



Panorama: the next generation of NIPT

Screens for:

Singleton pregnancies

- Trisomies 21, 18, 13
- Monosomy X
- Triploidy
- Sex chromosome trisomies*
- 22q11.2 deletion syndrome (optional)
- Additional microdeletion syndromes (optional)
- Fetal sex (optional)

Twin pregnancies

- Zygosity
- Trisomies 21, 18, 13
- Fetal sex for each twin (optional)

If screening reveals monozygotic twins, Panorama can additionally screen for:

- Monosomy X
- Sex chromosome trisomies*
- 22q11.2 deletion syndrome (optional)

Egg donor or surrogate pregnancies

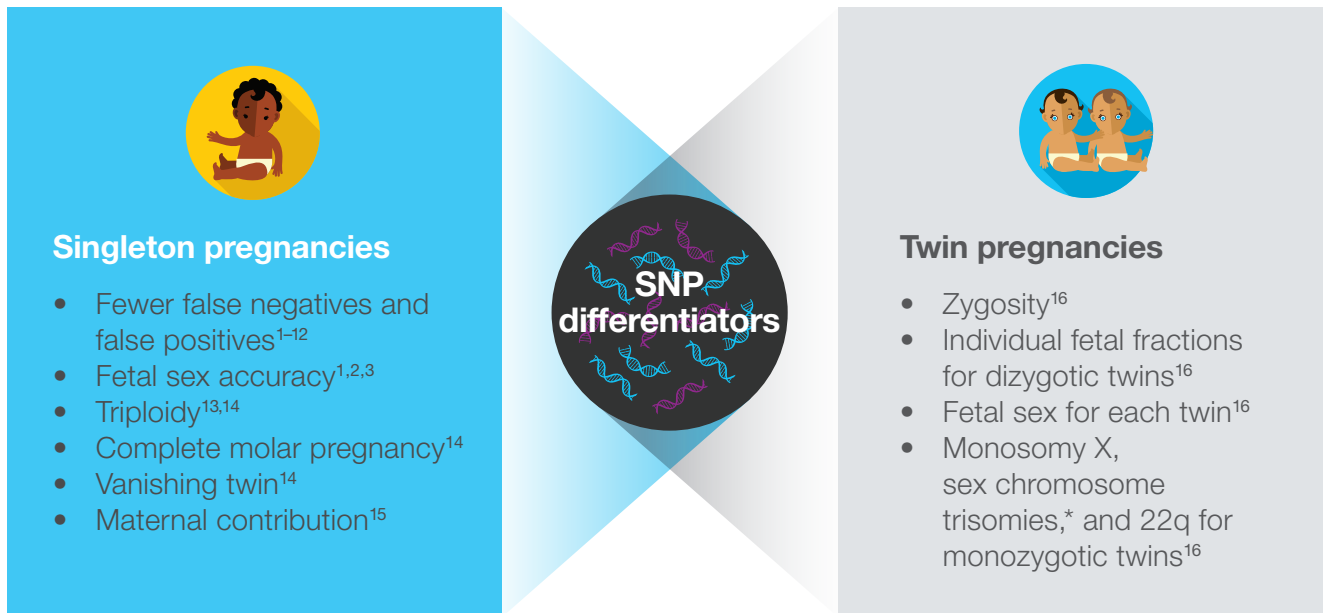
(Singleton pregnancies only)

- Trisomies 21, 18, 13
- Fetal sex (optional)

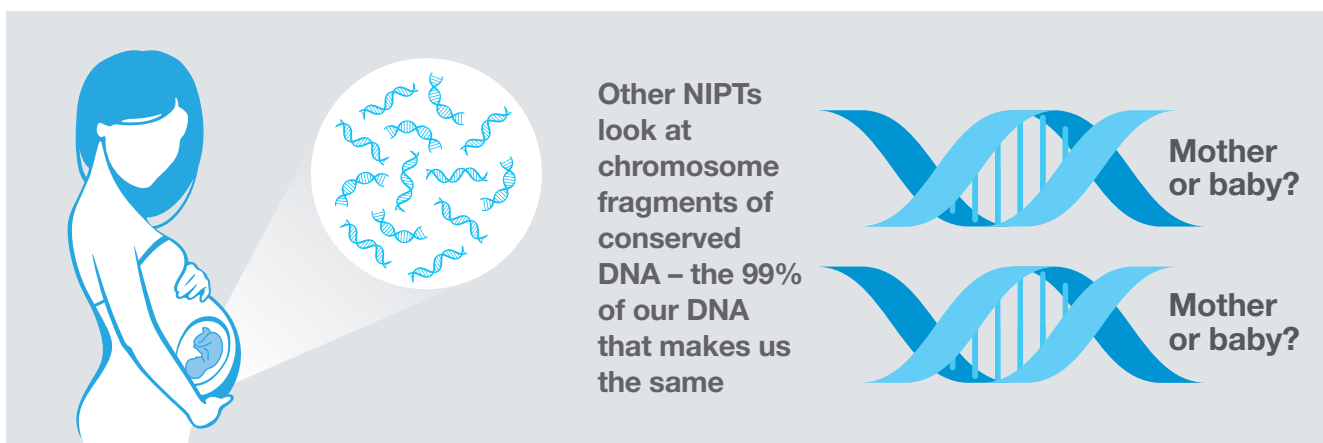
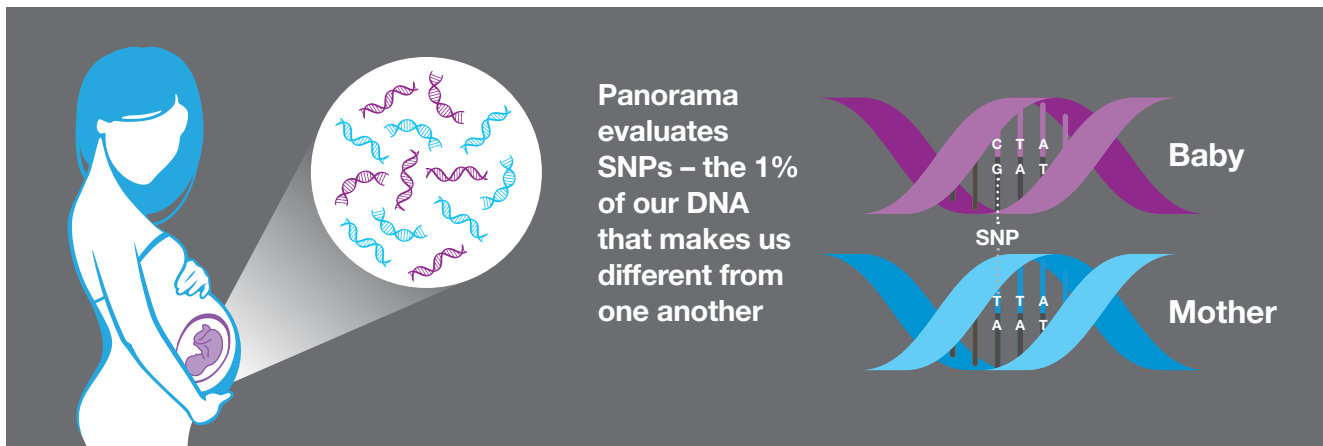
*Reported when suspected

Non-invasive prenatal screen

Panorama's unique SNP-based technology enables more comprehensive screening with greater accuracy in validation¹⁻¹⁶

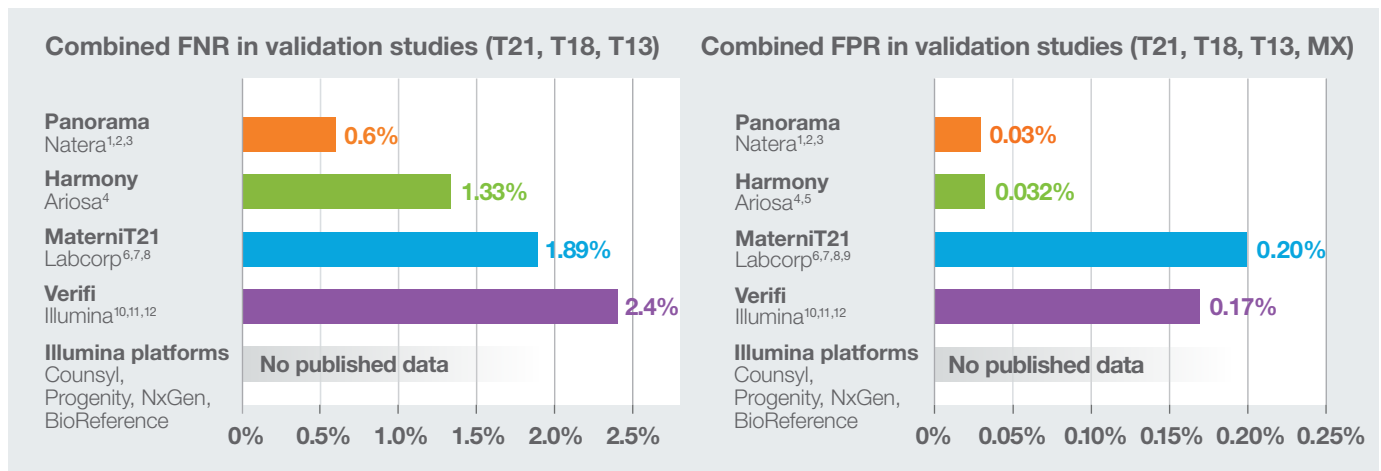


Only Panorama distinguishes between maternal and fetal (placental) DNA



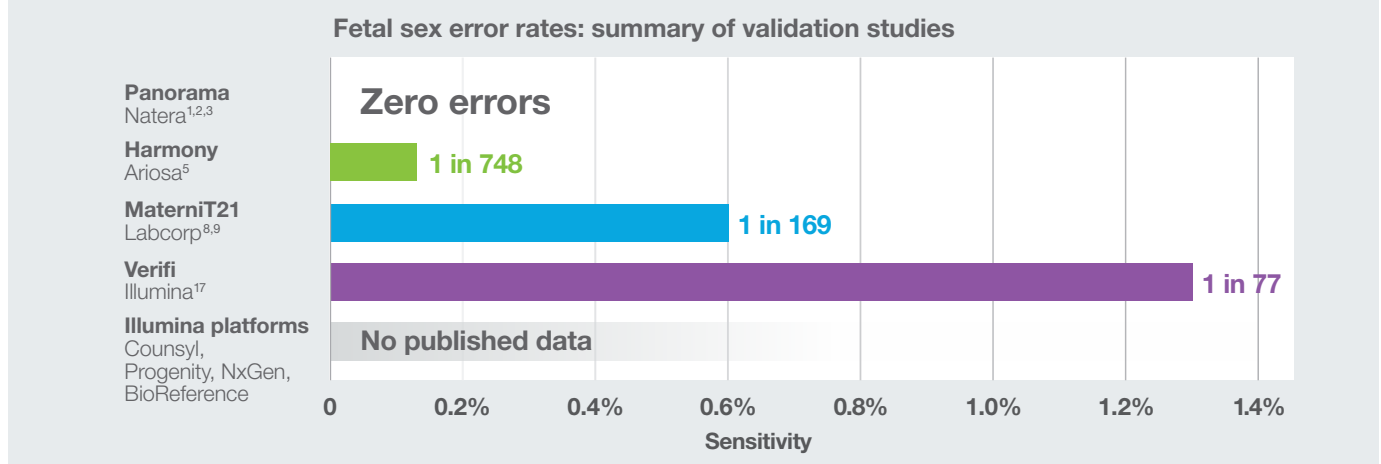
*Reported when suspected

Panorama reduces both false negative rates (FNR) and false positive rates (FPR) compared to other NIPTs



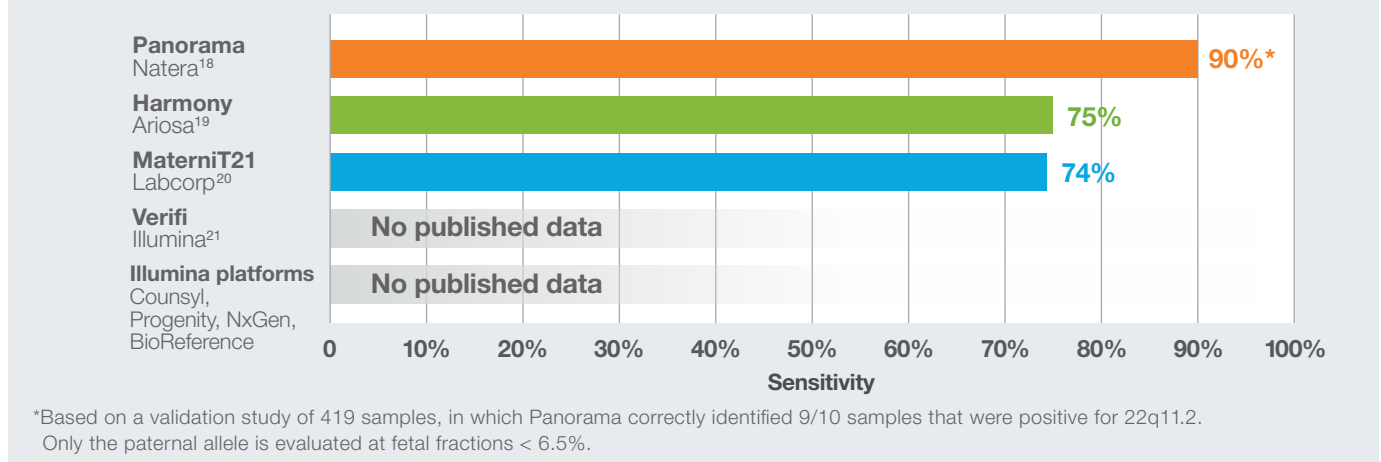
Panorama's SNP-based technology results in the highest fetal sex accuracy of any NIPT in validation studies

Other NIPTs may report incorrect gender for as many as 1 in 77 cases. A wrong call can lead to unnecessary clinical work-up and create anxiety for the patient.



Panorama's SNP-based approach yields the highest commercially available sensitivity for the most common ~3Mb 22q11.2 deletion

For small deletions like 22q11.2, Panorama's ability to evaluate unique DNA sequences within the region of interest enables better detection.



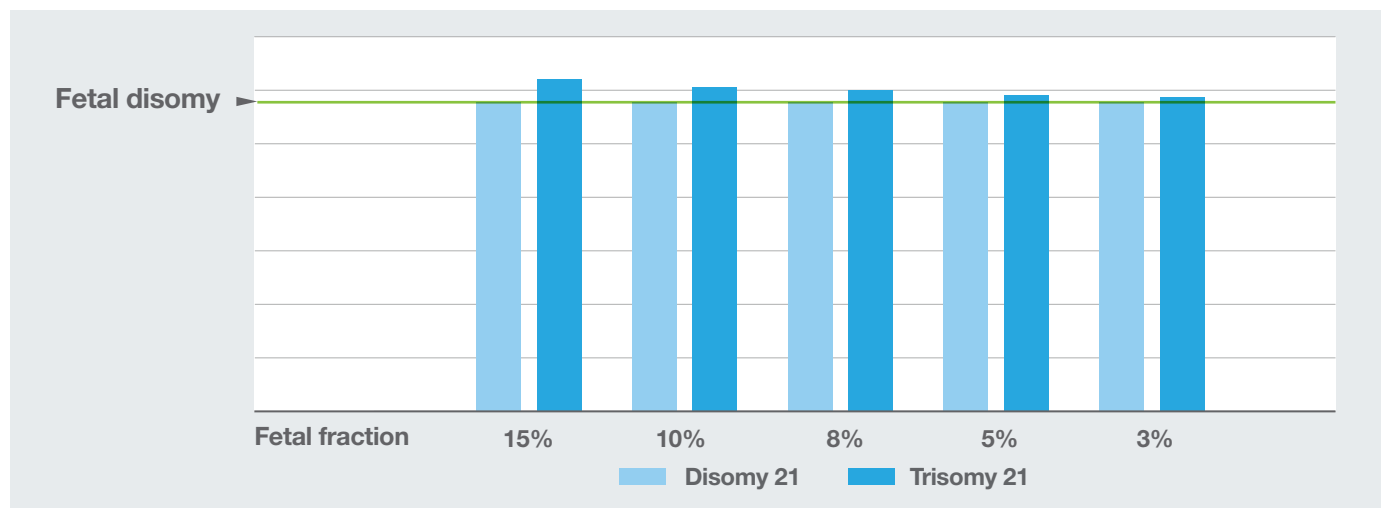
Accurate fetal fraction measurement is essential for accurate results²²

Panorama is the only NIPT that has always measured and reported fetal fraction

Panorama's SNP-based method is a gold standard in fetal fraction measurement

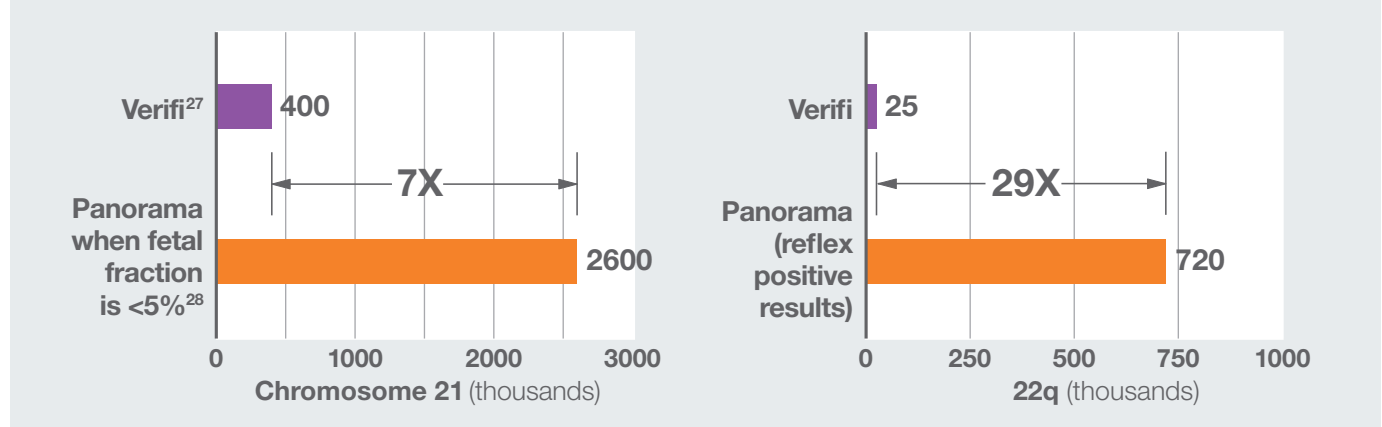
	Panorama ^{1,2,3}	Harmony ^{4,23}	MaterniT21 ^{5,6,7,24}	Verifi ^{10,11,12}	Illumina platforms
Method of fetal fraction measurement	13,392 SNPs	576 SNPs	Distribution of short (<150 bp) cfDNA	No data available on methodology or performance	No data available on methodology or performance
Combined false negative rate in validation studies (trisomies 21, 18, 13)	0.60%	1.33%	1.89%	2.40%	No published data

Counting methodologies' ability to detect abnormalities decreases below 8% fetal fraction, which may increase false negative results^{25,26}



Deeper sequencing on chromosomal regions of interest enables Panorama to maintain high-quality results at lower fetal fractions

Panorama's proprietary algorithm incorporates fetal fraction measurement and reflexes samples with lower fetal fraction to a higher depth of read.






Panorama for twin pregnancies

Only Panorama reports zygosity, individual fetal fractions for dizygotic twins, and fetal sex for each twin¹⁶

	Panorama ¹⁶	Harmony ^{29,30}	MaterniT21 ³¹	Verifi ³²	Illumina platforms ^{33,34,35,36}
Zygosity	✓	✗	✗	✗	✗
Individual fetal fractions	✓	✗	✗	✗	✗
Fetal sex for each twin	✓	✗	✗	✗	✗
Trisomies 21, 18, 13	✓	✓	✓	✓	✓
Monosomy X*	✓*	✗	✗	✗	✗
22q11.2 deletion syndrome*	✓*	✗	✓	✗	✗

*Available for monozygotic twins only

FINAL RESULTS SUMMARY: TWINS

<i>Result</i> LOW RISK 	<i>Zygosity</i> Dizygotic <i>FRATERNAL TWINS</i>	<i>Fetal Sex</i>  Male  Female	<i>Fetal Fraction(s)</i> 8.3%, 8.4%
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Notes by the clinical reviewer, if any, will be shown here.

Panorama allows clinicians to align their ultrasound findings with early and accurate zygosity information

Chorionicity is the strongest predictor for pregnancy complications in twins.³⁷ Studies have shown that up to 19% of monozygotic pregnancies are incorrectly classified as dichorionic using ultrasound.³⁸



Monozygotic

- Higher risk for twin-twin transfusion syndrome (TTTS), birth defects, etc. if monozygotic
- Consider early MFM referral to confirm chorionicity
- Develop tailored care plan for pregnancy



Dizygotic

- All dizygotic pregnancies are dichorionic
- Low risk for TTTS
- Continue standard care for pregnancy

Fetal fraction measurement and reporting in twins is important for reliable NIPT results



Fetal fraction is, on average, 30% higher in twin pregnancies, but fetal fraction per twin is lower compared to singleton pregnancies.²⁸

Provider support



Proactive billing outreach and price transparency



Complimentary mobile phlebotomy services and in-office phlebotomist for our testing*



Direct support from board-certified genetic counselors; call 650.249.9090 or email niptgc@natera.com for:

- Clinical questions
- Consultations on high-risk results
- Regional medical education and support



Our Natera Connect provider portal can be accessed online at connect.natera.com to:

- Order and track tests electronically
- Release low-risk results to patients (optional)

*Where permitted by state law

Patient support



Patients can learn about testing, book services, and track test results through our patient portal at my.natera.com



Complimentary mobile phlebotomy services can be scheduled by calling 855.271.1502 and pressing option 1



Complimentary pre- and post-test genetic information sessions with a board certified genetic counselor can be scheduled by calling 855.271.1502 and pressing option 2



Flexible payment plans, including assistance for financial hardship, are available

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The tests described have been developed and their performance characteristics determined by the CLIA-certified laboratory performing the tests. These tests have not been cleared or approved by the U.S. Food and Drug Administration (FDA). Although FDA does not currently clear or approve laboratory-developed tests in the U.S., certification of the laboratory is required under CLIA to ensure the quality and validity of the tests. © 2018 Natera, Inc. All Rights Reserved. PAN_MD_BR_2018_02_12_NAT-801513



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