation. Sex chromosome aneuploidies can only be performed for singleton pregnancies.

ICD-10 DIAGNOSIS CODES
Codes below are not exhaustive, provide additional codes as necessary.

UNITY Carrier Screen		UNITY Aneuploidy Screen	
Female for testing for genetic disease carrier status for procreative management	Z31.430	Supervision of elderly primigravida, first trimester	O09.511
Male for testing for genetic disease carrier status for procreative management	Z31.440	Supervision of elderly primigravida, second trimester	O09.512
Supervision of normal first pregnancy, unspecified trimester	Z34.00	Supervision of elderly multigravida, first trimester	O09.521
Supervision of normal first pregnancy, first trimester	Z34.01	Supervision of elderly multigravida, second trimester	O09.522
Supervision of normal first pregnancy, second trimester	Z34.02	Supervision of other high risk pregnancies, first trimester	O09.891
Supervision of other normal pregnancy, unspecified trimester	Z34.80	Supervision of other high risk pregnancies, second trimester	O09.892
Supervision of other normal pregnancy, first trimester	Z34.81	Abnormal ultrasonic finding on antenatal screening of mother	O28.3
Supervision of other normal pregnancy, second trimester	Z34.82	Abnormal chromosomal and genetic finding on antenatal screening of mother	O28.5
Supervision of normal pregnancy, unspecified, first trimester	Z34.91	Maternal care for (suspected) chromosomal abnormality in fetus	O35.1XX0
Family history of intellectual disabilities	Z81.0	Maternal care for (suspected) chromosomal abnormality in fetus 1	O35.1XX1
Family history of carrier genetic disease	Z84.81	Encounter for Rh incompatibility status	Z31.82
Family history of other specified conditions	Z84.89	Encounter for antenatal screening for chromosomal anomalies	Z36.0
		Family history of chromosomal abnormalities	Z82.79

PATIENT ACKNOWLEDGEMENT
Read and sign the front page.

I have been informed of and understand the details of the tests ordered herein for me by my healthcare provider, including the risks, benefits and alternatives, and consented to testing. I understand (1) the test results may inform me of a medical condition that may require follow-up and (2) a negative result does not rule out the possibility of such medical condition in the fetus, myself or my partner. I hereby authorize (1) the release to BillionToOne of any medical and insurance information necessary to process claims and recover reimbursement for services provided by BillionToOne and (2) BillionToOne to pursue all necessary appeals of any denials of payment in relation to services provided by BillionToOne. I understand that the test may not be (1) covered by my insurer/health plan, or (2) deemed medically necessary and I am responsible for any costs not paid by my plan directly to BillionToOne, including any copayments, deductibles or amounts deemed 'patient responsibility'. I acknowledge that I may be responsible for non-covered services. BillionToOne may (1) contact my healthcare provider to obtain more information regarding clinical correlation and confirmatory testing and (2) contact my provider or me for test status, billing/collection, quality assurance or research purposes.

BEFORE YOU SHIP, please ensure that:

✓

Test panel and ICD10 codes are selected

✓

Required fields on this form are completed

✓

Insurance card copies are included (front and back)

✓

Provided barcode is affixed to tubes and this form

✓

Requisition is signed

Call 1-800-463-3339 (1-800-GO FEDEX) to schedule a pickup

3200 Whipple Rd, Union City, CA 94587
E support@unityscreen.com
T 650.460.2551
F 1.833.915.0146
unityscreen.com
UN_TRF_085_2503
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