

PATIENT		SAMPLE		PROVIDER	
First Name	Jane	Sample Type	Blood	Provider	John Smith
Last Name	Doe	Date Collected	08/27/2022	Clinic Address	1035 O'Brien Drive Menlo Park, CA 94025
DOB	01/01/1984	Date Received	08/29/2022	Phone Number	650.460.2551
Ethnicity	African or African American	Accession ID	1234567K1234	Fax Number	833.915.0146
Gender	Female	Requisition ID	1234567K1234-5		
Gestational Age	19w3d	Date Reported	09/04/2022		
Medical Record #	001001				

UNITY™ Complete: Aneuploidy Screen

Singleton Gestation

SUMMARY OF RESULTS			
HIGH RISK Trisomy 21	MALE FETAL SEX	6.1% FETAL FRACTION	
CONDITIONS SCREENED	FETAL RISK <i>by</i> NIPT	RISK <i>Before</i> NIPT	RISK <i>After</i> NIPT
Trisomy 21	HIGH RISK	1 in 98	9 in 10
Trisomy 18	Low Risk	1 in 439	<1 in 10,000
Trisomy 13	Low Risk	1 in 1503	<1 in 10,000
Monosomy X	Low Risk	1 in 560	<1 in 10,000
Sex Chromosome Aneuploidy (XXX / XXY / XYY)	Not Detected		
RECOMMENDED FOLLOW-UP			
	PRENATAL DIAGNOSIS via chorionic villus sampling or amniocentesis is RECOMMENDED .		
	GENETIC COUNSELING is recommended to review the implications of this result. The patient may contact BillionToOne at (650) 460-2551 to schedule an appointment for a complimentary telephone genetic consultation to review these results. A genetic counselor can also be found at www.nsgc.org .		

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Patient Name Jane Doe	DOB 01/01/1983	Gestational Age 19w3d	Medical Record # 001001
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INTERPRETATION

UNITY™ Complete: Aneuploidy Screen Singleton Gestation

The fetus is HIGH RISK to be affected with TRISOMY 21. The estimated fetal fraction was 6.1%.

NIPT was performed to evaluate for fetal copy number of chromosomes 13, 18, 21, X & Y.

This NIPT result is valid only for a singleton gestation.

Prenatal diagnosis via chorionic villus sampling or amniocentesis is recommended. UNITY™ is not diagnostic. No irreversible decisions regarding the pregnancy should be made without confirmatory invasive prenatal testing. Genetic testing can also be performed postnatally.

Genetic counseling is recommended for this patient to review the implications of this result. The patient may contact BillionToOne at (650) 460-2551 to schedule an appointment for a complimentary telephone genetic consultation to review these results. Comprehensive genetic counseling is recommended for a patient with a family history of a chromosome abnormality or other genetic disorder so that risks can be accurately discussed, as well as additional testing options that may be available. A genetic counselor can also be found at www.nsgc.org.

METHODS AND LIMITATIONS

UNITY™ Complete: Aneuploidy Screen

Cell-free DNA (cfDNA) was isolated from 2-4 mL of plasma from whole blood collected in a Streck cell-free DNA tube. A paternal inheritance NIPT was performed as a multiplex PCR on common single nucleotide variants (SNVs) to measure the fraction of cell-free DNA of fetal origin. A modified fetal fraction calculation is performed for twin pregnancies and pregnancies conceived with a gestational carrier. The fetal risk for aneuploidy of chromosomes 13, 18, 21, X, & Y for singleton pregnancies or chromosomes 13, 18, & 21 for twin pregnancies, was determined by a separate multiplex PCR on cfDNA using no more than 99 amplicons per chromosome to perform relative chromosomal dosage analysis. When multiple blood tubes are analyzed for NIPT (e.g. for redraws), we report the combined reported fetal fraction by taking the arithmetic mean of fetal fractions across different tubes. Due to the tube-to-tube assay variability, the reported fetal fraction for the same patient can differ between single-gene NIPT and aneuploidy NIPT. Pre-test risks for aneuploidy ('Risk Before NIPT') are based on maternal age and gestational age. Post-test residual risks ('Risk After NIPT') for high-risk results are personalized and calculated based on the test sensitivity, test specificity, and prior risks, as determined by maternal age and gestational age. Post-test residual risks for low-risk results are not personalized and calculated based on the test sensitivity, test specificity, and prior-risk in the general population.

Test Limitations: Results may not be reported when the amount of cell-free fetal DNA in the blood sample is too low. Results from this test are highly accurate; however, discordant results may occur. Potential causes of discordant results include: maternal, fetal, or placental mosaicism, low fetal fraction, vanishing twin, maternal malignancy, maternal organ or bone marrow transplant, or other reasons. Findings of unknown significance will not be reported. UNITY™ does not screen for neural tube defects or abdominal wall defects. This test does not screen for microdeletions. UNITY™ NIPT is not diagnostic. No irreversible decisions regarding the pregnancy should be made without confirmatory invasive prenatal testing.

CONDITIONS SCREENED	SENSITIVITY	SPECIFICITY	POSITIVE PREDICTIVE VALUE	NEGATIVE PREDICTIVE VALUE
Trisomy 21	99.8% [98.9% - 100%]	99.9% [99.8% - 99.9%]	84.2%	> 99.9%
Trisomy 18	99.9% [99.0% - 100%]	> 99.9% [99.9% - 100%]	94.2%	> 99.9%
Trisomy 13	99.1% [97.4% - 100%]	> 99.9% [99.9% - 100%]	52.5%	> 99.9%
Monosomy X	97.7% [93.7% - 100%]	> 99.9% [99.9% - 100%]	85.5%	> 99.9%
Sex Chromosome Aneuploidy (XXX / XXY / XYY)	Reported when identified	> 99.9%	n/a	> 99.9%
Presence of Y Chromosome	> 99.9%	> 99.9%	n/a	> 99.9%

** Internal analytical validation data for singleton pregnancies. Performance metrics are estimated based on analytical data from contrived aneuploidy samples at 9% fetal fraction. Representative coefficients of variance from human plasma samples were applied to this analytical data to perform Monte Carlo simulations to calculate these metrics, assuming prior risks for a patient population of advanced maternal age. Mosaicism is not reflected in the validation.*

This NIPT was developed and its performance characteristics determined by the BillionToOne laboratory. It has not been cleared or approved by the U.S. Food and Drug Administration. The BillionToOne laboratory is regulated under CLIA. This test is used for clinical purposes. It should not be regarded as investigational or for research.

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Trisomy 21 - Patient Information Sheet

YOUR BABY'S RISK: HIGH CHANCE TO BE AFFECTED WITH TRISOMY 21

The testing performed by UNITY™ shows your baby's chance of being affected with trisomy 21 is **significantly increased**. Please reference your report to review personalized risk figures. **No irreversible pregnancy decisions, such as pregnancy termination, should be considered based on UNITY™ results alone.**

Follow-up testing and genetic counseling is recommended. You may contact BillionToOne at (650) 460-2551 to schedule an appointment for a complimentary telephone genetic consultation to review these results. A local genetic counselor can also be found at www.nsgc.org.

WHAT IS TRISOMY 21?

Individuals with trisomy 21, or Down syndrome, have three copies of chromosome 21. Children with trisomy 21 typically look like their parents, but also have some unique features such as a flat facial profile, small nose, and upward slanted eyes. In addition, they are at risk for various medical health issues. Approximately 50% of children have heart defects. Other issues may include hearing or vision impairment, feeding and digestion issues, and leukemia. Children with Down syndrome typically learn to walk and talk but are slower to achieve language and motor milestones than their siblings or peers. Almost all individuals with Down syndrome have some degree of intellectual disability.

Pregnancies affected with trisomy 21 may have anomalies identified on first or second trimester ultrasound. These pregnancies may have a higher risk for miscarriage or stillbirth.

Most health issues are treatable but may require surgery after birth. The average life expectancy is approximately 60 years. Most adults require support; however, some individuals may work or live in a supervised setting. The type and severity of symptoms varies from one person to another. Genetic testing cannot predict the severity or ultimate level of functioning.

WHAT CAUSES TRISOMY 21?

The average person has 46 chromosomes; males typically have an X and a Y chromosome, while females have two X chromosomes. An individual inherits one of each chromosome from each parent. An individual with trisomy 21 has an extra 21st chromosome. Trisomy 21 is not usually inherited. This is a mistake that happens at conception. There is nothing you could have done to cause or prevent trisomy 21.

CONFIRMATORY TESTING

Although UNITY™ is a highly accurate screening test, diagnostic testing is strongly recommended to determine whether your baby is affected with trisomy 21. UNITY™ uses an advanced technology to determine whether your baby's risk to have trisomy 21 is high or low. Confirmatory genetic testing during pregnancy or after birth can determine whether or not your baby is actually affected.

Before Birth

Testing during pregnancy can be performed by obtaining a sample of placenta (chorionic villus sampling or CVS) or fluid around your baby (amniocentesis) which both contain DNA that is representative of your baby. A CVS is ideally performed between 10 to 13 weeks pregnancy while an amniocentesis is performed between 15 to 24 weeks pregnancy. Both procedures are safe, but have a small risk for pregnancy complications, including miscarriage.

After Birth

Infants with Down syndrome typically have physical features of the condition. However, if confirmatory testing during pregnancy is declined, the baby's pediatrician should be informed of this result. An individual inherits one of each chromosome from each parent. An individual with trisomy 21 has an extra 21st chromosome. Trisomy 21 is not usually inherited. This is a mistake that happens at conception. There is nothing you could have done to cause or prevent trisomy 21.

RESOURCES

- National Down Syndrome Society - <https://www.ndss.org/>
- Down Syndrome Pregnancy - <https://www.downsyndromepregnancy.org>
- Global Down Syndrome Foundation - <https://www.globaldownsyndrome.org/>
- Lettercase, Understanding Down Syndrome - <https://understandingdownsyndrome.org/>
- National Society of Genetic Counselors - <https://www.nsgc.org>
- American College of Obstetricians and Gynecologists Guide to Prenatal Diagnosis - <https://www.acog.org/Patients/FAQs/Prenatal-Genetic-Diagnostic-Tests>