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Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review)

Badeau M, Lindsay C, Blais J, Nshimyumukiza L, Takwoingi Y, Langlois S, Légaré F, Giguère Y, Turgeon AF, Witteman W, Rousseau F

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[Diagnostic Test Accuracy Review]

Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women

Mylène Badeau¹, Carmen Lindsay¹, Jonatan Blais^{2,3}, Leon Nshimyumukiza⁴, Yemisi Takwoingi⁵, Sylvie Langlois⁶, France Légaré¹, Yves Giguère^{7,8}, Alexis F Turgeon^{9,10}, William Witteman¹, François Rousseau^{8,10}

¹Population Health and Optimal Health Practices Research Axis, CHU de Québec - Université Laval, Québec City, Canada. ²Department of Medical Biology, CHAU-Hôtel-Dieu de Lévis, Lévis, Canada. ³Department of Molecular Biology, Medical Biochemistry and Pathology, Faculty of Medicine, Université Laval, Quebec City, Canada. ⁴School of Public Health, University of Alberta, Edmonton, Canada. ⁵Institute of Applied Health Research, University of Birmingham, Birmingham, UK. ⁶Department of Medical Genetics, Faculty of Medicine, University of British Columbia, Vancouver, Canada. ⁷Reproductive, Mother and Child Health Research Axis, CHU de Québec - Université Laval, Québec City, Canada. ⁸Department of Molecular Biology, Medical Biochemistry and Pathology, Faculty of Medicine, Université Laval, Québec City, Canada. ⁹Department of Anesthesiology and Critical Care Medicine, Division of Critical Care Medicine, and Population Health and Optimal Health Practices Research Unit, CHU de Québec - Université Laval, Research Center, CHU de Québec - Université Laval, Université Laval, Quebec City, Canada. ¹⁰Population Health and Optimal Health Practices Research Axis, CHU de Québec Research Center, Université Laval, Québec City, Canada.

Contact address: François Rousseau, Population Health and Optimal Health Practices Research Axis, CHU de Québec Research Center, Université Laval, 1401, 18eme rue, Québec City, Québec, G1J 1Z4, Canada. francois.rousseau@mac.com, francois.rousseau@fmed.ulaval.ca.

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ABSTRACT

Background

Common fetal aneuploidies include Down syndrome (trisomy 21 or T21), Edward syndrome (trisomy 18 or T18), Patau syndrome (trisomy 13 or T13), Turner syndrome (45,X), Klinefelter syndrome (47,XXY), Triple X syndrome (47,XXX) and 47,XYY syndrome (47,XYY). Prenatal screening for fetal aneuploidies is standard care in many countries, but current biochemical and ultrasound tests have high false negative and false positive rates. The discovery of fetal circulating cell-free DNA (ccfDNA) in maternal blood offers the potential for genomics-based non-invasive prenatal testing (gNIPT) as a more accurate screening method. Two approaches used for gNIPT are massively parallel shotgun sequencing (MPSS) and targeted massively parallel sequencing (TMPS).

Objectives

To evaluate and compare the diagnostic accuracy of MPSS and TMPS for gNIPT as a first-tier test in unselected populations of pregnant women undergoing aneuploidy screening or as a second-tier test in pregnant women considered to be high risk after first-tier screening for common fetal aneuploidies. The gNIPT results were confirmed by a reference standard such as fetal karyotype or neonatal clinical examination.

Search methods

We searched 13 databases (including MEDLINE, Embase and Web of Science) from 1 January 2007 to 12 July 2016 without any language, search filter or publication type restrictions. We also screened reference lists of relevant full-text articles, websites of private prenatal diagnosis companies and conference abstracts.

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Selection criteria

Studies could include pregnant women of any age, ethnicity and gestational age with singleton or multifetal pregnancy. The women must have had a screening test for fetal aneuploidy by MPSS or TMPS and a reference standard such as fetal karyotype or medical records from birth.

Data collection and analysis

Two review authors independently carried out study selection, data extraction and quality assessment (using the QUADAS-2 tool). Where possible, hierarchical models or simpler alternatives were used for meta-analysis.

Main results

Sixty-five studies of 86,139 pregnant women (3141 aneuploids and 82,998 euploids) were included. No study was judged to be at low risk of bias across the four domains of the QUADAS-2 tool but applicability concerns were generally low. Of the 65 studies, 42 enrolled pregnant women at high risk, five recruited an unselected population and 18 recruited cohorts with a mix of prior risk of fetal aneuploidy. Among the 65 studies, 44 evaluated MPSS and 21 evaluated TMPS; of these, five studies also compared gNIPT with a traditional screening test (biochemical, ultrasound or both). Forty-six out of 65 studies (71%) reported gNIPT assay failure rate, which ranged between 0% and 25% for MPSS, and between 0.8% and 7.5% for TMPS.

In the population of unselected pregnant women, MPSS was evaluated by only one study; the study assessed T21, T18 and T13. TMPS was assessed for T21 in four studies involving unselected cohorts; three of the studies also assessed T18 and 13. In pooled analyses (88 T21 cases, 22 T18 cases, eight T13 cases and 20,649 unaffected pregnancies (non T21, T18 and T13)), the clinical sensitivity (95% confidence interval (CI)) of TMPS was 99.2% (78.2% to 100%), 90.9% (70.0% to 97.7%) and 65.1% (9.16% to 97.2%) for T21, T18 and T13, respectively. The corresponding clinical specificity was above 99.9% for T21, T18 and T13.

In high-risk populations, MPSS was assessed for T21, T18, T13 and 45,X in 30, 28, 20 and 12 studies, respectively. In pooled analyses (1048 T21 cases, 332 T18 cases, 128 T13 cases and 15,797 unaffected pregnancies), the clinical sensitivity (95% confidence interval (CI)) of MPSS was 99.7% (98.0% to 100%), 97.8% (92.5% to 99.4%), 95.8% (86.1% to 98.9%) and 91.7% (78.3% to 97.1%) for T21, T18, T13 and 45,X, respectively. The corresponding clinical specificities (95% CI) were 99.9% (99.8% to 100%), 99.8% to 100%), 99.8% (99.8% to 99.9%) and 99.6% (98.9% to 99.8%). In this risk group, TMPS was assessed for T21, T18, T13 and 45,X in six, five, two and four studies. In pooled analyses (246 T21 cases, 112 T18 cases, 20 T13 cases and 4282 unaffected pregnancies), the clinical sensitivity (95% CI) of TMPS was 99.2% (96.8% to 99.8%), 98.2% (93.1% to 99.6%), 100% (83.9% to 100%) and 92.4% (84.1% to 96.5%) for T21, T18, T13 and 45,X respectively. The clinical specificities were above 100% for T21, T18 and T13 and 99.8% (98.3% to 100%) for 45,X. Indirect comparisons of MPSS and TMPS for T21, T18 and 45,X showed no statistical difference in clinical sensitivity, clinical specificity or both. Due to limited data, comparative meta-analysis of MPSS and TMPS was not possible for T13.

We were unable to perform meta-analyses of gNIPT for 47,XXX, 47,XXY and 47,XYY because there were very few or no studies in one or more risk groups.

Authors' conclusions

These results show that MPSS and TMPS perform similarly in terms of clinical sensitivity and specificity for the detection of fetal T31, T18, T13 and sex chromosome aneuploidy (SCA). However, no study compared the two approaches head-to-head in the same cohort of patients. The accuracy of gNIPT as a prenatal screening test has been mainly evaluated as a second-tier screening test to identify pregnancies at very low risk of fetal aneuploidies (T21, T18 and T13), thus avoiding invasive procedures. Genomics-based non-invasive prenatal testing methods appear to be sensitive and highly specific for detection of fetal trisomies 21, 18 and 13 in high-risk populations. There is paucity of data on the accuracy of gNIPT as a first-tier aneuploidy screening test in a population of unselected pregnant women. With respect to the replacement of invasive tests, the performance of gNIPT observed in this review is not sufficient to replace current invasive diagnostic tests.

We conclude that given the current data on the performance of gNIPT, invasive fetal karyotyping is still the required diagnostic approach to confirm the presence of a chromosomal abnormality prior to making irreversible decisions relative to the pregnancy outcome. However, most of the gNIPT studies were prone to bias, especially in terms of the selection of participants.

PLAIN LANGUAGE SUMMARY

Accuracy of gNIPT for identifying genetic abnormalities in unborn babies

What is the issue?

How accurate is the new test (genomics-based non-invasive prenatal testing (gNIPT)) for detecting abnormal chromosome number in an unborn baby's genetic material (DNA) found in the mother's blood? We assessed the accuracy for the screening of Down syndrome (trisomy 21), Edward syndrome (trisomy 18), Patau syndrome (trisomy 13), Turner syndrome (45,X), Klinefelter syndrome (47,XXY), Triple X syndrome (47,XXX) and 47,XYY syndrome. There are different methods in use for gNIPT. We assessed MPSS (massively parallel shotgun sequencing) that tests whole DNA and TMPS (targeted massively parallel sequencing) that tests targeted DNA.



Background

There are 46 chromosomes (23 pairs) in humans. Abnormal numbers of chromosomes can cause genetic disorders for which there are no cures. Having an extra chromosome is called trisomy and an excess (or less) of sexual chromosome is called sex chromosome abnormality (SCA). The most common trisomy is Down syndrome which occurs in about one in 1000 babies. Children with Downs have slow growth, characteristic facial features and mild to moderate intellectual disability, with some requiring specialist education later in life. However, the symptoms vary from mild to severe so that some infants lead relatively normal lives. The other trisomy or SCA conditions have varying degrees of disability but the chance of a baby being affected is much less.

Current screening tests for these conditions require confirmation if the baby has the condition or not and for this an invasive test like amniocentesis is used. Amniocentesis is where fetal cells that float in the fluid surrounding the unborn baby are collected by putting a fine needle through the mother's abdomen and collecting the fluid. Alternatively, tissue can be collected from the placenta (chorionic villus sampling (CVS)). With these invasive tests, pregnant women are exposed to a higher chance of losing their baby even if the baby is unaffected by Down syndrome. So, this invasive test is only offered to women who are thought to have a higher chance of having an affected unborn baby

What we did

We looked for studies that included women of any age, ethnicity and gestational age who were carrying either a single baby or more than one. We searched for studies (up to July 2016) that assessed the accuracy of the new test.

What we found

We found 65 studies with a total of 86,139 pregnant women, including 3141 affected pregnancies. Forty-two studies (65%) enrolled pregnant women with a high chance of having babies with abnormal chromosome number. Forty-eight (74%) studies included only women with a singleton pregnancy. Forty-four studies (68%) used MPSS and 21 studies (32%) used TMPS.

gNIPT seems to be accurate for screening unborn babies (either singletons or twins), especially for detecting Down syndrome, trisomy 18 and trisomy 13. However, there were some problems with how the studies were conducted which makes us cautious about our findings. This may result in gNIPT appearing to perform better than it really does.

Other important information to consider

gNIPT method appears to perform well in identifying unborn babies with abnormal number of chromosomes. However, when a gNIPT detects an abnormal chromosome number, then a confirmation using invasive tests (like amniocentesis or CVS) is still needed before pregnancy-related decisions can be made.

It is important that pregnant women are given full information on the possible health problems that might arise for babies affected by an additional chromosome. For example, with Down syndrome though some children have considerable disability, others can lead relatively normal lives. In addition, in this review most studies enrolled pregnant women with increased chance of having babies with abnormal chromosome number, so our findings do not directly apply to general populations of pregnant women.

SUMMARY OF FINDINGS

Summary characteristics of incl	uded studies
Review question	What is the diagnostic accuracy of massively parallel shotgun sequencing (MPSS) and targeted massively parallel sequencing (TMPS) using circulating cell-free DNA (ccfDNA) in maternal blood for the detection of common fetal aneuploidies (T21, T18, T13, 45,X, 47,XXY, 47,XXX and 47,XYY) in pregnant women according to their prior risk of fetal aneuploidy?
Importance (rationale)	These new genomics-based non-invasive prenatal testing (gNIPT) approach report higher sensitivity and lower false positive rate than traditional screening tests. gNIPT is already advertised and marketed. How gNIPT should be used in clinical practice should be assessed in order to provide a framework for its use.
Study design	There were 40 prospective cohort studies, 8 retrospective cohort studies, 16 case-control studies and 1 prospective and retro- spective cohort study.
Population	Pregnant women of any age, ethnicity and gestational age, with singleton or multifetal pregnancy who had a screening test for fetal aneuploidy using gNIPT and received a reference standard. 42 studies enrolled pregnant women selected at high risk of fe- tal aneuploidy, 5 enrolled unselected pregnant women undergoing aneuploidy screening and 18 enrolled pregnant women from a mixed-risk population of fetal aneuploidy. 48 studies included only women with singleton pregnancy, 5 included only multife- tal pregnancies, 4 included either type of pregnancy and 8 did not report type of pregnancy. 10 studies included only women in the first trimester (15 weeks or less), 21 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters), 24 studies included women in the first 2 trimesters (29 weeks or less), 24 studies included women in the first 2 trimesters) age.
Index tests	gNIPT by MPSS (44 studies) or TMPS (21 studies), including 5 studies that compared a gNIPT with a traditional screening test. 37 studies were industry-funded or were written by 1 or more authors affiliated with a company who sells gNIPT. 22 studies were not reported to be funded by industry but samples were sequenced and analysed by a commercial laboratory. 3 studies had no links with industry.
Target conditions	36 studies reported results for only autosomes (T21, T18, T13), 4 for only SCA (45,X, 47,XXY, 47,XXX and 47,XYY), and 25 for both autosomes and SCA.
Reference standard	Fetal karyotyping performed on cells obtained from chorionic villi sampling, amniotic fluid, placental tissue, a fetus lost by mis- carriage or other equivalent and recognised methods on the same materials for autosomes and SCA. If fetal karyotyping was not performed, we used neonatal clinical examination or medical records from birth (for autosomes only). Only 1 reference standard was used for all pregnant women included in 36 studies while multiple reference standards were used in 29 studies.
Risk of bias	The QUality Assessment of Diagnostic Accuracy Studies (QUADAS-2) tool was used to assess the methodological quality of in- cluded studies.
	No study was assessed as being at low risk of bias across all domains. For the patient selection domain, no study was assessed as being at low risk of bias. For the index test, reference standard and flow and timing domains, the risk of bias was low for 94%, 77% and 23% of studies, respectively.

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Applicability concerns

Applicability was of low concern for all studies in the index test and reference standard domains because the studies matched the review question. In the patient selection domain, 47 (71%) studies were judged to be of low applicability concern because they included pregnant women matching the review question.

45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, T21: trisomy 21, T18: trisomy 18, T13: trisomy 13.

Summary of findings 2. Performance of gNIPT for detection of T21

Test strategy	Number of	Number of affected	Sensitivity	Specificity	Median	Missed	False
	studies	pregnancies (Number of	% (95% CI)	% (95% CI)	prevalence ^b	cases	positives
		unaffected pregnan- cies) ^a				(FN)¢	(FP) ^d
Unselected pro	egnant women						
MPSS	1	8 (1733)	100 (67.6 to 100)	100 (99.8 to 100)	0.46	0	0
TMPS	4	88 (20,679)	99.2 (78.2 to 100)	100 (> 99.9 to 100)	(0.24 to 5.21)	4	0
Tradition- al screening test ^e	1	38 (15,803)	78.9 (63.7 to 88.9)	94.6 (94.2 to 94.9)		97	5375
Implications	MPSS will cwith TMPS,	000 pregnancies expected to letect all cases and no pregna , 4 cases will be missed and n onal screening tests, 363 cas	ant woman will undergo a o pregnant woman will ur	ndergo unnecessary invasive	test; and	cessary invasive	test.
Selected high-	risk pregnant w	omen					
MPSS	30	1048 (15,937)	99.7 (98.0 to 100)	99.9 (99.8 to 100)	4.95	15	95
TMPS	6	246 (4380)	99.2 (96.8 to 99.8)	100 (99.8 to 100)	(0.44 to 27.66)	40	0

-0.03 (-0.11 to 0.04)

NA

0.53 (-0.73 to 1.78)

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Difference between MPSS and TMPS

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for detection of fetal chromosomal aneuploidy in pregnant women (Review)

- Implications 4950 of 100,000 pregnancies expected to be affected by T21;
 - 4936 and 4911 cases will be detected while 15 and 40 cases will be missed by MPSS and TMPS, respectively; and
 - of 95,050 expected pregnancies unaffected by T21, 95 and 0 pregnant women will undergo unnecessary invasive tests with MPSS and TMPS, respectively.

MPSS: massively parallel shotgun sequencing, NA; not applicable, TMPS: targeted massively parallel sequencing, T21: trisomy 21.

^aUnaffected pregnancies: we included patients with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies.

^bThe median prevalence and range were calculated by using all prospective or retrospective studies for each category considered.

^cMissed cases per 100,000 tested. FN: false negatives.

^{*d*}False positives per 100,000 tested. A false positive result may lead to unnecessary invasive tests depending on choices by the pregnant woman.

eTraditional screening tests are first-trimester combined test, second-trimester quadruple test, second-trimester fully integrated test, second-trimester sequential test or second-trimester triple test.

Summary of findings 3. Performance of gNIPT for detection of T18

Performance of gNIPT for detection of T18 Number of affected Sensitivity Specificity Median Missed False Test strategy Number of studies pregnancies (Number % (95% CI) % (95% CI) prevalenceb cases positives of % (range) (FN)c (FP)d unaffected pregnancies)a Unselected pregnant women MPSS 1 2 (1739) 100 (34.3 to 100) 99.9 (99.7 to 100) 0.11 0 100 (0.06 to 0.36) TMPS 3 22 (20,553) 90.9 (70.0 to 97.7) 100 (99.9 to 100) 10 0 1 22 Tradition-10 (15,831) 80.0 (49.0 to 94.3) 99.7 (99.6 to 99.8) 300 al screening teste Implications • 109 of 100,000 pregnancies expected to be affected by T18;

• MPSS will detect all cases and 100 unaffected pregnant women will undergo an unnecessary invasive test;

• with TMPS, 10 cases will be missed and no unaffected pregnant woman will undergo unnecessary invasive test; and

• with traditional screening tests, 87 cases will be detected, 22 will be missed and 300 unaffected pregnant women will undergo unnecessary invasive test.

Selected high-risk pregnant women

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IPSS	28	332 (16,180)	97.8 (92.5 to 99.4)	99.9 (99.8 to 100)	1.46	32	99	
TMPS	5 112 (4010)		98.2 (93.1 to 99.6) 100 (99.8 to 100)		(0.22 to 17.02)	26	0	
Difference be- tween MPSS and TMPS			-0.41 (-4.11 to 3.28)	-0.06 (-0.14 to 0.03)	NA			
Implications	• 1431 and 14		hile 32 and 26 cases will b	e missed by MPSS and TMPS will undergo unnecessary in		and TMPS, respe	ctively.	
Traditional scree	ening tests are fir	d. A false positive result may st-trimester combined test, s		le test, second-trimester fully			uential test or second-	
False positives p Traditional scree rimester triple to Summary of fin Performance o	ening tests are fir est. ndings 4. Perf	st-trimester combined test, s formance of gNIPT for de ection of T13	econd-trimester quadrup	ole test, second-trimester fully	y integrated test, secon	ıd-trimester seqı		
False positives p Traditional scree rimester triple te Summary of fi	ening tests are fir est. ndings 4. Perf	st-trimester combined test, s formance of gNIPT for de ection of T13 Number of affected pregnancies (Number	econd-trimester quadrup				uential test or second- False positives	
False positives p Traditional scree rimester triple to Summary of fin Performance o	ening tests are fir est. ndings 4. Perf of gNIPT for dete Number of	formance of gNIPT for de ection of T13 Number of affected	econd-trimester quadrup etection of T13 Sensitivity %	ole test, second-trimester fully Specificity %	y integrated test, secon Median	nd-trimester sequ	False	
False positives p Traditional scree rimester triple to Summary of fin Performance o	ening tests are fir est. ndings 4. Perf of gNIPT for dete Number of studies	formance of gNIPT for de ection of T13 Number of affected pregnancies (Number of unaffected pregnan-	econd-trimester quadrup etection of T13 Sensitivity %	ole test, second-trimester fully Specificity %	y integrated test, secon Median prevalence ^b	nd-trimester sequ Missed cases	False positives	
False positives p Traditional scree rimester triple to Summary of fi Performance o Test strategy	ening tests are fir est. ndings 4. Perf of gNIPT for dete Number of studies	formance of gNIPT for de ection of T13 Number of affected pregnancies (Number of unaffected pregnan-	econd-trimester quadrup etection of T13 Sensitivity %	ole test, second-trimester fully Specificity %	y integrated test, secon Median prevalence ^b	nd-trimester sequ Missed cases	False positives	
False positives p Traditional scree rimester triple to Summary of fi Performance o Test strategy Unselected pre	ening tests are fir est. ndings 4. Perf of gNIPT for dete Number of studies	formance of gNIPT for de ection of T13 Number of affected pregnancies (Number of unaffected pregnan- cies) ^a	econd-trimester quadrup etection of T13 Sensitivity % (95% CI)	Specificity % (95% CI)	y integrated test, secon Median prevalence ^b % (range)	Missed cases (FN) ^c	False positives (FP) ^d	

- Implications 118 of 100,000 pregnancies expected to be affected by T13;
 - MPSS will detect all cases and no unaffected pregnant woman will undergo an unnecessary invasive test;
 - with TMPS, 41 cases will be missed and no unaffected pregnant woman will undergo unnecessary invasive test; and
 - with traditional screening tests, 59 cases will be missed and 300 unaffected pregnant women will undergo unnecessary invasive test.

Selected hig	h-risk pregnant v	vomen						
MPSS	20	128 (13,810)	95.8 (86.1 to 98.9)	99.8 (99.8 to 99.9)	1.09	46	198	
TMPS	2	20 (293)	100 (83.9 to 100) ^{<i>f</i>}	100 (98.7 to 100) ^{<i>f</i>}	(0.04 to 3.54)	0	0	

- Implications 1087 of 100,000 pregnancies expected to be affected by T13;
 - 1041 and 1087 cases will be detected while 46 and 0 cases will be missed by MPSS and TMPS, respectively; and
 - of 98,913 expected unaffected by T13, 198 and 0 pregnant women will undergo unnecessary invasive test with MPSS and TMPS, respectively.

MPSS: massively parallel shotgun sequencing, NA: not applicable, TMPS: targeted massively parallel sequencing, T13: trisomy 13.

^aUnaffected pregnancies: we included patients with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies.

^bThe median prevalence and range were calculated by using all prospective or retrospective studies for each category considered.

^cMissed cases per 100,000 tested. FN: false negatives.

^dFalse positives per 100,000 tested. A false positive result may lead to unnecessary invasive tests depending on choices by the pregnant woman.

eTraditional screening tests are first-trimester combined test, second-trimester quadruple test, second-trimester fully integrated test, second-trimester sequential test or second-trimester triple test.

^fSimple pooling used to obtain summary estimates of sensitivity, specificity or both.

Summary of findings 5. Performance of gNIPT for detection of 45,X

Performance of gNIPT for detection of 45,X

Test	strategy	Number of	Number of affected	Sensitivity	Specificity	Median	Missed	False
		studies	pregnancies (Number of	% (95% CI)	% (95% CI)	prevalence ^b	cases	positives
			unaffected pregnancies) ^a			% (range)	(FN)¢	(FP) ^d
Seleo	cted high-r	risk pregnant w	omen					
MPSS	S	12	119 (7440)	91.7 (78.3 to 97.1)	99.6 (98.9 to 99.8)	1.04	86	396
TMPS	S	4	79 (985)	92.4 (84.1 to 96.5)	99.8 (98.3 to 100)	(0.27 to 18.58)	79	198
Diffe	rence betw	veen MPSS and T	MPS	-0.74 (-11.1 to 9.60)	-0.23 (-0.82 to 0.36)	NA		

(Review)

fetal chromosomal aneuploidy in pregnant women

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Genomics-based non-invasive prenatal testing for detection of

- 953 and 960 cases will be detected while 86 and 79 cases will be missed by MPSS and TMPS, respectively; and
- of 98,961 expected unaffected by 45X, 396 and 198 pregnant women will undergo unnecessary invasive test with MPSS and TMPS, respectively.

45,X: Turner syndrome, MPSS: massively parallel shotgun sequencing, NA: not applicable, TMPS: targeted massively parallel sequencing. ^aUnaffected pregnancies: we included patients with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies. ^bThe median prevalence and range were calculated by using all prospective or retrospective studies for each category considered.

^cMissed cases per 100,000 tested. FN: false negatives.

^dFalse positives per 100,000 tested. A false positive result may lead to unnecessary invasive tests depending on choices by the pregnant woman.

Summary of findings 6. Performance of gNIPT for detection of autosomes aneuploidies (T21, T18 and T13 combined)

Performance o	f gNIPT for dete	ction of autosomes aneuploid	lies (T21, T18 and T13 co	ombined)			
Test strategy	Number of	Number of affected	Sensitivity	Specificity	Median	Missed	False
	studies	pregnancies (Number of	% (95% CI)	% (95% CI)	prevalence ^b	cases	positives
		unaffected pregnancies) ^a			% (range)	(FN) ^c	(FP) ^d
Unselected pre	egnant women						
MPSS	1	11 (1730)	100 (74.1 to 100)	99.9 (99.7 to 100)	0,63	0	99
TMPS	4	118 (20,649)	94.9 (89.1 to 97.7)	99.9 (99.8 to 99.9)	(0.32 to 5.73)	32	99
Tradition- al screening test ^e	4	120 (22,247)	ND ^f			ND	
Implications	• 632 and 600	000 pregnancies expected to be) cases will be detected wherea naffected, 99 pregnant women	s 0 and 32 cases will be m	issed by MPSS and TMPS,			
Selected high-	risk pregnant w	omen					
MPSS	32	1508 (15,797)	98.8 (97.2 to 99.5)	99.9 (99.7 to 100)	5.85	70	94
TMPS	7	378 (4282)	98.9 (97.2 to 99.6)	99.9 (99.8 to 100)	(0.67 to 46.81)	64	94
Difference betw	veen MPSS and TI	MPS	-0.11	-0.08	NA		

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Implications • 5851 of 100,000 pregnancies expected to be affected by T21, T18 or T3;

• 5781 and 5787 cases will be detected, whereas 70 and 64 cases will be missed by MPSS and TMPS, respectively; and

(-1.58 to 1.35)

• of 94,149 unaffected, 94 pregnant women will undergo unnecessary invasive test with MPSS or TMPS.

MPSS: massively parallel shotgun sequencing, NA: not applicable, ND: no data available, TMPS: targeted massively parallel sequencing, T13: trisomy 13, T18: trisomy 18, T21: trisomy 21.

(-0.22 to 0.07)

^{*a*}Unaffected pregnancies: we included patients with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies.

^bThe median prevalence and range were calculated by using all prospective or retrospective studies for each category considered.

^cMissed cases per 100,000 tested. FN: false negatives.

^dFalse positives per 100,000 tested. A false positive result may lead to unnecessary invasive tests depending on choices by the pregnant woman.

^eTraditional screening tests are first-trimester combined test, second-trimester quadruple test, second-trimester fully integrated test, second-trimester sequential test or second-trimester triple test.

^fSummary sensitivity and specificity were not obtained for traditional screening tests because the four studies used different cut-offs to determine test positivity. Three of the four studies compared TMPS and traditional screening tests in the same population (direct comparison).

Summary of findings 7. Performance of gNIPT for detection of sex chromosome aneuploidies (45,X, 47,XXX, 47,XXY and 47,XYY combined)^a

Performance of gNIPT for detection of sex chromosome aneuploidies (45,X, 47,XXX, 47,XXY and 47,XYY combined)

	Test strategy	Number of	Number of affected	Sensitivity	Specificity	Median	Missed	False
•		studies	pregnancies (Number of	% (95% CI)	% (95% CI)	prevalence ^c	cases	positives
			unaffected pregnancies) ^b			% (range)	(FN) ^d	(FP) ^e
	Selected high-ı	risk pregnant wo	men					
	MPSS	12	151 (7452)	91.9 (73.8 to 97.9)	99.5 (98.8 to 99.8)	1.53	124	492
	TMPS	4	96 (968)	93.8 (86.8 to 97.2)	99.6 (98.1 to 99.9)	(0.45 to 18.58)	95	394
i	Difference betw	een MPSS and TM	IPS	-1.85 (-13.3 to 9.60)	-0.06 (-0.82 to 0.71)	NA		

Implications • 1535 of 100,000 pregnancies expected to be affected by SCA;

• 1411 and 1440 cases will be detected while 124 and 95 cases will be missed by MPSS and TMPS, respectively;

• of 98,465 unaffected by SCA, 492 and 394 pregnant women will undergo unnecessary invasive test with MPSS and TMPS, respectively.

45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, MPSS: massively parallel shotgun sequencing, NA: not applicable, ND: no data available, TMPS: targeted massively parallel sequencing

^aWe did not assess the accuracy of gNIPT individually for 47,XXX, 47,XXY and 47,XYY due to paucity data.

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(Review

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for detection of fetal chromosomal aneuploidy in pregnant women

Genomics-based non-invasive prenatal testing

^bUnaffected pregnancies: we included patients with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies.

^cThe median prevalence and range were calculated by using all prospective or retrospective studies for each category considered.

^dMissed cases per 100,000 tested. FN: false negatives.

^eFalse positives per 100,000 tested. A false positive result may lead to unnecessary invasive tests depending on choices by the pregnant woman.

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BACKGROUND

Aneuploidies^[1] are chromosomal abnormalities characterised by a different (additional or missing) number of chromosomes than the 23 pairs normally present in humans. These chromosomal anomalies are among the most common types of genetic disorders and they represent a significant cause of both childhood and adulthood morbidity or death. In addition, they may lead to perinatal complications (Wellesley 2012; Wu 2013a). The severity of associated symptoms is often variable and typically less severe in mosaic cases (not all cells affected) (Fishler 1991; Modi 2003; Zhu 2013). Although offering prenatal screening for fetal aneuploidies such as Down syndrome is now considered standard of care in routine antenatal care in most upper-middle and highincome countries, prenatal screening methods and strategies are evolving. Prenatal screening consists of blood-based biochemical testing or ultrasound measurements or a combination of both, in addition to maternal age (Alldred 2012). Because of the serious health consequences of various aneuploidies and given their incurable nature, prenatal screening is an option available to pregnant women. An invasive diagnostic test (e.g. amniocentesis) is offered to pregnant women found to be at high risk of fetal aneuploidy after prenatal screening, but there is a procedurerelated risk of miscarriage. The discovery of circulating cell-free DNA (ccfDNA) in maternal blood has enabled the development of genomics-based non-invasive prenatal testing (gNIPT) to analyse the fetal genome. Prenatal screening, and ultimately prenatal diagnosis, provides couples with the information necessary for taking informed decisions (the optimisation of medical intervention and psychological counselling for managing the identified condition or pregnancy termination). The decision to terminate pregnancy among women who received a positive diagnosis of fetal aneuploidy during the prenatal period varies between 86% and 97% (Choi 2012; Irving 2011). Many factors, such as religion, maternal age, gestational age at the time of diagnosis, number of existing children, past history of induced abortion and psychosocial factors (perceived parenting burden/reward, quality of life of a child with a chromosomal abnormality, attitudes toward, and comfort with individuals with disabilities, and support from others) influence women's decision making following prenatal anomaly detection (Choi 2012).

In this systematic review, we assessed the accuracy of gNIPT for the detection of common fetal aneuploidies in pregnant women according to their prior risk of fetal aneuploidy. More specifically, we evaluated and compared the diagnostic performance of two new next-generation sequencing approaches (i.e. massively parallel shotgun sequencing (MPSS) and targeted massively parallel sequencing (TMPS)) that have recently been proposed as methods of choice to detect fetal aneuploidies by analysing ccfDNA in maternal plasma. We also made comparisons between MPSS and TMPS or between gNIPT and their combination with other first-tier screening approaches. gNIPT could be used as a firsttier test in pregnant women without prior risk (i.e. in unselected pregnant women or the general population) or as a second-tier test after a positive result for traditional first-tier screening tests such as biochemical, ultrasound or both markers (with maternal age included in risk assessment) and previous maternal history when possible.

[1] For a glossary of terms, see Appendix 1. For a list of acronyms and abbreviations, see Appendix 2.

Target condition being diagnosed

The target conditions are fetal chromosomal abnormalities diagnosed in pregnant women. The seven target conditions assessed were Down syndrome (trisomy 21 or T21), Edward syndrome (trisomy 18 or T18), Patau syndrome (trisomy 13 or T13), Turner syndrome (45,X), Klinefelter syndrome (47,XXY), Triple X syndrome (47,XXX) and 47,XYY syndrome (47,XYY) (Table 1). The majority of aneuploidies are associated with an extra copy (trisomy) of one chromosome (e.g. three copies of chromosome 21 for T21 instead of two) or a loss of one chromosome (e.g. female 45,X). Chromosomal abnormality is usually caused by a chromosome division failure or a chromosomal translocation. For example, most cases (76.2%) of 45,X karyotype (all cells affected) are caused by paternal chromosome division failure (Uematsu 2002). The most common chromosomal abnormalities are T21 and 45,X, respectively. For T21, the prevalences reported for pregnant women are 0.11% and 0.44% at 25 and 35 years old, respectively at diagnosis procedure (Snijders 1999).

Clinical characteristics and spectrum of severity are variable among aneuploidies. It has been reported that 50% of 45,X cases are mosaic (Sybert 2004). During the past few decades, caring for children with T21 or sex chromosomal abnormalities and provision of counselling to their family has changed fundamentally. These changes, including medical and surgical advances, specific interventions in the classroom for those with learning disabilities, interventions and support for parents and family members, have helped individuals with T21 live longer and enjoy an improved quality of life (Van Riper 2001). Many health problems associated with T21, 45,X, 47,XXY, 47,XXX and 47,XYY aneuploidies can be treated but fetuses with T18 and T13 are most affected and usually die in utero. The age at diagnosis varies widely depending on the condition. T21, T18 and T13 are generally detected during the perinatal period, while detection of 45,X, 47,XXX and 47,XYY is often delayed, sometimes up to 60 years old (Stochholm 2006; Stochholm 2010a; Tartaglia 2010). Around 10% of fetuses with 47,XXY are diagnosed prenatally and the mean age at diagnosis is in the mid-30s. Most 47,XXY cases are never diagnosed (Groth 2013; Tyler 2004). The incidence, clinical features and prognosis of the target conditions are summarised in Table 1.

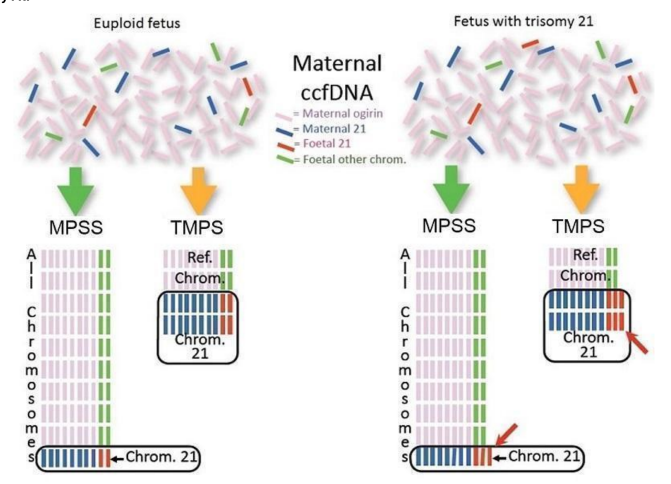
Index test(s)

Genomics-based non-invasive prenatal tests are based on the finding that placental cells continuously release detectable amounts of fetal ccfDNA into maternal blood. This fetal ccfDNA originates from normal placental cell death and consists mainly of relatively short fragments of < 300 base pairs (Bianchi 2004; Fan 2010). Proof-of-concept studies showed the feasibility of such tests to detect fetal aneuploidy in 2008 (Chiu 2008; Fan 2008).

We assessed these two gNIPT approaches (Figure 1):



Figure 1. Difference between massively parallel shotgun sequencing (MPSS) and targeted massively parallel sequencing (TMPS). Genomics-based non-invasive prenatal testing (gNIPT) aims to count the number of copies of DNA fragments from the chromosomes of interest (chromosome 21 (Chrom. 21) in this example) present in circulating cell-free DNA (ccfDNA) from a pregnant woman, relative to a reference set of chromosomes (Ref. Chrom.). DNA fragments circulating in maternal blood in the case of a euploid (left) and aneuploid (right) pregnancy are illustrated (top). MPSS produces a large number of sequence reads from all chromosomes while TMPS generates a larger proportion of reads from the chromosomes of interest (bottom). In both methods, sequence reads can be used to detect a slight excess of fetal genomic material coming from the chromosome of interest. Figure was created by FR.



- massively parallel shotgun sequencing (MPSS) which randomly analyses all DNA fragments of a sample; and
- targeted massively parallel sequencing (TMPS) which targets specific DNA fragments from the chromosomal regions of interest.

The fraction of the total ccfDNA in maternal circulation that is of fetal origin (the fetal fraction) is an important parameter for correctly identifying an aneuploid fetus by gNIPT (Canick 2013). Although the fetal ccfDNA fraction is a relatively small fraction (about 2% to 20%) of all ccfDNA in maternal blood, it can be detected from five weeks of gestation (Birch 2005; Canick 2013; Lo 1997; Lun 2008). Invasive procedures such as amniocentesis, may (Samura 2003) or may not be (Bussani 2011; Vora 2010) associated with a statistically significant increase of ccfDNA in maternal blood, which could affect fetal DNA concentration and affect gNIPT results. Therefore, in the context of clinical studies, maternal blood for gNIPT is usually collected either before or after waiting for a minimum of 24 hours following an invasive test. Indeed, the halflife of ccfDNA has been estimated to be less than one day (Lo 1999; Yu 2013). On average, euploid multifetal pregnancies have a higher fetal ccfDNA fraction than euploid singleton pregnancies (Attilakos 2011; Canick 2012). There is no reported difference in ccfDNA concentration between monochorionic and dichorionic multifetal pregnancies (Attilakos 2011). However, dichorionic pregnancies complicate gNIPT analysis by the presence of an additional genome (or more in the presence of more than two fetuses) as opposed to the two genomes of mother and fetus present in singleton or monochorionic twin pregnancies.

Next generation sequencing (NGS) applied on DNA extracted from the plasma of pregnant women generates millions of DNA sequences from both maternal and fetal genomes in relative proportion to their original abundance (for technical details see Appendix 3). The data thus produced can be used to detect a slight

excess (or loss) of fetal genomic material associated with cases of fetal aneuploidy (Papageorgiou 2012). These NGS technologies have paved the way for the development of gNIPT by alleviating the need for fetal-specific genetic markers and with potentially better test accuracy than current fetal aneuploidy screening methods.

Currently, gNIPT for the detection of common aneuploidies has been developed by companies in America, Asia and Europe and are commercially available. As part of their marketing material, these companies have published the diagnostic performance of their respective tests on their websites (Table 2). In addition, several research and clinical laboratories have developed in-house gNIPT.

Before taking a personal decision to accept or decline gNIPT, pregnant women should be given information on the screening process, which must include a discussion with a health professional (Gagnon 2010; Legare 2010; Legare 2011; St-Jacques 2008). Following screening, the results should be explained in the context of the harms and benefits of definitive diagnosis through non directive counselling (Benn 2013b). In their recent guideline, the American College of Obstetricians and Gynecologists (ACOG) recommends that gNIPT should not be used to replace diagnostic testing and that all pregnant women with a positive gNIPT result should have a diagnostic procedure before undertaking any irreversible action such as pregnancy termination. Guidelines also recommend that pregnant women with an unreported, indeterminate or uninterpretable gNIPT result should receive further genetic counselling and be offered comprehensive ultrasound evaluation and diagnostic testing (ACOG #163 2016).

Clinical pathway

Prior test(s)

Prenatal screening for fetal aneuploidy (mostly T21) is part of public health programs in most upper-middle and high-income countries and is typically offered to all pregnant women (Benn 2013b; Chitayat 2011). Up to now, screening tests for aneuploidies have relied on blood-based biochemical testing of placental markers with or without ultrasound imaging to assess for nuchal translucency thickness and other markers of fetal aneuploidy in the first trimester. The age of the pregnant woman is combined with levels of biomarkers and nuchal translucency as predictive markers for T21 in the first or second trimester (Benn 2011; Chitayat 2011; Summers 2007). Table 3 presents the various testing combinations (e.g. sequential, integrated or contingent algorithms) that have been described and are currently in use in prenatal clinics (Alldred 2017b). The screening performance of these algorithms is mostly related to the detection rates of different marker combinations and the accepted level of false positive rates. A large prospective Canadian study of 32,227 pregnant women showed that the detection rate of existing screening strategies for T21 can reach about 88.4%, with a screen-positive rate of 3.3% when applying the integrated prenatal screening procedure (Okun 2008).

A woman is classified as screen-positive if her risk is equal to or exceeds a predetermined threshold following prenatal screening result or due to some other factors such as personal or familial history of aneuploidies or translocations. Although these factors are considered to significantly increase the risk of fetal aneuploidy, the indications for invasive testing may vary between countries. To confirm the presence or absence of fetal aneuploidy in these high-risk pregnant women, a diagnostic test involving karyotyping by an invasive procedure such as amniocentesis or chorionic villi sampling (CVS) is offered (ACOG #88 2007; Benn 2011; Chitayat 2011). Karyotyping by traditional banding techniques of fetal cells obtained from amniotic fluid or placental tissue has been considered the standard of care for prenatal diagnosis of aneuploidies (ACOG #545 2012; Benn 2013a; ICFMM 2013). Fluorescence in situ hybridisation (FISH) and quantitative fluorescence polymerase chain reaction (QF-PCR) are appropriate standards of care for pregnant women at increased risk of common fetal aneuploidies based on screening results (Duncan 2011; Langlois 2011; South 2013). Microarray analysis by array comparative genomic hybridisation (aCGH) is recommended in pregnancies with fetal anomalies and it is increasingly replacing karyotyping (ACOG #682 2016).

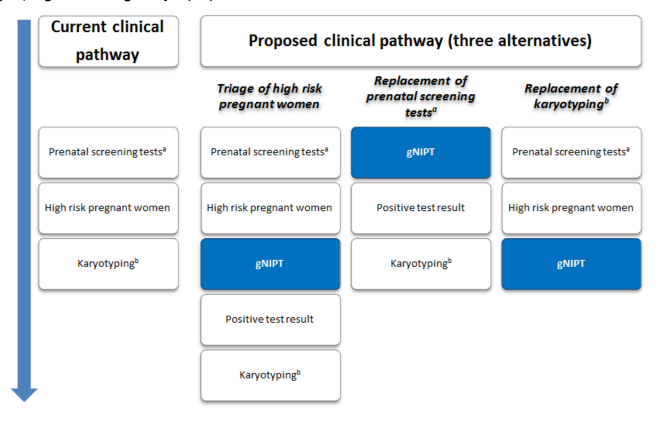
Five reviews published in the Cochrane Library examined serum, urine, ultrasound or a combination of these tests for T21 screening. For first-trimester serum tests (Alldred 2015a), the authors concluded that two markers in combination with maternal age, specifically pregnancy associated plasma protein A (PAPP-A) and free human chorionic gonadotropin (hCG) are significantly better than those involving single markers combined with or without maternal age. For second-trimester serum tests (Alldred 2012), the authors concluded that two or more markers, with or without inhibin A, in combination with maternal age are significantly more sensitive than one marker alone. Their review also showed that no test combination was superior to the others and therefore it was not possible to recommend a specific test combination. For first-trimester ultrasound tests alone of in combination with first-trimester serum tests (Alldred 2017a), the authors concluded that test strategies that combine ultrasound markers with serum markers, especially PAPP-A and free ßhCG, and maternal age were significantly better than those involving only ultrasound markers (with or without maternal age) except nasal bone. For first- and second-trimester serum tests with and without first-trimester ultrasound tests (Alldred 2017b), the authors concluded that tests involving first-trimester ultrasound with firstand second-trimester serum markers in combination with maternal age are significantly better than those without ultrasound, or those evaluating first-trimester ultrasound in combination with secondtrimester serum markers, without first-trimester serum markers. For first- and second-trimester urine tests (Alldred 2015b), the authors concluded that second-trimester ß-core fragment and oestriol with maternal age are significantly more sensitive than the single marker second-trimester ß-core fragment and maternal age. However, there were few studies and the evidence does not support the use of urine tests for T21 screening for the first 24 weeks of pregnancy.

Role of index test(s)

Genomics-based non-invasive prenatal testing such as MPSS or TMPS could be offered to pregnant women after a first-tier screening and before a diagnostic test in order to better identify which pregnant women at increased risk of fetal aneuploidy should be offered further testing (triage) (Figure 2). The use of such NGSbased approaches has also been suggested as a replacement for current first-tier screening tests (biochemical, ultrasound or both) or as potential diagnostic tests to replace current diagnostic test (karyotyping of fetal cells from amniocentesis or CVS) (Bianchi 2012).

Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review) Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

Figure 2. Current clinical pathway and three proposed uses of genomics-based non-invasive prenatal testing (gNIPT). Currently (on the left), pregnant women can have a prenatal screening test consisting of biomarkers or ultrasound, or both. For high-risk pregnant women, an invasive diagnostic test (karyotyping) is offered. In the present review, we propose 3 different clinical pathways. First, gNIPT could be used as a triage test, to decide which pregnant women should receive further testing. Second, gNIPT could be used to replace current prenatal screening tests. Finally, gNIPT could be used to replace current invasive diagnostic tests (if diagnostic performance permits). At any point in a clinical pathway, a pregnant woman may decide not to proceed with other tests (not shown in the figure). Figure was designed by CL, JB, MB and YT.



^aPrenatal screening test consists of serum biomarkers or ultrasound measurements or both offer to all pregnant women.

^bKaryotyping from amniocentesis or chorionic villus sampling.

Rationale

Current screening tests (biochemical, ultrasound or both) have relatively high false positive rates, which may result in undue anxiety for many pregnant women who will be offered an invasive diagnostic procedure. For example, at a prenatal screening risk cut-off of 1:300, fetal aneuploidy is confirmed by karyotyping in only about 1/34 to 1/14 (3% to 7%) screen-positive cases (Renshaw 2013; Wald 2005). As a result, many more women will undergo invasive diagnostic testing following positive screening tests than the number carrying a fetus with aneuploidy. In France, each year, about 800,000 pregnant women opt for prenatal T21 biochemical screening, ultrasound measurements or both, and about 24,000 of them (3%) will have karyotype testing (Basset 2013). Invasive testing methods for prenatal diagnosis of aneuploidy identify pregnancies with fetal chromosomal abnormalities, but contribute to an additional procedure-related fetal loss rate (Wilson 2007). A recent meta-analysis showed that weighted pooled procedurerelated risks of miscarriage of invasive testing methods before 24 weeks' gestation were 0.11% for amniocentesis and 0.22% for CVS (Akolekar 2015). The risk of miscarriage of normal fetuses associated with such invasive procedures has fostered the development of alternative screening and diagnostic approaches.

The discovery of fetal circulating cells and fetal ccfDNA in maternal blood during pregnancy has enabled the development of noninvasive methods to analyse the fetal genome (Birch 2005; Lo 1997; Wright 2009). Fetal DNA offers advantages over circulating fetal cells because it is more easily extracted from maternal plasma samples and it disappears within hours after birth (undetectable about one to two days postpartum), as compared to the paucity and persistence of fetal cells in maternal blood over several consecutive pregnancies (up to 27 years) (Wright 2009; Yu 2013). At present, the analysis of ccfDNA by NGS technologies seems to be the most

promising alternative gNIPT approach for the detection of fetal aneuploidies from maternal blood. This allows sequencing of tens of millions of these DNA fragments simultaneously, paving the way for the development of a non-invasive, less psychologically stressful method potentially able of detecting fetal aneuploidies earlier and with better accuracy than current screening programs. As such, NGS technologies have the potential to radically change prenatal screening for fetal aneuploidy. Indeed, a study exploring the impact of gNIPT on prenatal care showed that more pregnant women with positive first-trimester screening opt for further testing (from 47.2% to 78.8%) than before the introduction of gNIPT, while the rate of invasive diagnostic testing has decreased significantly (from 47.2% to 39.2%). Additionally, fewer pregnant women declined follow-up testing when gNIPT was an option (from 52.8% to 21.2%) (Chetty 2013). Another study suggested that gNIPT could reduce procedure-related fetal losses in high-risk women by up to 88% (O'Leary 2013).

For instance, the new gNIPT approach is reported to detect aneuploidy with high sensitivity to select a subset of pregnant women for an invasive diagnostic procedure and could be performed in high-risk pregnant women (as a second-tier test) following a positive screening result (Benn 2013a). The major expected advantage of gNIPT by NGS over current (biochemical, ultrasound or both) screening tests is the significant decrease in false positive results and thus the reduction of invasive procedures and their associated normal fetus losses. Also, it was reported that a reduction of invasive prenatal procedures with the introduction gNIPT has indeed been documented (Chetty 2013; Larion 2014; Tiller 2014). Assessment of how NGS should be used in clinical practice for an uploidy detection is currently being studied. NGS approaches could also be performed in general obstetrical population (as first-tier test), in place of current screening algorithms (biochemical, ultrasound or both) (Figure 2). However, the field is moving rapidly. From January to July 2014, around 60 NIPT studies were published in PubMed compared to 70 studies in 2013 and 40 studies in 2012.

Up to now, no comprehensive systematic review including metaanalyses has analysed and compared the diagnostic accuracy of MPSS and TMPS methods for the detection of fetal aneuploidies, either as a second-tier test (i.e. in women at increased risk of fetal aneuploidy after current screening procedures) or as a firsttier test (i.e. in all pregnant women). Benn 2013b published a review on gNIPT focused on providing the information needed by clinicians and public health providers before implementation of this technology in routine clinical practice. However, their review included only T21 and T18. Mersy 2013 published a systematic review on guality and outcome of diagnostic test accuracy studies on non-invasive detection of fetal T21 only. One updated metaanalysis (Gil 2015a) pooled all gNIPT methods but did not assess the relative performance of MPSS and TMPS technologies separately. More recently, Taylor-Phillips 2016 published a meta-analysis on gNIPT accuracy for major autosomal anomalies (T21, T18 and T13) without sex chromosome aneuploidies (SCAs) assessment and using restrictive inclusion criteria for included publications (e.g. limited to the English language, cohorts of more than 50 pregnant women) and including studies with incomplete follow-up (pregnant women without reference standard). In the meta-analysis of Mackie 2017, multifetal pregnancies and case-control study design were excluded. In the meta-analysis published by the Haute Autorité de Santé in France (HAS 2015), the accuracy of gNIPT was evaluated for T21 only and included studies with pregnant women selected at high risk of fetal aneuploidy as well as studies with pregnant women unselected for their risk (general population). Only studies published in English were included. The review of Agarwal 2013 described the properties of commercial tests available (e.g. type of gNIPT method, costs, turnaround times and reimbursement), intellectual property, commercialisation, patenting, patenting litigation and licensing landscape of technologies underlying these tests.

Genomics-based non-invasive prenatal tests are already advertised and marketed to North-American, European and Asian healthcare providers. Leading companies are summarised in Table 2. Other entities are trying to make their way into the market (Birmingham Women's NHS; Counsyl; GENDIA; Genesis Genetics; Integrated Genetics; NIPD Genetics; Progenity; Quest Diagnostics; RAVGEN; Xcelom). Some of these assays have yet to be approved by the US Food and Drug Administration. There is significant pressure for increasing their use in clinical practice, but comparative effectiveness and cost-effectiveness studies, as well as studies of the ethical, legal and social issues are scarce. Furthermore, tools needed for their patient value-based implementation are not available or have not been validated.

OBJECTIVES

To evaluate and compare the diagnostic accuracy of massively parallel shotgun sequencing (MPSS) and targeted massively parallel sequencing (TMPS) using circulating cell-free DNA (ccfDNA) in maternal blood for the detection of common fetal aneuploidies (T21, T18, T13, 45,X, 47,XXY, 47,XXX and 47,XYY) according to their prior risk of fetal aneuploidy. The genomics-based non-invasive prenatal testing (gNIPT) results were confirmed by a reference standard such as fetal karyotype or neonatal clinical examination.

To evaluate the screening performance of MPSS and TMPS as triage tests (a second-tier screening test) for identifying which pregnant women at increased risk of fetal aneuploidy should be offered further testing, that is, after a first-tier screening, but before a diagnostic test.

To assess the screening performance of MPSS and TMPS as a firsttier test in pregnant women without prior risk (i.e. in unselected pregnant women or general population) as a replacement for current offered first-tier tests (biochemical, ultrasound or both).

To assess the diagnostic performance of MPSS and TMPS as a second-tier test as potential diagnostic tests to replace current invasive diagnostic tests.

Secondary objectives

To investigate potential sources of heterogeneity that may influence the diagnostic accuracy of MPSS and TMPS such as gestational age at the time of blood collection and type of reference standard used.

METHODS

Criteria for considering studies for this review

Types of studies

We included studies that met the following inclusion criteria:



- randomised studies where pregnant women were randomised to receive one gNIPT (MPSS or TMPS) as well as the reference standard:
- retrospective and prospective cohort studies where all pregnant women were tested with one or more gNIPT methods and the reference standard (including head-to-head studies); and
- retrospective and prospective case-control studies comparing one or more of the gNIPT methods with the reference standard.

Although studies with a retrospective or case-control design are prone to biases, we included such studies because we anticipated a paucity of other study designs. When data were sufficient, we explored the effect of excluding case-control studies in sensitivity analyses.

We excluded studies for which it was not possible to extract or derive the number of true positives, false positives, false negatives and true negatives.

Participants

We included women of any age, ethnicity and gestational age with a singleton or multifetal (monochorionic and dichorionic) pregnancy.

Index tests

Genomics-based non-invasive prenatal tests based on plasma ccfDNA in maternal blood, analysis by either MPSS or TMPS methods.

Target conditions

We considered seven fetal aneuploidies, namely T21, T18, T13, 45,X, 47,XXY, 47,XXX and 47,XYY.

Reference standards

We considered the following test as reference standard: fetal karyotyping performed on cells obtained from chorionic villi sampling (CVS), amniotic fluid, placental tissue, a fetus lost by miscarriage or other equivalent and recognised methods on the same materials. By "fetal karyotyping" we mean traditional banding techniques, spectral karyotyping, fluorescence in situ hybridisation (FISH), array comparative genomic hybridisation (aCGH) or quantitative fluorescence polymerase chain reaction (QF-PCR). If fetal karyotyping was not performed, we used neonatal clinical examination or medical records from birth as a secondary reference standard for T21, T18 or T13. For sex chromosome aneuploidies (SCA), only fetal karyotype was an appropriate reference standard because newborns usually have a normal phenotype.

Search methods for identification of studies

Electronic searches

We used a sensitive search strategy that included the following three sets of search terms and synonyms:

- index test (e.g. cell-free DNA, sequencing, non-invasive and genetic diagnosis);
- participants' description (e.g. pregnant women, fetus and prenatal); and
- target condition (e.g. aneuploidy and chromosome anomalies).

We combined free-text words and subject headings used within each set with the Boolean operator OR and then combined the three sets using AND. We reviewed publications from 1st January 2007 because MPSS and TMPS were introduced in the literature in 2008 (Chiu 2008; Fan 2008). We did not limit our search by language, search filter or publication type (e.g. journal article, clinical trial, validation study, review and comment).

We applied a comparable search strategy (Appendix 4) with adaptations for each of the following databases:

- MEDLINE (Ovid) (January 2007 to July 2016);
- Embase (January 2007 to July 2016);
- Web of Science (ISI) (January 2007 to July 2016);
- Cochrane Register of Diagnostic Test Accuracy Studies, Cochrane Library (January 2007 to October 2016);
- ClinicalTrials.gov (January 2007 to September 2016);
- European Clinical Trials Register (January 2007 to September 2016);
- WHO ICTRP (January 2007 to September 2016);
- The National Technical Information Service (NTIS) (January 2007 to September 2016);
- OpenGrey (January 2007 to October 2016); and
- National Guideline Clearing House (January 2007 to September 2016).

Searching other resources

We examined references cited in potentially relevant full-text papers and those cited in previous reviews by cross-checking bibliographies. We examined grey literature by searching data available on the websites of private prenatal diagnosis companies (Ariosa Diagnostics 2016; BGI 2016; Berry Genomics 2016; Genoma 2016; Genome Care 2016; Illumina 2016; LabGenomics 2016; LifeCodexx 2016; Natera 2016; Genesupport 2016; Premaitha Health plc 2016; Sequenom 2016) using gNIPT technologies (January 2007 to December 2016). We also searched for conference abstracts and theses in appropriate sources (e.g. TheseNet, Theses Canada Portal) (January 2007 to October 2016).

Data collection and analysis

We used the methods suggested by the Cochrane Diagnostic Test Accuracy Working Group (Deeks 2013). For selection of studies, data extraction and assessment of methodological quality, we conducted a pilot using 20 randomly selected articles to trial our forms in order to ensure criteria were applied consistently.

None of the review authors involved in conducting a gNIPT primary study (FL, FR, SL and YG) took part in the selection of studies, nor in any decisions/analyses related to their own studies. Furthermore, by the final date of data collection, these authors had not published a primary gNIPT study.

Selection of studies

Two review authors (MB and CL) independently identified relevant studies by screening the titles and abstracts of all studies identified by the search strategy. We obtained the full-text version of all potentially relevant studies and assessed them for inclusion by using a study eligibility table based on prespecified inclusion criteria. The data collection form (Excel® format) for classifying studies during the full-text assessment is presented in Appendix



5. We considered all comments, statements or errata related to included studies. We excluded studies that did not match the inclusion criteria and we recorded the reason(s) for exclusion. If results from the same study cohort were reported in multiple publications, we considered all the publications and included results from the most relevant and comprehensive publications. We excluded papers with preliminary results whose full published results were available. We resolved any disagreement between assessors (MB and CL) by iteration, discussion and consensus. If required, we consulted a third review author (JB or LN).

Data extraction and management

Two review authors (MB and CL, JB or LN) independently extracted information and data from each included study by using a data extraction form that we developed in Excel[®] format. We included the following items:

- study characteristics (e.g. reference details allowing identification of the publication, language and study design);
- population characteristics (e.g. gestational age, maternal age, ethnicity, total number of pregnant women, number of aneuploid cases, number of euploid cases, recruitment location (country, geographic locations or regions), recruitment period and other relevant tests carried out prior to index test (e.g. ultrasonography, biochemical screening));
- features of the reference standard (e.g. fetal karyotyping, chromosome analysis or clinical examination);
- features of the index test (e.g. technical details, commercial or in-house gNIPT, cutpoint, failure rate, blood sample collection time (before or after reference standard) and first-tier test or second-tier test); and
- data for constructing two-by-two tables (number of true positives, false positives, false negatives and true negatives) or summary statistics from which the data were derived. In the two-by-two tables, the true negative cases were patients with any other aneuploidy than the one under analysis and all euploid cases were considered unaffected. When data were presented in three-by-two tables due to unclassified index test results (defined as grey zone between positive and negative test results), we constructed two-by-two tables by considering all unclassified gNIPT results as test positives. This is because in practice such results will lead to further testing and investigation to ensure a case of fetal aneuploidy is not missed.

We cross-checked all extracted and recorded data and we resolved any disagreement by iteration, discussion and consensus between two review authors (MB and CL, JB or LN). If required, we consulted a third author (JB, LN or CL). We wrote to the study contact author if information was missing or unclear or to clarify potential overlap between publications based on the same dataset to avoid including the same women more than once. If an article presented results including other aneuploidies than the ones under review, we considered only the subset of the cohort with the aneuploidies of interest.

Assessment of methodological quality

We used the revised QUality Assessment of Diagnostic Accuracy Studies (QUADAS-2) tool for assessment of methodological quality of included studies (Whiting 2011). We tailored the tool to this review question using the operational criteria detailed in Appendix 6 to answer signalling questions and make the overall judgment of risk of bias and applicability concerns for each domain of the tool. We answered each signalling question with a 'yes', 'no' or 'unclear' response for each included study and we recorded the reason for the judgment made. If a study was recorded as 'yes' on all signalling questions related to risk of bias, then it was deemed appropriate to have an overall judgment of 'low risk of bias'. If a study is recorded 'no' or 'unclear' on one or more signalling questions in a domain, then it was judged as having 'high or unclear risk of bias'. Judgments about applicability concern were rated as 'low', 'high' or 'unclear' in relation to our review question. 'Unclear concern' was used only if insufficient information was available. Two review authors (MB and CL, JB or LN) independently applied the QUADAS-2 tool to each included study and we resolved any disagreement by iteration, discussion and consensus. If required, we consulted a third review author (JB, LN or CL).

Statistical analysis and data synthesis

The unit of analysis was the pregnant woman irrespective of the type of pregnancy (multifetal or singleton pregnancy). We evaluated the performance of MPSS and TMPS for the detection of each type of aneuploidy under study both individually and globally for any type of aneuploidy (all autosomal aneuploidies combined and all sex chromosomal aneuploidies combined). We distinguished between each of the following groups of pregnant women and performed separate analyses for each subgroup:

- unselected pregnant women undergoing aneuploidy screening (first-tier gNIPT, i.e. offered to all pregnant women) and women selected at high risk of fetal aneuploidy (second-tier gNIPT);
- women with singleton and multifetal pregnancy because ccfDNA's fetal fraction in multifetal pregnancy is higher than in singleton pregnancy (Attilakos 2011; Canick 2012); and
- pregnant women who underwent gNIPT during the first trimester (15 weeks or less), the first or second trimester (29 weeks or less) or at any time during pregnancy (42 weeks or less).

For each gNIPT method, we used Review Manager® to produce coupled forest plots of sensitivity and specificity, together with their 95% confidence intervals (CIs). We also plotted studyspecific estimates of sensitivity and specificity in receiver operating characteristic (ROC) space. All gNIPTs are laboratory-developed tests based on differently calibrated assays with specific cutpoints to classify samples as euploid or aneuploid. There is no consensus on the cutpoints to use in practice. For this reason, we had planned to use a modelling strategy that focuses on the estimation of summary ROC curves (Macaskill 2010; Rutter 2001) and to estimate summary points (summary sensitivity and specificity) if a sufficient number of studies reported common cutpoints. However, given the qualitative nature of the cutpoints, which is highly dependent on each laboratory's developed gNIPT and study populations, it was not possible to identify a common cutpoint. Therefore, we reasoned that this was a special case where we can assume gNIPT results were binary (positive or negative). The rationale was further strengthened by the lack of apparent threshold effect when we examined the studies in ROC space. If a study reported more than one cutpoint, we considered all cutpoints and chose one cutpoint, the most commonly reported across all studies, such that only one pair of sensitivity and specificity from a study was included in metaanalysis.

Due to limited or absence of threshold effect, there was no requirement to account for correlation between sensitivity



and specificity across studies in meta-analysis. Therefore, we removed the correlation parameter from the bivariate model (Chu 2006), thus simplifying the model to two univariate random-effects logistic regression models for separate meta-analyses of sensitivities and specificities (Takwoingi 2015). In cases where there were few studies in the meta-analysis or a random-effects analysis failed to converge, we used fixed-effect logistic regression models. Where all studies in the meta-analysis reported 100% sensitivity or 100% specificity, these fixed-effect models fail as the prediction is perfect. Therefore, in such situations we used simple pooling by summing up the numbers of true positives and total cases to compute sensitivity, and the numbers of true negatives and unaffected pregnancies to compute specificity. Cls were obtained using the Wilson method (Newcombe 1998).

We compared the diagnostic accuracy of MPSS and TMPS by first using all available data (indirect comparison). If studies that compared MPSS and TMPS in the same population (head-to-head or direct comparison) were available, we had planned a second set of analyses restricted to direct comparisons. Comparative meta-analyses were done by adding a covariate for test type to random-effects or fixed-effect models. We used likelihood ratio tests to assess the statistical significance of differences between tests by comparing models that included covariate terms for test type with models that did not include the terms. If data were available, comparisons between gNIPTs and traditional screening approaches were planned using a similar strategy to that described above. Meta-analyses were performed using the xtmelogit and blogit functions in the Stata software package (version 13; StataCorp, College Station, Texas 77845, USA). When meta-analyses of direct comparisons were not possible, we examined individual study results. For each comparative study, we computed differences in sensitivity and specificity, and 95% CIs were calculated for the differences using the Newcombe-Wilson method without continuity correction (Newcombe 1998).

Investigations of heterogeneity

We examined forest plots of sensitivity and specificity and summary ROC plots for each gNIPT method to visually assess heterogeneity. If sufficient data were available for meta-regression (by adding a covariate to a logistic regression model to explore its effect on sensitivity and specificity), we had planned to investigate the effect of the following:

- study population (e.g. ethnicity, gestational age at blood collection); and
- type of reference standard (i.e. karyotype or mixed reference standard).

However, formal investigations using meta-regression were not possible due to limited data and little or no heterogeneity in test accuracy.

Sensitivity analyses

We performed sensitivity analyses to assess the effect of excluding case-control studies and studies with a small number of cases of aneuploidy (less than 10 cases) on the summary estimates of test accuracy.

We had planned to also assess the effect of:

- studies where pregnant women received an invasive diagnostic test less than one day before blood collection for gNIPT;
- third trimester gestational age at the moment of blood collection for gNIPT;
- studies available only as abstracts; and
- studies at 'high or unclear risk of bias' according to the QUADAS-2 assessment tool.

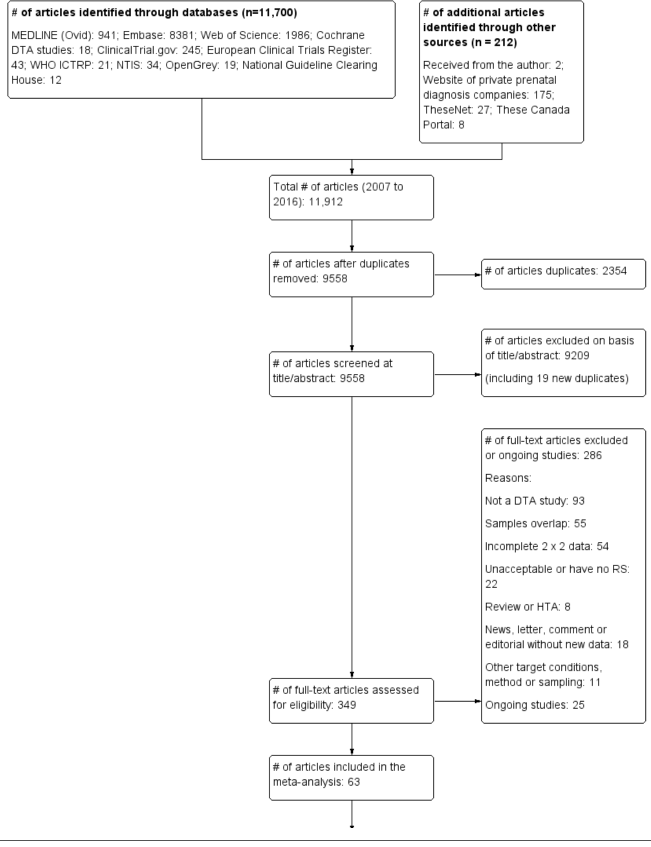
However, due to lack of data or lack of variability in estimates of sensitivity and specificity, only assessments of the impact of study design and number of cases were performed.

RESULTS

Results of the search

We found a total of 11,912 articles through our electronic searches from January 2007 to October 2016 (see PRISMA study flow diagram in Figure 3). A total of 11,700 articles were identified through databases (941 through MEDLINE, 8381 through Embase, 1986 through Web of Science, 18 through Cochrane Diagnostic Test Accuracy register of studies, 245 through ClinicalTrial.gov, 43 through European Clinical Trials Register, 21 through WHO ICTRP, 34 through NTIS, 19 through OpenGrey and 12 through the National Guideline Clearing House). We found 212 publications through other sources (two articles received from the author, 175 from gNIPT company's website, 27 from TheseNet and eight from These Canada Portal). After removing 2354 duplicates, two review authors independently screened the titles and abstracts of 9558 publications. Of the 9558 publications, 9209 were deemed irrelevant to our review question. We retrieved the full texts of the remaining 349 articles to assess their eligibility. After resolving disagreement between two or three review authors, 261 articles were excluded (see Characteristics of excluded studies for details) and 63 articles fulfilled our inclusion criteria (see Characteristics of included studies for details). Among these 63 articles, 62 were journal articles and one was a letter to the editor with sufficient information to be included (Jackson 2014). From the 63 articles, two articles presented two studies (two different cohort, two 2x2 tables). At all, we included 65 studies of 86,139 pregnant women (3141 aneuploids and 82,998 euploids). No studies are awaiting classification. We identified 25 ongoing trials through clinical trials databases (see Characteristics of ongoing studies for details). We will consider these trials in future updates.

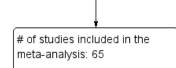
Figure 3. PRISMA flow diagram for selection of studies from January 2007 to October 2016. #: number, DTA: diagnostic test accuracy, NTIS: The National Technical Information Service and WHO ICTRP: World Health Organization International Clinical Trials Registry Platform.



Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review) Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.



Figure 3. (Continued)



Basic features of the included studies

The clinical characteristics of pregnant women and sequencing method were generally well described or referenced. Some studies did not clearly report how patient selection was done and which inclusion and exclusion criteria were applied. Patients' enrolment flow-charts, pregnancy outcome flow-chart and 2 x 2 tables were unclear in many studies. We therefore contacted study authors to clarify unclear information, to obtain missing data or to clarify potential overlap of patients between publications.

We described the characteristics of included studies in Characteristics of included studies table and provided a summary in Table 4. Forty-two studies (65%) enrolled pregnant women selected at high risk of fetal aneuploidy (Alberti 2015; Ashoor 2012; Benachi 2015; Bianchi 2012; Bianchi 2013; Bijok 2014; Canick 2012; Chen 2011; Ehrich 2011; Hall 2014; Hooks 2014; Hou 2012; Huang 2014; Jeon 2014; Jiang 2012; Johansen 2016; Ke 2015; Kim 2016; Lee 2015; Lefkowitz 2016; Liang 2013; Liu 2012; Mazloom 2013; Nicolaides 2013; Nicolaides 2014a; Norton 2012; Palomaki 2012; Papageorghiou 2016a; Papageorghiou 2016b; Persico 2016; Poon 2016; Porreco 2014; Sehnert 2011; Song 2015; Sparks 2012a; Stumm 2014; Sukhikh 2015; Sung-Hee 2015; Verweij 2013; Wang 2014; Wang 2015a; Zhang 2016); five studies (8%) enrolled pregnant women without prior risk of fetal aneuploidy (del Mar Gil 2014; Nicolaides 2012; Norton 2015; Quezada 2015; Song 2013); and 18 studies (28%) enrolled pregnant women from a mixed risk cohort of fetal aneuploidy (Ashoor 2013; Bevilacqua 2015; Bianchi 2014a; Chiu 2011; Comas 2015; Fiorentino 2016; Gil 2016; Jackson 2014; Korostelev 2014; Lau 2012; Ma 2016; Pergament 2014; Samango-Sprouse 2013; Shaw 2014; Tynan 2016; Yao 2014; Zhou 2014a; Zhou 2014b). Mixed-risk samples included a mixture of selected pregnant women with low, high or no prior risk of fetal aneuploidy. Such samples do not represent the real-life situation (i.e. using gNIPT as a first-tier screening test or as a second-tier test) and so such studies were not used for addressing our research objectives. Nevertheless, as we did not pre-specify exclusion of such studies, we analysed the data and the results are presented in Appendix 7.

The studies assessed MPSS and TMPS using various algorithms and cutpoints. Table 4 describes the specific gNIPT assay used in the included studies. Each assay was developed and validated by the testing laboratory. Among the 65 studies, 44 studies (68%) used a whole genome sequencing method (MPSS) (Alberti 2015; Benachi 2015; Bianchi 2012; Bianchi 2013; Bianchi 2014a; Bijok 2014; Canick 2012; Chen 2011; Chiu 2011; Ehrich 2011; Fiorentino 2016; Hou 2012; Huang 2014; Jeon 2014; Jiang 2012; Johansen 2016; Ke 2015; Kim 2016; Lau 2012; Lee 2015; Lefkowitz 2016; Liang 2013; Liu 2012; Ma 2016; Mazloom 2013; Palomaki 2012; Papageorghiou 2016a; Papageorghiou 2016b; Poon 2016; Porreco 2014; Sehnert 2011; Shaw 2014; Song 2013; Song 2015; Stumm 2014; Sukhikh 2015; Sung-Hee 2015; Tynan 2016; Wang 2014; Wang 2015a; Yao

2014; Zhang 2016; Zhou 2014a; Zhou 2014b), and 21 (32%) used a targeted method (TMPS) (Ashoor 2012; Ashoor 2013; Bevilacqua 2015; Comas 2015; del Mar Gil 2014; Gil 2016; Hall 2014; Hooks 2014; Jackson 2014; Korostelev 2014; Nicolaides 2012; Nicolaides 2013; Nicolaides 2014a; Norton 2012; Norton 2015; Pergament 2014; Persico 2016; Quezada 2015; Samango-Sprouse 2013; Sparks 2012a; Verweij 2013). Of the 65 studies, five studies compared gNIPT with traditional screening tests (Bianchi 2014a; Nicolaides 2012; Norton 2015; Quezada 2015; Song 2013). MPSS studies involved 50,864 pregnant women, TMPS studies involved 35,275 pregnant women and traditional screening tests involved 24,279 pregnant women. The most commonly (15 studies) used cutpoint for gNIPT assays was a chromosomal ratio Z score of 3. Thirteen studies used the FORTE risk score, eight studies used a normalised chromosome value (NCV) and 13 studies did not report their cutpoint. The remaining studies used other cutpoints (Table 4). Timing of blood sampling for gNIPT was before invasive procedure in 55 studies, before or more than 24 hours after invasive sampling in four studies (Ashoor 2013; Lefkowitz 2016; Pergament 2014; Samango-Sprouse 2013), and was not reported in six studies (Bevilacqua 2015; Jiang 2012; Song 2013; Sparks 2012a; Wang 2014; Zhang 2016).

Among all aneuploidies considered, 36 studies (55%) reported analyses only for autosomes, four (6%) for only sex chromosome aneuploidies (SCA) and 25 studies (39%) for both autosomes and SCA. Fifty-seven studies (82,620 pregnant women) evaluated T21, 50 studies (79,322 pregnant women) evaluated T18, 39 studies (68,958 pregnant women) evaluated T13, 20 studies (10,081 pregnant women) evaluated 45,X, seven studies (6035 pregnant women) evaluated 47,XXX, 12 studies (7609 pregnant women) evaluated 47,XXY and 10 studies (6987 pregnant women) evaluated 47,XYY (Table 4). Among all 65 included studies, there are a total of 2004 T21 cases, 634 T18 cases, 215 T13 cases, 232 45,X cases, 14 47,XXX cases, 25 47,XXY cases and 16 47,XYY cases. All 65 studies used an appropriate reference standard such as fetal or neonatal karyotype, genetic testing, neonatal clinical examination or medical records from birth. In 36 studies (55%), only one reference standard was used while 29 studies (45%) used more than one reference standard (Table 4).

Among the 65 studies, 40 (62%) studies were prospective cohort studies (Ashoor 2013; Bevilacqua 2015; Bianchi 2014a; Bijok 2014; Comas 2015; Fiorentino 2016; Gil 2016; Hou 2012; Huang 2014; Jackson 2014; Jeon 2014; Jiang 2012; Johansen 2016; Ke 2015; Kim 2016; Korostelev 2014; Lau 2012; Lee 2015; Liang 2013; Liu 2012; Mazloom 2013; Nicolaides 2013; Norton 2012; Norton 2015; Pergament 2014; Persico 2016; Porreco 2014; Quezada 2015; Samango-Sprouse 2013; Shaw 2014; Song 2013; Song 2015; Stumm 2014; Sukhikh 2015; Verweij 2013; Wang 2014; Wang 2015a; Zhang 2016; Zhou 2014a; Zhou 2014b), eight (12%) studies were retrospective cohort studies (Benachi 2015; Bianchi 2013; del Mar Gil 2014; Nicolaides 2012; Sehnert 2011; Sung-Hee 2015; Tynan



2016; Yao 2014), one (1%) study was a prospective and retrospective cohort study (Ma 2016) and 16 (25%) studies used a case-control design (Alberti 2015; Ashoor 2012; Bianchi 2012; Canick 2012; Chen 2011; Chiu 2011; Ehrich 2011; Hall 2014; Hooks 2014; Lefkowitz 2016; Nicolaides 2014a; Palomaki 2012; Papageorghiou 2016b; Poon 2016; Sparks 2012a) (Table 4).

Forty-eight (74%) studies included only singleton pregnancies, while five (8%) studies included only multifetal pregnancies. Four (6%) studies included women with either type of pregnancy and eight (12%) studies did not report the type of pregnancy. Ten (15%) studies included only pregnant women in the first trimester (15 weeks or less), 21 (33%) studies included pregnant women in the first two trimesters (29 weeks or less), 24 studies (37%) included pregnant women in the three trimesters (42 weeks or less) and 10 studies (15%) did not report gestational age. Eighteen studies (28%) had more than 50% Caucasian women in their cohort, 21 studies (32%) had more than 50% Asian women and 26 studies (40%) did not report ethnicity.

Thirty-seven studies (57%) were industry-funded or were written by one or more author affiliated with a company who sells gNIPT (Benachi 2015; Bianchi 2012; Bianchi 2013; Bianchi 2014a; Canick 2012; Chen 2011; Chiu 2011; Ehrich 2011; Hall 2014; Hooks 2014; Huang 2014; Jackson 2014; Jiang 2012; Kim 2016; Lau 2012; Lee 2015; Lefkowitz 2016; Ma 2016; Mazloom 2013; Nicolaides 2012; Nicolaides 2013; Norton 2012; Norton 2015; Palomaki 2012; Papageorghiou 2016a; Papageorghiou 2016b; Pergament 2014; Persico 2016; Porreco 2014; Samango-Sprouse 2013; Sehnert 2011; Shaw 2014; Sparks 2012a; Stumm 2014; Tynan 2016; Verweij 2013; Yao 2014); 22 studies (34%) were not reported to be funded by industry but samples were sequenced and analysed by a commercial laboratory (Ashoor 2012; Ashoor 2013; Bevilacqua 2015; Bijok 2014; Comas 2015; del Mar Gil 2014; Fiorentino 2016; Gil 2016; Hou 2012; Jeon 2014; Ke 2015; Korostelev 2014; Liang 2013; Poon 2016; Quezada 2015; Song 2013; Song 2015; Sung-Hee 2015; Wang 2014; Wang 2015a; Zhou 2014a; Zhou 2014b); three studies (4.5%) had no link with industry (Alberti 2015; Johansen 2016; Sukhikh 2015); and the funding source was not reported for three studies (4.5%) (Liu 2012; Nicolaides 2014a; Zhang 2016). Table 5 describes the specific gNIPT assay used in the included studies. Of the 65 studies, 61 (94%) used a commercial gNIPT (15 from Ariosa Diagnostics, Inc., 12 from Bejing Genomics Institute, four from Illumina (or Verinata Health), six from Natera, nine from Sequenom and 15 from other companies) (Table 5). It appears that, for three of the commercially available assays, there are nine studies or more adding up to a large number of cases and unaffected cases analysed. Further, only two assays (one TMPS and one MPSS) were used in one of the five studies involving unselected pregnant women and one assay (Ariosa's Harmony[™] test) was used in four of them. Twelve studies (19%) included their entire cohort in the analyses, 36 studies (55%) included between 80% to 99.9%, and 17 studies (26%) included less than 80%. We found 54 (83%) studies where patient exclusions and failed samples were reported (Table 6; Table 7).

Summary of excluded studies

We described the excluded studies in the PRISMA flow diagram (Figure 3) as well as in Characteristics of excluded studies. After full-text assessment, we excluded 261 articles.

Of these 261:

- 93 (36%) studies were not diagnostic test accuracy studies (e.g. implementation study, simulation model, method development, proof-of-concept, method without sequencing approach);
- 55 (21%) studies had overlapping samples and were excluded to avoid double counting;
- 54 (21%) studies had incomplete 2 X 2 data or insufficient information to derive a 2 X 2 table;
- 22 (8%) studies had either an inappropriate or no reference standard;
- 8 (3%) studies were identified as reviews or Health Technology Assessment reports;
- 11 (4%) studies had target conditions, methods or sampling schemes other than those specified in our review; and
- 18 (7%) studies were news, letters, comments, notes, replies or editorials without new data.

The 25 ongoing studies are described in Characteristics of ongoing studies.

Methodological quality of included studies

Figure 4 and Figure 5 show the risk of bias and applicability concerns for each included study for MPSS and TMPS, respectively. In Figure 6, the quality assessment results are summarised across all studies.

		<u> </u>	<u> </u>		 ·	· ·		
	F	Risk o	of Bias	5	 Appli	cabilit	y Con	icerns
	Patient Selection	Index Test: MPSS	Reference Standard	Flow and Timing	Patient Selection	Index Test: MPSS	Reference Standard	
Alberti 2015		•	•		•	•	•	
Benachi 2015	•	•	•	•	•	•	•	
Bianchi 2012	•	•	•	•	•	•	•	
Bianchi 2013	•	٠	•	•	•	•	•	
Bianchi 2014a		•	•		•	•	•	
Bijok 2014	?	•	•	•	•	•	٠	
Canick 2012		•	•	•	•	•	•	
Chen 2011		?	•	•	•	•	•	
Chiu 2011		•	?		•	•	•	
Ehrich 2011		•	•		•	•	•	
Fiorentino 2016	•	•	?	•	•	•	•	
Hou 2012	•	•	•	•	•	•	•	
Huang 2014	•	•	•	•	•	•	•	
Jeon 2014	•	•	•	•	•	٠	•	
Jiang 2012	•	•	•	?	•	•	•	
Johansen 2016	•	•	•	•	•	•	•	
Ke 2015	•	•	?	•	•	•	•	
Kim 2016	•	•	•	•	•	•	•	
Lau 2012	•	•	•	•	•	•	•	
Lee 2015	•	•	•	•	•	•	•	
Lefkowitz 2016	•	•	•	•	•	•	•	
Liang 2013	•	•	•	•	•	•	•	
Liu 2012	?	•	•	•	•	•	•	
Ma 2016	•	•	•	•	•	•	•	
Mazloom 2013	•	•	•	•	•	•	•	
Palomaki 2012	•	•	•	•	•	•	•	

Figure 4. Risk of bias and applicability concerns summary: review authors' judgements about each domain for each of the studies included for massively parallel shotgun sequencing (MPSS).

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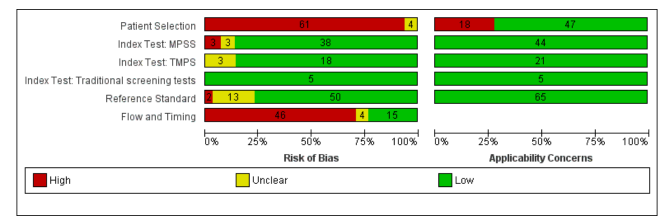
Figure 4. (Continued)

Papageorghiou 2016a
Poon 2016 • <
Porreco 2014 • <t< td=""></t<>
Sehnert 2011 Image: Constraint of the sentence o
Shaw 2014 •
Song 2013
Song 2015 😑 🔁 🙂 🙂 🙂
Stumm 2014 🔴 😑 😁 🕒 🙂 🙂
Sukhikh 2015 🔴 ? ? 🔮 🚯 🔮
Sung-Hee 2015 😑 😗 😑 😑 🔮 🔮
Tynan 2016 😑 🔹 😑 😑 🔮
Wang 2014 🔴 😗 ? ?
Wang 2015a 😑 😗 😗 🔮 🔮 🖤
Yao 2014 😑 😍 ? 😑 🕒 🙂
Zhang 2016 😑 😬 😗 🕐 🕑 🖤
Zhou 2014a 😐 😗 😨 🛑 🖤 🕐
Zhou 2014b 😑 ? ? 😑 🕒 😬
🛑 High 🥐 Unclear 📀 Low

r										_
		Risk o	of Bias	s	Applic	cabilit	ty Cor	cerns	•	
	Patient Selection	Index Test: TMPS	Reference Standard	Flow and Timing	Patient Selection	Index Test TMPS	Reference Standard			
Ashoor 2012	•	•	•		•	•	•			
Ashoor 2013	•	•	•	•	•	•	•			
Bevilacqua 2015	•	•	?		•	•	•			
Comas 2015	•	•	•	•	•	•	•			
del Mar Gil 2014	?	•	•		•	•	•			
Gil 2016	•	•				•	•			
Hall 2014	•	?	?		•	•	•			
Hooks 2014	•	?	•		•	•	•			
Jackson 2014	?	•	•	•	•	٠	•			
Korostelev 2014	•	•	?	•		•	•			
Nicolaides 2012	•	•	•	•	•	٠	•			
Nicolaides 2013	•	•	•		•	•	•			
Nicolaides 2014a	•	•	•		•	•	•			
Norton 2012	•	•	•		•	٠	•			
Norton 2015	•	•	•		•	•	•			
Pergament 2014	•	•	•	•	•	•	•			
Persico 2016	•	•	•		•	•	•			
Quezada 2015	•	•	?	•	•	•	•			
Samango-Sprouse 2013	•	•	•	•	•	•	•			
Sparks 2012a	•	?	•	?	•	•	•			
Verweij 2013	•	•	•		•	•	•			
😑 High	?	Uncle	ar		•	ow				
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Figure 5. Risk of bias and applicability concerns summary: review authors' judgements about each domain for each study included for targeted massively parallel sequencing (TMPS).

Figure 6. Risk of bias and applicability concerns (all tests included): review authors' judgements about each domains presented as percentages across included studies. MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing.



Risk of bias

No study was assessed as being at low risk of bias across all domains (Figure 4). For the patient selection domain, the 'Risk of bias' judgement was influenced mainly by inappropriate exclusions than the other signalling questions in this domain. Of the 61 studies judged to be at high risk of bias, 57 (93%) had inappropriate exclusions. The exclusions were mainly due to multifetal pregnancy, gestational age limits, and the prior risk of fetal aneuploidy. The remaining four (7%) studies were judged to be at unclear risk of bias (Figure 6).

In the index test domain, the risk of bias was considered to be low in 38 (58%) of the 44 MPSS studies and unclear in three (5%) studies. The remaining three (5%) MPSS studies were judged to be at high risk of bias because the index test was performed knowing the results of the reference standard or the threshold was not prespecified. The risk of bias was low in 18 (27%) of the 21 TMPS studies. The remaining three (5%) TMPS studies were judged to be at unclear risk of bias. All five studies that assessed traditional screening approaches were judged to be at low risk of bias for the index test domain (Figure 6).

In the reference standard domain, all studies used a reference standard likely to correctly classify the target condition. We considered 50 (77%) studies to be at low risk of bias because the studies stated that the reference standard results were interpreted without knowledge of the results of the index test. Of the remaining 15 studies, two (3%) studies were at high risk of bias because the reference standard was performed knowing the results of the index test while it was unclear what was done in the other 13 (20%) studies (Figure 6).

For the flow and timing domain, 46 (71%) studies were considered to be at high risk of bias because some pregnant women were excluded from 2 x 2 tables because gNIPT failed during the sequencing process. Fifteen (23%) studies were judged to be at low risk of bias. For the remaining four (6%) studies, information about the appropriate interval between the index test and reference standard was not provided (Figure 6).

Applicability concerns

We judged all studies to be of low applicability concern in the index test and reference standard domains because the studies matched the review question (Figure 4; Figure 6). All studies used a gNIPT method with ccfDNA in maternal blood and appropriate reference standard for the detection of common fetal aneuploidies. In the patient selection domain, 47 (72%) studies included cohort of pregnant women selected at high risk of fetal aneuploidy or cohort of unselected pregnant women and were judged to be of low applicability concern. In the other 18 (28%) studies, the cohorts comprised pregnant women with different prior risk of fetal aneuploidy (mixed risk cohorts). This population did not represent the real-life situation and those cohorts were judged to be of high applicability concern.

Findings

The characteristics of the studies are summarised in Table 4 and Summary of findings 1. Results are presented separately for each of the main fetal aneuploidies (T21, T18, T13 and 45,X) and globally for all autosomes or all sex chromosome aneuploidies (SCA) combined (Summary of findings 2; Summary of findings 3; Summary of findings 4; Summary of findings 5; Summary of findings 6; Summary of findings 7). For each aneuploidy, results are presented according to the prior risk of chromosomal abnormality as high risk or unselected population and according to MPSS and TMPS methods. Results from mixed-risk populations are summarised in Appendix 7. No study directly compared the accuracy of MPSS and TMPS. There were insufficient data to separately consider monochorionic and dichorionic pregnancies and four of the nine studies did not report chorionicity.

1. Trisomy 21 (T21 or Down syndrome)

A total of 57 studies assessed gNIPT for T21 in 2004 affected and 80,616 non T21 pregnancies. Five studies enrolled an unselected population of pregnant women undergoing aneuploidy screening, 36 studies enrolled pregnant women selected at high risk of fetal aneuploidy and 16 studies enrolled pregnant women with various prior risk and no a priori risk of fetal aneuploidy (mixed risk). Of the 57 studies, 41 assessed MPSS and 16 assessed TMPS. The results are summarised in Summary of findings 2.



a. Unselected population of pregnant women undergoing aneuploidy screening

Five cohort studies evaluated gNIPT in an unselected population of pregnant women undergoing aneuploidy screening. The studies

included 22,412 non T21 pregnancies and 96 (0.43%) T21 cases. MPSS was assessed in one study and TMPS was assessed in four studies (Figure 7).

Figure 7. Forest plot of MPSS and TMPS for T21 in unselected pregnant women undergoing aneuploidy screening. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS T21											
Study TP	FP	FN	TN	Case	control	Sensi	itivity (95% CI)	Speci	ficity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Song 2013 8	0	0	1733		No	1.	00 [0.63, 1.00]	1.	00 [1.00, 1.00]		
MPSS T18										0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study TP	FP	FN	TN	Case	control	Sensi	itivity (95% CI)	Speci	ficity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Song 2013 2	1	0	1738		No	1.	00 [0.16, 1.00]	1.	00 [1.00, 1.00]		
MPSS T13										0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study TP	FP	FN	TN	Case	-control	Sensi	itivity (95% CI)	Speci	ficity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Song 2013 1	0	0	1740		No	1.	00 [0.03, 1.00]	1.	00 [1.00, 1.00]		
TMPS T21										0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study	TP	FP	FN	TN	Case-c	ontrol	Sensitivity (9	5% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Norton 2015	38	9	0	15794		No	1.00 (0.91		1.00 [1.00, 1.00]	-	
Quezada 2015	32	: 1	0	2752		No	1.00 (0.89		1.00 [1.00, 1.00]		
del Mar Gil 2014	9	0	1	182		No	0.90 [0.55	, 1.00]	1.00 [0.98, 1.00]		•
Nicolaides 2012	8	0	0	1941		No	1.00 [0.63	, 1.00]	1.00 [1.00, 1.00]		
TMPS T18										0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study	TP	FP	FN	TN	Case-c	ontrol	Sensitivity (9	5% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Norton 2015	9	1	1	15830		No	0.90 (0.55	. 1.00]	1.00 [1.00, 1.00]		•
Quezada 2015	9	5	1	2770		No	0.90 (0.55		1.00 [1.00, 1.00]		
Nicolaides 2012	2	2	0	1945		No	1.00 [0.16	1.00]	1.00 [1.00, 1.00]		
TMPS T13										0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study	TP	FP	FN	TN	Case-c	ontrol	Sensitivity (9	5% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Norton 2015	2	_	-	11181		No	1.00 [0.16		1.00 [1.00, 1.00]		•
Quezada 2015	2			2778		No	0.40 [0.05	• •	1.00 [1.00, 1.00]		•
del Mar Gil 2014	1	0	0	191		No	1.00 [0.03	, 1.00]	1.00 [0.98, 1.00]		
											1 112 017 010 010 1

i. MPSS

One prospective cohort study included eight T21 cases and 1733 non T21 pregnancies (Song 2013). The sensitivity (95% confidence interval (CI)) of MPSS was 100% (67.6% to 100%) and the specificity (95% CI) was 100% (99.8% to 100%).

ii. TMPS

TMPS was evaluated in four studies comprising 20,679 non T21 pregnancies and 88 T21 cases (del Mar Gil 2014; Nicolaides 2012; Norton 2015; Quezada 2015). The summary sensitivity (95% CI) was 99.2% (78.2% to 100%) and the summary specificity (95% CI) was 100% (> 99.9% to 100%).

iii. Comparative accuracy of MPSS and TMPS

It was not possible to compare the accuracy of MPSS and TMPS in a meta-analysis because of limited data.

b. Selected population of pregnant women at high risk of fetal aneuploidy

Overall, 36 studies included pregnant women selected at high risk of fetal aneuploidy involving 20,317 non T21 pregnancies and 1294 (6.37%) T21 cases. MPSS was assessed in 30 studies and TMPS in six studies (Figure 8).

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Figure 8. Forest plot of MPSS and TMPS for T21 in pregnant women selected at high risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MDSS	T21
	-

Study	ТР	FP	FN	TN	Sensitivity (95% Cl	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% Cl)
Johansen 2016	27	1	1	144	0.96 [0.82, 1.00	0.99 [0.96, 1.00]		•
Kim 2016	5	0	0	96	1.00 [0.48, 1.00	1.00 [0.96, 1.00]		e
Lefkowitz 2016	84	1	0	1081	1.00 [0.96, 1.00	1.00 [0.99, 1.00]		
Papageorghiou 2016a	42	0	0	384	1.00 [0.92, 1.00	1.00 [0.99, 1.00]	-	• • •
Papageorghiou 2016b	1	0	0	10	1.00 [0.03, 1.00	1.00 [0.69, 1.00]		·•
Poon 2016	35	0	0	206	1.00 (0.90, 1.00	1.00 [0.98, 1.00]		
Zhang 2016	3	0	0	84	1.00 [0.29, 1.00	1.00 [0.96, 1.00]		
Alberti 2015	47	0	0	136	1.00 [0.92, 1.00	1.00 [0.97, 1.00]	-	
Benachi 2015	76	1	0	809	1.00 [0.95, 1.00	1.00 [0.99, 1.00]	-	
Ke 2015	17	0	0	2323	1.00 [0.80, 1.00	1.00 [1.00, 1.00]		
Lee 2015	5	0	0	87	1.00 [0.48, 1.00	1.00 [0.96, 1.00]		
Song 2015	2	0	0	202	1.00 [0.16, 1.00	1.00 [0.98, 1.00]		
Sukhikh 2015	17	0	2	181	0.89 [0.67, 0.99	1.00 [0.98, 1.00]		
Sung-Hee 2015	4	0	0	897	1.00 [0.40, 1.00	1.00 [1.00, 1.00]		
Wang 2015a	25	0	0	892	1.00 [0.86, 1.00	1.00 [1.00, 1.00]		
Huang 2014	9	0	0	180	1.00 [0.66, 1.00	1.00 [0.98, 1.00]		
Jeon 2014	11	0	0	144	1.00 [0.72, 1.00	1.00 [0.97, 1.00]		
Porreco 2014	137	3	0	3182	1.00 [0.97, 1.00	1.00 [1.00, 1.00]	•	
Stumm 2014	40	0	2	430	0.95 [0.84, 0.99	1.00 [0.99, 1.00]		
Wang 2014	3	0	0	133	1.00 [0.29, 1.00	1.00 [0.97, 1.00]		
Bianchi 2013	30	1	0	82	1.00 [0.88, 1.00	0.99 [0.93, 1.00]		•
Liang 2013	40	0	0	372	1.00 [0.91, 1.00	1.00 [0.99, 1.00]	-	
Bianchi 2012	93	6	0	404	1.00 [0.96, 1.00	0.99 [0.97, 0.99]	-	
Canick 2012	7	0	0	20	1.00 [0.59, 1.00	1.00 [0.83, 1.00]		
Hou 2012	2	0	0	203	1.00 [0.16, 1.00	1.00 [0.98, 1.00]		
Jiang 2012	16	0	0	887	1.00 [0.79, 1.00	1.00 [1.00, 1.00]		•
Liu 2012	1	1	0	151	1.00 [0.03, 1.00	0.99 [0.96, 1.00]		• •
Palomaki 2012	210	1	2	1758	0.99 [0.97, 1.00	1.00 [1.00, 1.00]	•	•
Ehrich 2011	39	1	0	409	1.00 [0.91, 1.00	1.00 [0.99, 1.00]	-	•
Sehnert 2011	13	0	0	34	1.00 [0.75, 1.00	1.00 [0.90, 1.00]		
							0 0.2 0.4 0.6 0.8 1	
TMPS T21								
Study TP	FP F	N	TN	Sensi	tivity (95% CI) Spec	ificity (95% Cl)	Sensitivity (95% CI)	Specificity (95% CI)
Persico 2016 35	0	1	213	0.9	97 [0.85, 1.00] 1	.00 [0.98, 1.00]		
Nicolaides 2013 25	0	0	204	1.0	00 [0.86, 1.00] 1	.00 [0.98, 1.00]		•
Verweij 2013 17	0	1	486	0.9	94 [0.73, 1.00] 1	.00 [0.99, 1.00]		
Ashoor 2012 50	0	0	347	1.0	00 [0.93, 1.00] 1	.00 [0.99, 1.00]	-	•
Norton 2012 81	1	0 2	998	1.0	00 [0.96, 1.00] 1	.00 [1.00, 1.00]	-	•
Sparks 2012a 36	0	0	131	1.0	00 [0.90, 1.00] 1	.00 [0.97, 1.00]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

i. MPSS

The 30 MPSS studies included 15,937 non T21 pregnancies and 1048 T21 cases (Alberti 2015; Benachi 2015; Bianchi 2012; Bianchi 2013; Canick 2012; Ehrich 2011; Hou 2012; Huang 2014; Jeon 2014; Jiang 2012; Johansen 2016; Ke 2015; Kim 2016; Lee 2015; Lefkowitz 2016; Liang 2013; Liu 2012; Palomaki 2012; Papageorghiou 2016a; Papageorghiou 2016b; Poon 2016; Porreco 2014; Sehnert 2011; Song 2015; Stumm 2014; Sukhikh 2015; Sung-Hee 2015; Wang 2014; Wang 2015a; Zhang 2016). The summary sensitivity (95% CI) was 99.7% (98.0% to 100%) and the summary specificity (95% CI) was 99.9% (99.8% to 100%).

ii. TMPS

Six studies evaluated TMPS in 4380 non T21 pregnancies and 246 T21 cases (Ashoor 2012; Nicolaides 2013; Norton 2012; Persico 2016; Sparks 2012a; Verweij 2013). The summary sensitivity (95% CI) was 99.2% (96.8% to 99.8%) and the summary specificity (95% CI) was 100% (99.8% to 100%).

iii. Comparative accuracy of MPSS and TMPS

An indirect comparison of the 30 MPSS and six TMPS studies showed no statistical evidence of a difference in sensitivity or specificity or both (P value = 0.52). The differences in sensitivity and specificity were negligible (Summary of findings 2).



2. Trisomy 18 (T18)

ii. TMPS

Fifty studies assessed T18 in 634 cases and 78,688 non T18 pregnancies. Four studies enrolled unselected population of pregnant women undergoing aneuploidy screening, 33 studies enrolled pregnant women selected at high risk of fetal aneuploidy and 13 studies enrolled a cohort with mixed prior risk. Of the 50 studies, 38 evaluated MPSS and 12 evaluated TMPS. The results are summarised in Summary of findings 3.

a. Unselected population of pregnant women undergoing aneuploidy screening

Four studies, comprising 22,292 non T18 pregnancies and 24 (0.11%) T18 cases, assessed gNIPT for fetal aneuploidy in unselected pregnant women. One study assessed MPSS and three studies assessed TMPS (Figure 7).

i. MPSS

One MPSS study evaluated two T18 cases and 1739 non T18 pregnancies (Song 2013). The sensitivity (95% CI) was 100% (34.3% to 100%) and the specificity (95% CI) was 99.9% (99.7% to 100%).

Three studies evaluated TMPS in 20,553 non T18 pregnancies and 22 T18 cases (Nicolaides 2012; Norton 2015; Quezada 2015). The summary sensitivity (95% CI) was 90.9% (70.0% to 97.7%) and the summary specificity (95% CI) was 100% (99.9% to 100%).

iii. Comparative accuracy of MPSS and TMPS

It was not possible to compare the accuracy of MPSS and TMPS in a meta-analysis because data were sparse.

b. Selected population of pregnant women at high risk of fetal aneuploidy

A total of 33 studies included pregnant women selected at high risk of fetal aneuploidy involving 444 (2.20%) T18 cases and 20,190 non T18 pregnancies. Of these, 28 studies assessed MPSS and five studies assessed TMPS (Figure 9). Figure 9. Forest plot of MPSS and TMPS for T18 in pregnant women selected at high risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS T18

Study	TP	FP	FN	TN	Sensitivity (95% C	l) Specificity (95% Cl)	Sensitivity (95% CI)	Specificity (95% CI)
Johansen 2016	4	0	0	169	1.00 [0.40, 1.0))] 1.00 [0.98, 1.00]		
Lefkowitz 2016	27	0	0	1139	1.00 [0.87, 1.0)] 1.00 [1.00, 1.00]		L
Papageorghiou 2016a	9	0	0	417	1.00 [0.66, 1.0)] 1.00 [0.99, 1.00]		
Papageorghiou 2016b	1	0	0	10	1.00 [0.03, 1.0))] 1.00 [0.69, 1.00]		·
Poon 2016	4	0	0	237	1.00 [0.40, 1.0))] 1.00 [0.98, 1.00]		•
Zhang 2016	1	0	0	86	1.00 [0.03, 1.0))] 1.00 [0.96, 1.00]		
Benachi 2015	22	1	3	860	0.88 [0.69, 0.9]	7] 1.00 [0.99, 1.00]		
Ke 2015	6	0	0	2324	1.00 [0.54, 1.0)] 1.00 [1.00, 1.00]		
Lee 2015	2	0	0	90	1.00 [0.16, 1.0)			
Song 2015	1	0	0	203	1.00 [0.03, 1.0))] 1.00 [0.98, 1.00]		
Sukhikh 2015	8	0	0	192	1.00 [0.63, 1.0))] 1.00 [0.98, 1.00]		•
Sung-Hee 2015	2	0	0	899	1.00 [0.16, 1.0))] 1.00 [1.00, 1.00]		
Wang 2015a	3	1	0	913	1.00 [0.29, 1.0)] 1.00 [0.99, 1.00]		•
Bijok 2014	1	0	0	8	1.00 [0.03, 1.0)] 1.00 [0.63, 1.00]		· · · · •
Huang 2014	1	0	1	187	0.50 [0.01, 0.9	1.00 [0.98, 1.00]		
Jeon 2014	5	0	0	150	1.00 [0.48, 1.0)] 1.00 [0.98, 1.00]		
Porreco 2014	36	0	3	3283	0.92 [0.79, 0.9]	3] 1.00 [1.00, 1.00]		
Stumm 2014	8	1	0	463	1.00 [0.63, 1.0)] 1.00 [0.99, 1.00]		
Wang 2014	1	0	0	135	1.00 [0.03, 1.0))] 1.00 [0.97, 1.00]		•
Bianchi 2013	10	0	0	103	1.00 [0.69, 1.0)] 1.00 [0.96, 1.00]		•
Liang 2013	14	0	0	398	1.00 [0.77, 1.0			
Bianchi 2012	38	3	1	460	0.97 [0.87, 1.0])] 0.99 [0.98, 1.00]		
Hou 2012	3	0	0	202	1.00 [0.29, 1.0))] 1.00 [0.98, 1.00]		•
Jiang 2012	12	1	0	890	1.00 [0.74, 1.0))] 1.00 [0.99, 1.00]		•
Liu 2012	1	0	0	152	1.00 [0.03, 1.0))] 1.00 [0.98, 1.00]		•
Palomaki 2012	59	- 5	0	1907	1.00 [0.94, 1.0))] 1.00 [0.99, 1.00]	-	•
Chen 2011	34	- 5	3	247	0.92 [0.78, 0.9)	3] 0.98 (0.95, 0.99)		-
Sehnert 2011	8	0	0	39	1.00 [0.63, 1.0)] 1.00 [0.91, 1.00]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
TMPS T18								
Study TP	FP I	FN	TN	Sens	itivity (95% Cl) Sp	ecificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Persico 2016 13	0	0	236	1.	.00 [0.75, 1.00]	1.00 [0.98, 1.00]		
Nicolaides 2013 3	0	0	226	1.	.00 [0.29, 1.00]	1.00 [0.98, 1.00]		
Ashoor 2012 49	0	1	347	0	.98 [0.89, 1.00]	1.00 [0.99, 1.00]	-4	
Norton 2012 37	2	1	3040		.97 [0.86, 1.00]	1.00 [1.00, 1.00]		
Sparks 2012a 8	0	0	159		.00 [0.63, 1.00]	1.00 [0.98, 1.00]	· · · · · · · · · · · · · · · · · · ·	
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

i. MPSS

Twenty-eight studies evaluated MPSS in 16,180 non T18 pregnancies and 332 T18 cases (Benachi 2015; Bianchi 2012; Bianchi 2013; Bijok 2014; Chen 2011; Hou 2012; Huang 2014; Jeon 2014; Jiang 2012; Johansen 2016; Ke 2015; Lee 2015; Lefkowitz 2016; Liang 2013; Liu 2012; Palomaki 2012; Papageorghiou 2016a; Papageorghiou 2016b; Poon 2016; Porreco 2014; Sehnert 2011; Song 2015; Stumm 2014; Sukhikh 2015; Sung-Hee 2015; Wang 2014; Wang 2015a; Zhang 2016). The summary sensitivity (95% CI) was 97.8% (92.5% to 99.4%) and the summary specificity (95% CI) was 99.9% (99.8% to 100%).

ii. TMPS

Five studies evaluated TMPS in 4010 non T18 pregnancies and 112 T18 cases (Ashoor 2012; Nicolaides 2013; Norton 2012; Persico 2016; Sparks 2012a). The summary sensitivity (95% CI) was 98.2% (93.1% to 99.6%) and the summary specificity (95% CI) was 100% (99.8% to 100%).

iii. Comparative accuracy of MPSS and TMPS

An indirect comparison of the 28 MPSS and five TMPS studies showed no statistical evidence of a difference in sensitivity, specificity or both (P value = 0.47). The differences in sensitivity and specificity were negligible (Summary of findings 3).

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3. Trisomy 13 (T13)

ii. TMPS

T13 was assessed in 39 studies comprising 215 affected and 68,743 non T13 pregnancies. Four studies evaluated unselected population of pregnant women undergoing fetal aneuploidy screening, while 22 studies evaluated women at high risk of fetal aneuploidy and 13 studies evaluated mixed prior risk cohorts. Of the 39 studies, 29 assessed MPSS and 10 assessed TMPS. The results are summarised in Summary of findings 4.

a. Unselected population of pregnant women undergoing aneuploidy screening

Four studies assessed gNIPT for T13 in unselected pregnant women. The studies included 15,894 non T13 pregnancies and nine (0.06%) T13 cases. Three studies evaluated TMPS and one study evaluated MPSS (Figure 7).

i. MPSS

One study evaluated MPSS in one T13 case and 1740 non T13 pregnancies (Song 2013). The sensitivity (95% CI) was 100% (20.7% to 100%) and the specificity (95% CI) was 100% (99.8% to 100%).

Three studies evaluated TMPS in 14,154 non T13 pregnancies and eight T13 cases (del Mar Gil 2014; Norton 2015; Quezada 2015). The summary sensitivity (95% CI) was 65.1% (9.2% to 97.2%) and the summary specificity (95% CI) was 100% (99.9% to 100%).

iii. Comparative accuracy of MPSS and TMPS

It was not possible to compare the accuracy of MPSS and TMPS in a meta-analysis because data were sparse.

b. Selected population of pregnant women at high risk of fetal aneuploidy

A total of 22 studies evaluated pregnant women selected at high risk of fetal aneuploidy. The studies included 14,103 non T13 pregnancies and 148 (1.05%) T13 cases. Twenty studies assessed MPSS and two studies assessed TMPS (Figure 10).

Figure 10. Forest plot of MPSS and TMPS for T13 in pregnant women selected at high risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS T13

Study	ТР	FP	FN	TN	Sonsitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% Cl)
2							Sensitivity (95% CI)	specificity (95% Ci)
Johansen 2016	3	0	0	170	1.00 [0.29, 1.00]	1.00 [0.98, 1.00]		
Lefkowitz 2016	15	0	0	1151	1.00 [0.78, 1.00]	1.00 [1.00, 1.00]		
Papageorghiou 2016a	5	0	0	421	1.00 [0.48, 1.00]	1.00 [0.99, 1.00]		
Poon 2016	2	0	0	239	1.00 [0.16, 1.00]	1.00 [0.98, 1.00]		
Benachi 2015	12	1	0	873	1.00 [0.74, 1.00]	1.00 [0.99, 1.00]		
Ke 2015	1	0	0	2339	1.00 [0.03, 1.00]	1.00 [1.00, 1.00]		
Lee 2015	1	0	0	91	1.00 [0.03, 1.00]	1.00 [0.96, 1.00]		•
Song 2015	1	0	0	203	1.00 [0.03, 1.00]	1.00 [0.98, 1.00]		-
Sukhikh 2015	1	1	0	198	1.00 [0.03, 1.00]	0.99 [0.97, 1.00]		-
Porreco 2014	14	0	2	3306	0.88 [0.62, 0.98]	1.00 [1.00, 1.00]		-
Stumm 2014	5	0	0	467	1.00 [0.48, 1.00]	1.00 [0.99, 1.00]		-
Bianchi 2013	3	0	1	109	0.75 [0.19, 0.99]	1.00 [0.97, 1.00]	_	-
Liang 2013	4	1	0	407	1.00 [0.40, 1.00]	1.00 [0.99, 1.00]		
Bianchi 2012	13	0	3	485	0.81 [0.54, 0.96]	1.00 [0.99, 1.00]		-
Canick 2012	1	0	0	26	1.00 [0.03, 1.00]	1.00 [0.87, 1.00]		
Jiang 2012	2	0	0	901	1.00 [0.16, 1.00]	1.00 [1.00, 1.00]		-
Liu 2012	1	0	0	152	1.00 [0.03, 1.00]	1.00 [0.98, 1.00]		
Palomaki 2012	11	16	1	1943	0.92 [0.62, 1.00]	0.99 [0.99, 1.00]		
Chen 2011	25	3	0	261	1.00 [0.86, 1.00]	0.99 [0.97, 1.00]		
Sehnert 2011	1	0	0	46	1.00 [0.03, 1.00]	1.00 [0.92, 1.00]		
0000002011	•	Ŭ	Ŭ		1.00 [0.00] 1.00]			
TMPS T13							0 0.2 0.4 0.0 0.0 1	0 0.2 0.4 0.0 0.0 1
Study TP FF	FN	TN	Se	ensitivit	y (95% CI) Specificit	y (95% Cl)	Sensitivity (95% CI)	Specificity (95% CI)
Persico 2016 5 () ()	244		1.00 [0.48, 1.00] 1.00 [0.98, 1.00]		
Hall 2014 15 0) ()	49		•		0.93, 1.00]	· · · · · · •	
				•			0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

i. MPSS

Twenty studies evaluated MPSS in 13,810 non T13 pregnancies and 128 T13 cases (Benachi 2015; Bianchi 2012; Bianchi 2013;

Canick 2012; Chen 2011; Jiang 2012; Johansen 2016; Ke 2015; Lee 2015; Lefkowitz 2016; Liang 2013; Liu 2012; Palomaki 2012; Papageorghiou 2016a; Poon 2016; Porreco 2014; Sehnert 2011;



Song 2015; Stumm 2014; Sukhikh 2015). The summary sensitivity (95% CI) was 95.8% (86.1% to 98.9%) and the summary specificity (95% CI) was 99.8% (99.8% to 99.9%).

ii. TMPS

Two studies evaluated TMPS in 293 non T13 pregnancies and 20 T13 cases (Hall 2014; Persico 2016). The summary sensitivity (95% Cl) was 100% (83.9% to 100%) and the summary specificity (95% Cl) was 100% (98.7% to 100%).

iii. Comparative accuracy of MPSS and TMPS

It was not possible to compare the accuracy of MPSS and TMPS in a meta-analysis because data were sparse.

4. Turner syndrome (45,X)

Turner syndrome (45,X) was assessed in 20 studies, comprising 232 affected and 9849 non 45,X pregnancies. Among these studies, 16

enrolled pregnant women selected at high risk of fetal aneuploidy and four enrolled a cohort of pregnant women with mixed prior risk. Of the 20 studies, 14 evaluated MPSS and six evaluated TMPS. The results are summarised in Summary of findings 5.

a. Unselected population of pregnant women undergoing aneuploidy screening

No study assessed 45,X in this population.

b. Selected population of pregnant women at high risk of fetal aneuploidy

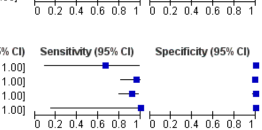
Sixteen studies included 198 (2.35%) affected and 8421 non 45,X pregnancies. MPSS and TMPS were assessed by 12 and four studies respectively (Figure 11).

Figure 11. Forest plot of MPSS and TMPS for 45,X in pregnant women selected at high risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS 45,X

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Lefkowitz 2016	19	- 7	0	1118	1.00 [0.82, 1.00]	0.99 [0.99, 1.00]		•
Song 2015	0	1	1	202	0.00 [0.00, 0.97]	1.00 [0.97, 1.00]		•
Sukhikh 2015	4	1	0	195	1.00 [0.40, 1.00]	0.99 [0.97, 1.00]		•
Porreco 2014	9	11	0	3258	1.00 [0.66, 1.00]	1.00 [0.99, 1.00]		•
Bianchi 2013	20	0	1	92	0.95 [0.76, 1.00]	1.00 [0.96, 1.00]		•
Liang 2013	5	1	3	403	0.63 [0.24, 0.91]	1.00 [0.99, 1.00]	_	•
Mazloom 2013	20	1	1	389	0.95 [0.76, 1.00]	1.00 [0.99, 1.00]		•
Bianchi 2012	21	46	6	416	0.78 [0.58, 0.91]	0.90 [0.87, 0.93]		•
Hou 2012	1	1	0	203	1.00 [0.03, 1.00]	1.00 [0.97, 1.00]		•
Jiang 2012	3	1	1	898	0.75 [0.19, 0.99]	1.00 [0.99, 1.00]	_	•
Liu 2012	1	0	0	152	1.00 [0.03, 1.00]	1.00 [0.98, 1.00]		•
Sehnert 2011	3	1	0	43	1.00 [0.29, 1.00]	0.98 [0.88, 1.00]		
TMPS 45,X							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study	٦	I PI	FP I	N TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)

Study					Sensitivity (55% Ci)	specificity (55
Persico 2016	2	0	1	246	0.67 [0.09, 0.99]	1.00 [0.99, 1
Hooks 2014	26	2	1	385	0.96 [0.81, 1.00]	0.99 [0.98, 1
Nicolaides 2014a	43	0	4	125	0.91 [0.80, 0.98]	1.00 [0.97, 1
Nicolaides 2013	2	0	0	227	1.00 [0.16, 1.00]	1.00 [0.98, 1



i. MPSS

Twelve studies evaluated MPSS in 119 affected and 7440 non 45,X pregnancies (Bianchi 2012; Bianchi 2013; Hou 2012; Jiang 2012; Lefkowitz 2016; Liang 2013; Liu 2012; Mazloom 2013; Porreco 2014; Sehnert 2011; Song 2015; Sukhikh 2015). The summary sensitivity (95% CI) was 91.7% (78.3% to 97.1%) and the summary specificity (95% CI) was 99.6% (98.9% to 99.8%).

ii. TMPS

Four studies evaluated TMPS in 79 affected and 985 non 45,X pregnancies (Hooks 2014; Nicolaides 2013; Nicolaides 2014a; Persico 2016). The summary sensitivity (95% CI) was 92.4% (84.1% to 96.5%) and the summary specificity (95% CI) was 99.8% (98.3% to 100%).

iii. Comparative accuracy of MPSS and TMPS

An indirect comparison of the 12 MPSS and four TMPS studies showed no statistical evidence of a difference in sensitivity,



specificity or both (P value = 0.40). The differences in sensitivity and specificity were negligible (Summary of findings 5).

5. Triple X syndrome (47,XXX)

Seven studies assessed 47,XXX, comprising 14 (0.23%) affected and 6021 non 47,XXX pregnancies (Hooks 2014; Lefkowitz 2016; Liang

2013; Mazloom 2013; Nicolaides 2014a; Porreco 2014; Song 2015). The studies enrolled pregnant women selected at high risk of fetal aneuploidy. Five studies evaluated MPSS and two studies evaluated TMPS. (Figure 12; Table 8). We did not perform a separate metaanalysis for 47,XXX due to sparse data (very few cases or studies, or one or more subgroups had no study).

Figure 12. Forest plot of MPSS and TMPS for 47,XXX, 47,XXY and 47,XYY in pregnant women selected at high risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS 47, XXX									
Study	ТР	FP	FN		TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% Cl)	Specificity (95% CI)
Lefkowitz 2016	1	0	0	11	43	1.00 [0.03, 1.00]	1.00 [1.00, 1.00]		
Song 2015	0	0	1	2	203	0.00 [0.00, 0.97]	1.00 [0.98, 1.00]		•
Porreco 2014	4	3	0	32	271	1.00 [0.40, 1.00]	1.00 [1.00, 1.00]		■ .
Liang 2013	1	0	0	4	111	1.00 [0.03, 1.00]	1.00 [0.99, 1.00]		• • • • • • • • • • • • • • • • • • •
Mazloom 2013	1	0	0	4	10	1.00 [0.03, 1.00]	1.00 [0.99, 1.00]		
MPSS 47,XXY								0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study	ТР	FP	FN		TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Lefkowitz 2016	3	0	0	11	41	1.00 [0.29, 1.00]	1.00 [1.00, 1.00]		• • • • • • • • • • • • • • • • • • •
Song 2015	0	0	1	2	203	0.00 [0.00, 0.97]	1.00 [0.98, 1.00]		
Porreco 2014	1	2	0	31	98	1.00 [0.03, 1.00]	1.00 [1.00, 1.00]		• • • • • • • • • • • • • • • • • • •
Liang 2013	1	0	0		111	1.00 [0.03, 1.00]	1.00 [0.99, 1.00]		•
Mazloom 2013	5	Ō	0		106	1.00 [0.48, 1.00]	1.00 [0.99, 1.00]		•
Hou 2012	Ō	Ō	1		204	0.00 [0.00, 0.97]	1.00 [0.98, 1.00]		
Jiang 2012	2	Ő	0		901	1.00 [0.16, 1.00]	1.00 [1.00, 1.00]		
ording 2012	2				.01	1.00 [0.10, 1.00]	1.00 [1.00, 1.00]		
MPSS 47,XYY								0 0.2 0.1 0.0 0.0 1	0 0.2 0.1 0.0 0.0 1
Study	TP	FP	FN		TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Lefkowitz 2016	3	0	0	11	41	1.00 [0.29, 1.00]	1.00 [1.00, 1.00]		• • • • • • • • • • • • • • • • • • •
Porreco 2014	1	0	0	32	200	1.00 [0.03, 1.00]	1.00 [1.00, 1.00]		•
Liang 2013	1	0	0	4	111	1.00 [0.03, 1.00]	1.00 [0.99, 1.00]		•
Mazloom 2013	3	0	0	4	108	1.00 [0.29, 1.00]	1.00 [0.99, 1.00]		
Hou 2012	0	0	1	2	204	0.00 [0.00, 0.97]	1.00 [0.98, 1.00]		
Jiang 2012	1	0	Ó		902	1.00 [0.03, 1.00]	1.00 [1.00, 1.00]		•
Liu 2012	1	0	0		52	1.00 [0.03, 1.00]	1.00 [0.98, 1.00]		
		-	Ĩ					0 0.2 0.4 0.6 0.8 1	
TMPS 47,XXX									
Study		TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Hooks 2014		1	2		411	1.00 [0.03, 1.00]	1.00 [0.98, 1.00]		•
Nicolaides 2014	а	5	1	0	166	1.00 [0.48, 1.00]	0.99 [0.97, 1.00]		
TMPS 47,XXY								0 0.2 0.4 0.6 0.8 1	1 1 1 1 1 1 1 0 0.2 0.4 0.6 0.8 1
Study		TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Persico 2016		1	0	0	248	1.00 [0.03, 1.00]		,	
Hooks 2014		6	0	0	408	1.00 [0.54, 1.00]			
	_		-	-	171				
Nicolaides 2014	a	1	U	U	171	1.00 [0.03, 1.00]	1.00 [0.98, 1.00]		
TMPS 47,XYY								0 0.2 0.4 0.0 0.0 1	0 0.2 0.4 0.0 0.8 1
Study		TP	FP	FN	TN	Sensitivity (95% Cl)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Nicolaides 2014	а	3	0	0	169	1.00 [0.29, 1.00]	1.00 [0.98, 1.00]	· · · · · · · · · · · · · · · · · · ·	· · · · · · ·
		-	-	-				0 0.2 0.4 0.6 0.8 1	

6. Klinefelter syndrome (47,XXY)

Twelve studies assessed 47,XXY in 25 (0.33%) affected and 7584 non 47,XXY pregnancies (Hooks 2014; Hou 2012; Jiang 2012; Lau 2012; Lefkowitz 2016; Liang 2013; Mazloom 2013; Nicolaides 2014a; Persico 2016; Porreco 2014; Samango-Sprouse 2013; Song 2015).

Ten studies enrolled pregnant women selected at high risk of fetal aneuploidy (Figure 12; Table 8) and two studies enrolled pregnant women with mixed risk (See Finding section 11). No study assessed 47,XXY in an unselected population of pregnant women undergoing aneuploidy screening. Eight studies assessed MPSS



MPSS. autosomes

and four studies assessed TMPS. We did not perform a separate meta-analysis for 47,XXY due to sparse data (very few cases or studies, or one or more subgroups had no study).

7.47,XYY

Ten studies assessed 47,XYY in 16 (0.23%) affected and 6971 non 47,XYY pregnancies (Hou 2012; Jiang 2012; Lefkowitz 2016; Liang 2013; Liu 2012; Mazloom 2013; Nicolaides 2014a; Porreco 2014; Samango-Sprouse 2013; Shaw 2014). Eight studies enrolled pregnant women selected at high risk of fetal aneuploidy (Figure 12; Table 8) and two studies enrolled pregnant women with mixed risk (See Finding section 11). Eight studies used MPSS and two studies used TMPS. We did not perform a separate meta-analysis for 47,XXX due to sparse data (very few cases or studies, or one or more subgroups had no study).

8. All autosomes combined

Autosomal aneuploidies were assessed in 61 studies. The studies included 84,954 pregnant women of which 2853 were T21, T18 or T13 pregnancies and 82,073 were unaffected. Among these 61 studies, 43 assessed MPSS and 18 assessed TMPS. Of the 61 studies, five enrolled unselected pregnant women, 39 enrolled high-risk pregnant women and 17 enrolled a cohort of mixed prior risk. The results are summarised in Summary of findings 6. The results for mixed risk cohorts are summarised in Appendix 7.

a. Unselected population of pregnant women undergoing aneuploidy screening

Five studies assessed 129 (0.58%) affected and 22,379 unaffected (non T21, T18 and T13) pregnancies. Of the five studies, one study assessed MPSS and four studies assessed TMPS (Figure 13).

Figure 13. Forest plot of MPSS and TMPS for autosomes (T21, T18 and T13 combined) in unselected pregnant women undergoing aneuploidy screening. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

Study Song 2013	TP 11	FP 1	FN O	TN 1729		3. , ,	ificity (95% Cl) 00 [1.00, 1.00]	Sensitivity (95% CI)	Specificity (95% Cl)
TMPS, autoso	me	5							
Study		TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Norton 2015		49	12	1	15779	0.98 [0.89, 1.00]	1.00 [1.00, 1.00]		
Quezada 2016	5	43	8	4	2730	0.91 [0.80, 0.98]	1.00 [0.99, 1.00]		
del Mar Gil 20	14	10	0	1	181	0.91 [0.59, 1.00]	1.00 [0.98, 1.00]		
Nicolaides 20	12	10	2	0	1937	1.00 [0.69, 1.00]	1.00 [1.00, 1.00]		

i. MPSS

Only one study assessed MPSS (Song 2013). The study evaluated 1730 unaffected (non T21, T18 and T13) pregnancies and 11 cases in women with singleton pregnancy. The sensitivity (95% CI) was 100% (74.1% to 100%) and the specificity (95% CI) was 99.9% (99.7% to 100%).

ii. TMPS

Four studies assessed TMPS in 20,649 unaffected (non T21, T18 and T13) pregnancies and 118 cases (del Mar Gil 2014; Nicolaides 2012; Norton 2015; Quezada 2015). Of the four studies, three studies included only women with singleton pregnancy and the remaining study included only women with multifetal pregnancy (Table 9).

Based on the four studies, the summary sensitivity (95% CI) was 94.9% (89.1% to 97.7%) and the summary specificity (95% CI) was 99.9% (99.8% to 99.9%).

iii. Comparative accuracy of MPSS and TMPS

It was not possible to compare the accuracy of MPSS and TMPS in a meta-analysis due to limited data.

b. Selected population of pregnant women at high risk of fetal aneuploidy

A total of 39 studies included 1886 (9.39%) affected and 20,079 unaffected (non T21, T18 and T13) pregnancies. Of the 39 studies, 32 assessed MPSS and seven assessed TMPS (Figure 14).

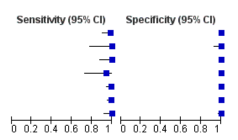
Figure 14. Forest plot of MPSS and TMPS for autosomes (T21, T18 and T13) in pregnant women selected at high risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS, autosomes

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Johansen 2016	34	1	1	137	0.97 [0.85, 1.00]	0.99 [0.96, 1.00]		
Kim 2016	5	Ó	Ó	96	1.00 [0.48, 1.00]	1.00 [0.96, 1.00]		•
Lefkowitz 2016	126	1	Õ	1039	1.00 [0.97, 1.00]	1.00 [0.99, 1.00]		
Papageorghiou 2016a	56	Ó	-	370	1.00 [0.94, 1.00]	1.00 [0.99, 1.00]	-	
Papageorghiou 2016b	2	Ō	Ō	9	1.00 [0.16, 1.00]	1.00 [0.66, 1.00]		
Poon 2016	41	Ō	Ō	200	1.00 [0.91, 1.00]	1.00 [0.98, 1.00]		
Zhang 2016	4	Ō	Ō	83	1.00 [0.40, 1.00]	1.00 [0.96, 1.00]		
Alberti 2015	47	Ō	Ō	136	1.00 [0.92, 1.00]	1.00 [0.97, 1.00]		•
Benachi 2015	110	1	3	744	0.97 [0.92, 0.99]	1.00 [0.99, 1.00]	-	
Ke 2015	24	Ó	0	2316	1.00 [0.86, 1.00]	1.00 [1.00, 1.00]		
Lee 2015	- 8	0	Ō	84	1.00 [0.63, 1.00]	1.00 [0.96, 1.00]		-
Song 2015	4	0	0	200	1.00 [0.40, 1.00]	1.00 [0.98, 1.00]		
Sukhikh 2015	26	1	2	171	0.93 [0.76, 0.99]	0.99 [0.97, 1.00]		
Sung-Hee 2015	6	0	0	895	1.00 [0.54, 1.00]	1.00 [1.00, 1.00]		
Wang 2015a	28	1	0	888	1.00 [0.88, 1.00]	1.00 [0.99, 1.00]		
Bijok 2014	1	0	0	8	1.00 [0.03, 1.00]	1.00 [0.63, 1.00]		
Huang 2014	10	0	1	178	0.91 [0.59, 1.00]	1.00 [0.98, 1.00]	_	
Jeon 2014	16	0	0	139	1.00 [0.79, 1.00]	1.00 [0.97, 1.00]		•
Porreco 2014	187	3	5	3127	0.97 [0.94, 0.99]	1.00 [1.00, 1.00]	•	
Stumm 2014	53	1	2	416	0.96 [0.87, 1.00]	1.00 [0.99, 1.00]		
Wang 2014	4	0	0	132	1.00 [0.40, 1.00]	1.00 [0.97, 1.00]		•
Bianchi 2013	43	1	1	68	0.98 [0.88, 1.00]	0.99 [0.92, 1.00]		-
Liang 2013	58	1	0	353	1.00 [0.94, 1.00]	1.00 [0.98, 1.00]	-	
Bianchi 2012	144	9	4	346	0.97 [0.93, 0.99]	0.97 [0.95, 0.99]	•	
Canick 2012	8	0	0	19	1.00 [0.63, 1.00]	1.00 [0.82, 1.00]		
Hou 2012	5	0	0	200	1.00 [0.48, 1.00]	1.00 [0.98, 1.00]		
Jiang 2012	30	1	0	872	1.00 [0.88, 1.00]	1.00 [0.99, 1.00]		
Liu 2012	3	1	0	149	1.00 [0.29, 1.00]	0.99 [0.96, 1.00]		•
Palomaki 2012	280	22	3	1716	0.99 [0.97, 1.00]	0.99 [0.98, 0.99]		•
Chen 2011	59	8	3	219	0.95 [0.87, 0.99]	0.96 [0.93, 0.98]		
Ehrich 2011	39	1	0	409	1.00 [0.91, 1.00]	1.00 [0.99, 1.00]		•
Sehnert 2011	22	0	0	25	1.00 [0.85, 1.00]	1.00 [0.86, 1.00]		
THDC							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

TMPS, autosomes

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)
Persico 2016	53	0	1	165	0.98 [0.90, 1.00]	1.00 [0.98, 1.00]
Hall 2014	15	0	0	49	1.00 [0.78, 1.00]	1.00 [0.93, 1.00]
Nicolaides 2013	28	0	0	201	1.00 [0.88, 1.00]	1.00 [0.98, 1.00]
Verweij 2013	17	0	1	486	0.94 [0.73, 1.00]	1.00 [0.99, 1.00]
Ashoor 2012	99	0	1	297	0.99 [0.95, 1.00]	1.00 [0.99, 1.00]
Norton 2012	118	3	1	2958	0.99 [0.95, 1.00]	1.00 [1.00, 1.00]
Sparks 2012a	44	0	0	123	1.00 [0.92, 1.00]	1.00 [0.97, 1.00]



i. MPSS

Thirty-two MPSS studies evaluated 15,797 unaffected (non T21, T18 and T13) pregnancies and 1508 cases (Alberti 2015; Benachi 2015; Bianchi 2012; Bianchi 2013; Bijok 2014; Canick 2012; Chen 2011; Ehrich 2011; Hou 2012; Huang 2014; Jeon 2014; Jiang 2012; Johansen 2016; Ke 2015; Kim 2016; Lee 2015; Lefkowitz 2016; Liang 2013; Liu 2012; Palomaki 2012; Papageorghiou 2016a; Papageorghiou 2016b; Poon 2016; Porreco 2014; Sehnert 2011; Song 2015; Stumm 2014; Sukhikh 2015; Sung-Hee 2015;

Wang 2014; Wang 2015a; Zhang 2016). Of the 32 studies, 19 evaluated only singleton pregnancies, three evaluated only multifetal pregnancies, three evaluated singleton and multifetal pregnancies, and the remaining seven studies did not report type of pregnancy. Based on the 32 studies, the summary sensitivity (95% CI) was 98.8% (97.2% to 99.5%) and the summary specificity (95% CI) was 99.9% (99.7% to 100%). Results are presented separately for singleton and multifetal pregnancy studies in Table 9. The sensitivity tends to be lower in multifetal pregnancies but there are

no enough studies in this subgroup to compare MPSS performance according to pregnancy type.

ii. TMPS

Seven TMPS studies evaluated 378 cases and 4282 unaffected (non T21, T18 and T13) pregnancies in women with singleton pregnancy (Ashoor 2012; Hall 2014; Nicolaides 2013; Norton 2012; Persico 2016; Sparks 2012a; Verweij 2013). The summary sensitivity (95% CI) was 98.9 (97.2% to 99.6%) and the summary specificity (95% CI) was 99.9% (99.8% to 100%) (Table 9).

iii. Comparative accuracy of MPSS and TMPS

An indirect comparison of the 32 MPSS and seven TMPS studies showed no statistical evidence of a difference in sensitivity, specificity or both (P value = 0.11). The differences in sensitivity and specificity were negligible (Summary of findings 6).

9. All sex chromosome aneuploidies (SCA) combined

The sex chromosome aneuploidies (45,X, 47,XXX, 47,XXY and 47,XYY) were considered together as one target condition. SCA was

assessed in 20 studies, comprising 286 affected cases and 9839 non SCA pregnancies. MPSS and TMPS were assessed by 14 and six studies, respectively. Among the 20 studies, 16 enrolled pregnant women selected at high risk of fetal aneuploidy and four enrolled a cohort of pregnant women with mixed prior risk. The results are summarised in Summary of findings 7. The results for mixed risk cohorts are summarised in Appendix 7.

a. Unselected population of pregnant women undergoing aneuploidy screening

No study assessed SCA in an unselected population of pregnant women.

b. Selected population of pregnant women at high risk of fetal aneuploidv

Sixteen studies involving 247 (2.93%) affected and 8420 non SCA pregnancies were included. MPSS and TMPS were assessed by 12 and four studies respectively (Figure 15).

Figure 15. Forest plot of MPSS and TMPS for SCA (45,X, 47,XXX, 47,XXY and 47,XYY combined) in pregnant women selected at high risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS, SCA

TMPS, SCA

Hooks 2014

Nicolaides 2014a

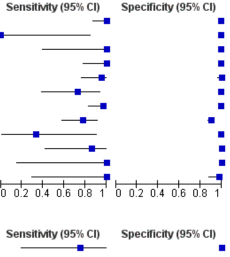
Nicolaides 2013

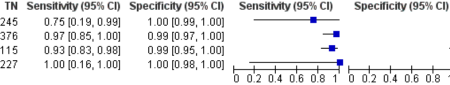
Study Persico 2016

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)
Lefkowitz 2016	26	- 7	0	1111	1.00 [0.87, 1.00]	0.99 [0.99, 1.00]
Song 2015	0	1	2	201	0.00 [0.00, 0.84]	1.00 [0.97, 1.00]
Sukhikh 2015	4	1	0	195	1.00 [0.40, 1.00]	0.99 [0.97, 1.00]
Porreco 2014	15	16	0	3291	1.00 [0.78, 1.00]	1.00 [0.99, 1.00]
Bianchi 2013	20	0	1	92	0.95 [0.76, 1.00]	1.00 [0.96, 1.00]
Liang 2013	8	1	3	400	0.73 [0.39, 0.94]	1.00 [0.99, 1.00]
Mazloom 2013	29	1	1	380	0.97 [0.83, 1.00]	1.00 [0.99, 1.00]
Bianchi 2012	21	46	6	416	0.78 [0.58, 0.91]	0.90 [0.87, 0.93]
Hou 2012	1	1	2	201	0.33 [0.01, 0.91]	1.00 [0.97, 1.00]
Jiang 2012	6	1	1	895	0.86 [0.42, 1.00]	1.00 [0.99, 1.00]
Liu 2012	2	0	0	151	1.00 [0.16, 1.00]	1.00 [0.98, 1.00]
Sehnert 2011	3	1	0	43	1.00 [0.29, 1.00]	0.98 [0.88, 1.00]

115

0 227





i. MPSS

Twelve MPSS studies evaluated 151 affected and 7452 non SCA pregnancies (Bianchi 2012; Bianchi 2013; Hou 2012; Jiang 2012; Lefkowitz 2016; Liang 2013; Liu 2012; Mazloom 2013; Porreco 2014; Sehnert 2011; Song 2015; Sukhikh 2015). Of the 12 studies, seven included only women with singleton pregnancy, one evaluated singleton and multifetal pregnancies, and the remaining four studies did not report type of pregnancy. Results are presented

TP FP FN

3 0 1 245

33 4 1 376

52 1 4

> 2 0

> > separately for singleton and multifetal pregnancy studies in Table 9. Based on all 12 studies, the summary sensitivity (95% CI) was 91.9% (73.8% to 97.9%) and the summary specificity (95% CI) was 99.5% (98.8% to 99.8%).

ii. TMPS

Four TMPS studies evaluated 96 affected and 968 non SCA pregnancies in women with singleton pregnancy (Hooks 2014;

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0.75 [0.19, 0.99]

0.97 [0.85, 1.00]

0.93 [0.83, 0.98]

1.00 [0.16, 1.00]

Nicolaides 2013; Nicolaides 2014a; Persico 2016). The summary sensitivity (95% CI) was 93.8% (86.8% to 97.2%) and the summary specificity (95% CI) was 99.6% (98.1% to 99.9%).

iii. Comparative accuracy of MPSS and TMPS

An indirect comparison of the 12 MPSS and four TMPS studies showed no statistical evidence of a difference in sensitivity, specificity or both (P value = 0.41). The differences in sensitivity and specificity were negligible (Summary of findings 7).

10. gNIPT approach (MPSS or TMPS) against traditional screening tests

Five studies directly compared a gNIPT approach (MPSS or TMPS) and traditional screening tests for autosomal aneuploidies by using cohorts of pregnant women who were tested by both methods. Three studies compared TMPS and traditional screening tests, and two studies compared MPSS and traditional screening tests. The results are summarised in Summary of findings 2, Summary of findings 3, Summary of findings 4 and Summary of findings 6.

a. Unselected population of pregnant women undergoing aneuploidy screening

Only one study that compared TMPS and a traditional screening test evaluated T21, T18 and T13 individually in an unselected population of pregnant women undergoing aneuploidy screening (Norton 2015). This study evaluated 38, 10 and two cases of T21, T18 and T13, respectively and 15,803, 15,831 and 11,183 non T21, T18 and T13, respectively (Figure 16). Direct comparisons between gNIPT and traditional screening tests were not possible because there was only one study but authors observed eight, two and one cases of T21, T18 and T13 respectively missed by traditional screening test and only one T18 case missed by TMPS.

Figure 16. Forest plot of traditional screening tests for T21, T18 and T13 in unselected pregnant women undergoing aneuploidy screening. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

Traditional screening tests T21

Study	ТР	FP	FN	I TN	Sensitivity (95% Cl)) Specificity (95% Cl)	Sensitivity (95% CI)	Specificity (95% CI)
Norton 2015	30	854	4 8	3 14949	0.79 [0.63, 0.90]] 0.95 [0.94, 0.95]		
Traditional sc	reeni	ng te	sts	T18			0 0.2 0.4 0.0 0.8 1	0 0.2 0.4 0.0 0.8 1
Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Norton 2015	8	49	2	15782	0.80 [0.44, 0.97]	1.00 [1.00, 1.00]		
Traditional sc	reeni	ng te	sts	T13			0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Norton 2015	1	28	1	11155	0.50 [0.01, 0.99]	1.00 [1.00, 1.00]		
							0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1

Four studies compared a gNIPT approach with a traditional screening test for autosomal aneuploidies (T21, T18 and T13 combined) in 22,367 unselected pregnant women (Figure 17). Three studies (Nicolaides 2012; Norton 2015; Quezada 2015) compared TMPS and first-trimester combined test (Figure 18), and one study (Song 2013) compared MPSS and a second-trimester triple test. The three TMPS studies had similar characteristics. Meta-analyses of direct comparisons between gNIPT and traditional screening tests

were not possible because traditional screening tests used different cutpoints and there were very few studies to enable estimation of summary sensitivity and specificity at specific cutpoints. Individual study results are presented in Table 10. Overall, 16 aneuploid cases were missed by traditional screening test and only five cases were missed by gNIPT approach. While specificity was consistently higher for TMPS than traditional screening tests, sensitivity was not consistently higher as shown in Figure 18.

Figure 17. Forest plot of traditional screening tests for autosomes (T21, T18 and T13 combined) in unselected pregnant women undergoing aneuploidy screening. FN: false negative, FP: false positive, TN: true negative and TP: true positive.

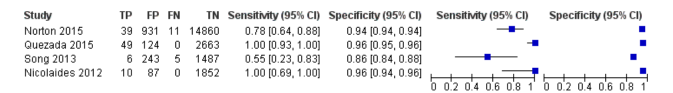


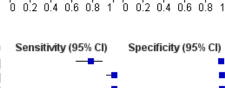


Figure 18. Forest plot of comparative studies of TMPS and traditional screening tests for autosomes (T21, T18 and T13 combined) in unselected pregnant women undergoing aneuploidy screening. FN: false negative, FP: false positive, TN: true negative and TP: true positive.

TMPS, autosomes

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% Cl)
Norton 2015	49	12	1	15779	0.98 [0.89, 1.00]	1.00 [1.00, 1.00]	-	I I
Quezada 2015	43	8	4	2730	0.91 [0.80, 0.98]	1.00 [0.99, 1.00]		•
Nicolaides 2012	10	2	0	1937	1.00 [0.69, 1.00]	1.00 [1.00, 1.00]		
Traditional screer	ning t	ests	, aut	osomes				

Study TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Norton 2015 39 931 11 14860 0.78 [0.64, 0.88] 0.94 [0.94, 0.94] 1.00 [0.93, 1.00] 0.96 [0.95, 0.96] Quezada 2015 49 124 0 2663 Nicolaides 2012 10 87 0 1852 1.00 [0.69, 1.00] 0.96 [0.94, 0.96]



b. Mixed-risk cohort of fetal aneuploidy

One study compared MPSS and traditional screening test for autosomal aneuploidies (T21, T18 and T13 combined) in a cohort with mixed prior risk of fetal aneuploidy including 1908 non T21, T18 and T31 pregnancies and four cases of autosomal aneuploidy (Bianchi 2014a). Traditional screening tests included first-trimester combined test or a second-trimester result (quadruple, serum integrated, fully integrated or sequential) (Figure 19). Overall, 80 unaffected pregnancies were detected as affected by traditional screening test against 12 for TMPS.

Figure 19. Forest plot of traditional screening tests for autosomes (T21, T18 and T13 combined) in pregnant women with mixed prior risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

Traditional screening tests, autosomes

Study	ΤР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI) Specificity (95% CI)
Bianchi 2014a	4	80	0	1828	1.00 [0.40, 1.00]	0.96 [0.95, 0.97]	
Traditional scre	ening	j tes	ts T2	21			'o o.'2 o.'4 o.'6 o.'8 1' 'o o.'2 o.'4 o.'6 o.'8 1'
Study	тр	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95% CI) Specificity (95% CI)
Bianchi 2014a	3	69	0	1840	1.00 [0.29, 1.00]	0.96 [0.95, 0.97]	
Traditional scre	ening) tes	ts T1	8			0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
Study	тр	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% Cl)	Sensitivity (95% Cl) Specificity (95% Cl)
Bianchi 2014a	1	11	0	1894	1.00 [0.03, 1.00]	0.99 [0.99, 1.00]	

11. Pregnant women with mixed prior risk of fetal aneuploidy

Summary sensitivities and specificities for cohorts of pregnant women with mixed prior risk of fetal aneuploidy are presented in Appendix 7. For autosomal aneuploidies, 17 studies included 838 cases and 39,615 unaffected (non T21, T18 and T13) pregnancies. Of the 17 studies, 10 assessed MPSS and seven assessed TMPS (Figure 20). For T21, 16 studies included 614 cases (1.6%) and 37,887 non T21 pregnancies. Of the 16 studies, 10 assessed MPSS and six assessed TMPS. For T18, 13 studies included 166 cases (0.5%) and 36,206 non T18 pregnancies. Of the 13 studies, nine assessed MPSS and four assessed TMPS. For T13, 13 studies included 58 cases (0.1%) and 38,746 non T13 pregnancies. Eight of the 13 studies assessed MPSS and the other five assessed TMPS (Figure 21). For SCA, four studies included 39 cases and 1419 non SCA pregnancies; two of the studies assessed MPSS and the other two assessed TMPS (Figure 22). For 45,X, four studies included 34 cases (2.4%) and 1424 non 45,X pregnancies. Of the four studies, two studies assessed



MPSS and two studies assessed TMPS. For 47,XXY, two studies (one of MPSS and one of TMPS) included three cases (1%) and 291 non 47,XXY pregnancies. For 47,XYY, two studies included two cases (0.5%) and 384 non 47,XYY pregnancies; one study assessed MPSS

and the other study assessed TMPS. No study assessed gNIPT for 47,XXX in cohorts of pregnant women with mixed prior risk of fetal aneuploidy (Figure 23).

Figure 20. Forest plot of MPSS and TMPS for autosomes (T21, T18 and T13 combined) in pregnant women with mixed prior risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS, autosomes

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Fiorentino 2016	105	2	0	6975	1.00 [0.97, 1.00]	1.00 [1.00, 1.00]	-	•
Ma 2016	211	4	0	10364	1.00 [0.98, 1.00]	1.00 [1.00, 1.00]	•	•
Tynan 2016	37	0	0	1011	1.00 [0.91, 1.00]	1.00 [1.00, 1.00]		•
Bianchi 2014a	8	12	0	1932	1.00 [0.63, 1.00]	0.99 [0.99, 1.00]		•
Shaw 2014	22	0	0	178	1.00 [0.85, 1.00]	1.00 [0.98, 1.00]		•
Yao 2014	38	2	0	5490	1.00 [0.91, 1.00]	1.00 [1.00, 1.00]		
Zhou 2014a	5	0	0	296	1.00 [0.48, 1.00]	1.00 [0.99, 1.00]		
Zhou 2014b	50	6	0	3894	1.00 [0.93, 1.00]	1.00 [1.00, 1.00]		
Lau 2012	23	0	0	85	1.00 [0.85, 1.00]	1.00 [0.96, 1.00]		
Chiu 2011	68	6	18	565	0.79 [0.69, 0.87]	0.99 [0.98, 1.00]	·····	
	_						0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.6 0.8 1
TMPS autosomes	2							

TMPS, autosomes

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)
Gil 2016	66	9	3	3555	0.96 [0.88, 0.99]	1.00 [1.00, 1.00]
Bevilacqua 2015	16	0	1	323	0.94 [0.71, 1.00]	1.00 [0.99, 1.00]
Comas 2015	4	0	0	308	1.00 [0.40, 1.00]	1.00 [0.99, 1.00]
Jackson 2014	3	- 4	2	1152	0.60 [0.15, 0.95]	1.00 [0.99, 1.00]
Korostelev 2014	52	0	1	632	0.98 [0.90, 1.00]	1.00 [0.99, 1.00]
Pergament 2014	94	1	1	870	0.99 [0.94, 1.00]	1.00 [0.99, 1.00]
Ashoor 2013	8	1	2	1938	0.80 [0.44, 0.97]	1.00 [1.00, 1.00]

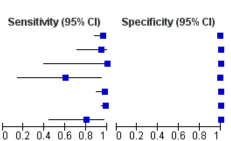


Figure 21. Forest plot of MPSS and TMPS for T21, T18 or T13 in pregnant women with mixed prior risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS T21							
Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI) Specificity (95% CI)
Fiorentino 2016	76	1	0	7005	1.00 [0.95, 1.00]	1.00 [1.00, 1.00]	
Ma 2016	162	2	0	10415	• • •		
Tynan 2016	21	0	0	1027			
Bianchi 2014a	5	6	0	1941	1.00 [0.48, 1.00]		
Shaw 2014	11	0	0	189	1.00 [0.72, 1.00]	1.00 [0.98, 1.00]	
Yao 2014	31	0	0	5499			
Zhou 2014a	4	0	0	297			
Zhou 2014b	38	2	0	3910			
Lau 2012	11	0	0	97			
Chiu 2011	68	6	18	565	0.79 [0.69, 0.87]	0.99 [0.98, 1.00]	
MPSS T18							0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI) Specificity (95% CI)
Fiorentino 2016	20	1	0	7061	1.00 [0.83, 1.00]	1.00 [1.00, 1.00]	
Ma 2016	46	2	0	10531	1.00 [0.92, 1.00]	1.00 [1.00, 1.00]	
Tynan 2016	10	0	0	1038	1.00 [0.69, 1.00]	1.00 [1.00, 1.00]	
Bianchi 2014a	2	3	0	1947	1.00 [0.16, 1.00]	1.00 [1.00, 1.00]	
Shaw 2014	8	0	0	192	1.00 [0.63, 1.00]	1.00 [0.98, 1.00]	
Yao 2014	6	1	0	5523	1.00 [0.54, 1.00]	1.00 [1.00, 1.00]	
Zhou 2014a	1	0	0	300	1.00 [0.03, 1.00]	1.00 [0.99, 1.00]	
Zhou 2014b	10	2 0	0	3938	1.00 [0.69, 1.00]	1.00 [1.00, 1.00]	
Lau 2012	10	U	0	98	1.00 [0.69, 1.00]	1.00 [0.96, 1.00]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1
MPSS T13							
Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI) Specificity (95% CI)
Study	IF				Sensitivity (55% CI)	specificity (55% ci)	Sensitivity (95% CI) Specificity (95% CI)
Fiorentino 2016	9	0	0	7073	1.00 [0.66, 1.00]	1.00 [1.00, 1.00]	
2			0				
Fiorentino 2016 Ma 2016 Tynan 2016	9	0 0 0	0 0 0	7073 10576 1042	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a	9 3 6 1	0 0 0 3	0 0 0 0	7073 10576 1042 1910	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014	9 3 6 1 3	0 0 0 3 0	0 0 0 0	7073 10576 1042 1910 197	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.03, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014	9 3 6 1 3	0 0 3 0	0 0 0 0 0	7073 10576 1042 1910 197 5528	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [0.98, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b	9 3 6 1 3 1 2	0 0 3 0 1 2	0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.03, 1.00] 1.00 [0.16, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012	9 3 6 1 3	0 0 3 0	0 0 0 0 0	7073 10576 1042 1910 197 5528	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [0.98, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b	9 3 6 1 3 1 2	0 0 3 0 1 2	0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.03, 1.00] 1.00 [0.16, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012	9 3 6 1 3 1 2 2	0 0 3 0 1 2	0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.03, 1.00] 1.00 [0.16, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21	9 3 6 1 3 1 2 2	0 0 3 0 1 2 0	0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106	$\begin{array}{c} 1.00 & [0.66, 1.00] \\ 1.00 & [0.29, 1.00] \\ 1.00 & [0.54, 1.00] \\ 1.00 & [0.03, 1.00] \\ 1.00 & [0.29, 1.00] \\ 1.00 & [0.03, 1.00] \\ 1.00 & [0.03, 1.00] \\ 1.00 & [0.16, 1.00] \\ 1.00 & [0.16, 1.00] \end{array}$	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study	9 3 6 1 3 1 2 2 TP	0 0 3 0 1 2 0	0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016	9 3 6 1 3 1 2 2 TP 43	0 0 3 0 1 2 0 FP	0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] Sensitivity (95% CI) 0.98 [0.88, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] Specificity (95% Cl) 1.00 [1.00, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015	9 3 6 1 3 1 2 2 TP 43 11	0 0 3 0 1 2 0 FP 1 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] Sensitivity (95% CI) 0.98 [0.88, 1.00] 0.92 [0.62, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] Specificity (95% Cl) 1.00 [1.00, 1.00] 1.00 [0.99, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014	9 3 6 1 3 1 2 2 TP 43 11 4 3 47	0 0 3 0 1 2 0 FP 1 0 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] Sensitivity (95% CI) 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] Specificity (95% Cl) 1.00 [1.00, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014	9 3 6 1 3 1 2 2 TP 43 11 4 3	0 0 3 0 1 2 0 FP 1 0 0 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] Specificity (95% Cl) 1.00 [1.00, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00]	Sensitivity (95% Cl)
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014	9 3 6 1 3 1 2 2 TP 43 11 4 3 47	0 0 3 0 1 2 0 FP 1 0 0 0 0 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99] 1.00 [0.92, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] Specificity (95% Cl) 1.00 [1.00, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00]	
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014 Pergament 2014	9 3 6 1 3 1 2 2 TP 43 11 4 3 47	0 0 3 0 1 2 0 FP 1 0 0 0 0 0 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638 905	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.54, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99] 1.00 [0.94, 1.00] 1.00 [0.94, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] Specificity (95% Cl) 1.00 [1.00, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00]	Sensitivity (95% Cl)
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014 Pergament 2014	9 3 6 1 3 1 2 2 7 7 7 4 3 11 4 3 47 58 7 7	0 0 3 0 1 2 0 FP 1 0 0 0 0 0 0 0 FP	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638 905 TN	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99] 1.00 [0.94, 1.00] 1.00 [0.94, 1.00] 1.00 [0.94, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] 1.00 [0.97, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00]	Sensitivity (95% Cl)
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014 Pergament 2014 TMPS T18 Study Gil 2016	9 3 6 1 3 1 2 2 4 3 11 4 3 47 58	0 0 3 0 1 2 0 FP 1 0 0 0 0 0 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638 905	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99] 1.00 [0.94, 1.00] 1.00 [0.94, 1.00] Sensitivity (95% CI) 1.00 [0.84, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] 1.00 [0.97, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00]	Sensitivity (95% Cl)
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014 Pergament 2014 TMPS T18 Study	9 3 6 1 3 1 2 2 7 7 7 7 7 8 7 7 8 7 7 7 7 8 7 7 7 7	0 0 3 0 1 2 0 FP 1 0 0 0 0 0 0 0 0 FP 4	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638 905 TN 3608	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99] 1.00 [0.94, 1.00] 1.00 [0.94, 1.00] 1.00 [0.94, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] 1.00 [0.97, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00]	Sensitivity (95% Cl)
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014 Pergament 2014 TMPS T18 Study Gil 2016 Bevilacqua 2015	9 3 6 1 3 1 2 2 7 7 7 7 7 8 7 7 8 7 7 7 8 7 7 7 8 7 7 7 7 8 7 7 7 7 7 8 7	0 0 3 0 1 2 0 FP 1 0 0 0 0 0 0 0 0 FP 4 0 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638 905 TN 3608 335	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.3, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99] 1.00 [0.92, 1.00] 1.00 [0.94, 1.00] 1.00 [0.84, 1.00] 1.00 [0.84, 1.00] 1.00 [0.48, 1.00] 1.00 [0.48, 1.00]	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] 1.00 [0.97, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [0.99, 1	Image: Second control of the second
Fiorentino 2016 Ma 2016 Tynan 2016 Bianchi 2014a Shaw 2014 Yao 2014 Zhou 2014b Lau 2012 TMPS T21 Study Gil 2016 Bevilacqua 2015 Comas 2015 Jackson 2014 Korostelev 2014 Pergament 2014 TMPS T18 Study Gil 2016 Bevilacqua 2015 Korostelev 2014	9 3 6 1 3 1 2 2 7 7 7 4 3 11 4 3 47 58 7 7 8 7 7 2 1 5 2	0 0 3 0 1 2 0 FP 1 0 0 0 0 0 0 0 0 FP 4 0 0	0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	7073 10576 1042 1910 197 5528 3946 106 TN 3588 328 308 1157 638 905 TN 3608 335 683	1.00 [0.66, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.03, 1.00] 1.00 [0.29, 1.00] 1.00 [0.29, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 1.00 [0.16, 1.00] 0.98 [0.88, 1.00] 0.92 [0.62, 1.00] 1.00 [0.40, 1.00] 0.75 [0.19, 0.99] 1.00 [0.92, 1.00] 1.00 [0.94, 1.00] 1.00 [0.84, 1.00] 1.00 [0.84, 1.00] 1.00 [0.48, 1	1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.98, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.97, 1.00] 1.00 [0.97, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [1.00, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00] 1.00 [0.99, 1.00]	Sensitivity (95% Cl)



Figure 21. (Continued)

TMPS T13							U U.Z U.4 U.0 U.8 I	U U.Z U.4 U.0 U.0 I
Study	ТР	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% CI)
Gil 2016	2	4	2	3625	0.50 [0.07, 0.93]	1.00 [1.00, 1.00]		•
Jackson 2014	0	1	1	1159	0.00 [0.00, 0.97]	1.00 [1.00, 1.00]		•
Korostelev 2014	3	0	1	681	0.75 [0.19, 0.99]	1.00 [0.99, 1.00]		•
Pergament 2014	12	0	0	953	1.00 [0.74, 1.00]	1.00 [1.00, 1.00]		•
Ashoor 2013	8	1	2	1938	0.80 [0.44, 0.97]	1.00 [1.00, 1.00]	0 0.2 0.4 0.6 0.8 1	

Figure 22. Forest plot of MPSS and TMPS for SCA (45,X, 47,XXX, 47,XXY and 47,XYY combined) in pregnant women with mixed prior risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

MPSS, SCA														
Study	TP	FP	FN	TN	Se	ensiti	vity (9	95% CI)	Specific	ity (9	5% CI)		Sensitivity (95% CI)	Specificity (95% CI)
Shaw 2014	4	0	1	195		0.80	0.28	3, 0.99]	1.00	0.98	3, 1.00]			
Lau 2012	9	0	0	99		1.00	0.60	6, 1.00]	1.00	[0.96	6, 1.00]			
TMPS, SCA													0 0.2 0.4 0.0 0.0 1	0 0.2 0.4 0.0 0.0 1
Study				ΤР	FP	FN	ΤN	Sensit	ivity (95%	6 CI)	Specificity (95% C	CI)	Sensitivity (95% CI)	Specificity (95% CI)
Pergament 2014			9	1	1	953	0.9	0 [0.55, 1	.00]	1.00 (0.99, 1.0	[00			
Samango-Sp	rous	e 20	13	14	0	1	171	0.9	13 [0.68, 1	.00]	1.00 [0.98, 1.0	00]		



Figure 23. Forest plot of MPSS and TMPS for 45,X, 47,XXY or 47,XYY in pregnant women with mixed prior risk of fetal aneuploidy. FN: false negative, FP: false positive, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, TN: true negative and TP: true positive.

TN Sensit	ivity (95% CI) Specificity (95% Cl) Sensitivity (95% CI)	Specificity (95% Cl)
196 0.7	5 [0.19, 0.99] 1.00 [0.98, 1.00	ıj — — — — — — — — — — — — — — — — — — —	
100 1.0	0 [0.63, 1.00] 1.00 [0.96, 1.00		
		0 0.2 0.4 0.0 0.8 1	0 0.2 0.4 0.8 0.8 1
TN Sensitiv	ity (95% CI) Specificity (95% CI)	Sensitivity (95% CI)	Specificity (95% Cl)
107 1.00	[0.03, 1.00] 1.00 [0.97, 1.00]		
		0 0.2 0.4 0.6 0.8 1	0 0.2 0.4 0.8 0.8 1
TN Sensit	ivity (95% CI) Specificity (95% Cl) Sensitivity (95% CI)	Specificity (95% CI)
199 1.0	0 [0.03, 1.00] 1.00 [0.98, 1.00		
		0 0.2 0.4 0.0 0.8 1	0 0.2 0.4 0.0 0.6 1
TP FP FN	TN Sensitivity (95% CI) Spec	ificity (95% CI) Sensitivity (95% CI)	Specificity (95% CI)
911	953 0.90 [0.55, 1.00] 1	.00 [0.99, 1.00]	•
11 0 1	174 0.92 [0.62, 1.00] 1		
		0 0.2 0.7 0.0 0.0 1	0 0.2 0.4 0.0 0.0 1
TP FP FN	TN Sensitivity (95% CI) Spec	ificity (95% CI) Sensitivity (95% CI)	Specificity (95% CI)
200	184 1.00 [0.16, 1.00] 1		
		0 0.2 0.4 0.0 0.8 1	0 0.2 0.4 0.0 0.8 1
TP FP FN	TN Sensitivity (95% CI) Spec	ificity (95% CI) Sensitivity (95% CI)	Specificity (95% Cl)
100	185 1.00 [0.03, 1.00] 1		
	196 0.7 100 1.0 TN Sensitiv 07 1.00 TN Sensitiv 199 1.00 TP FP 9 1 11 0 TP FP FN 2 2 0 TP FP FN 2 FP FN	196 0.75 [0.19, 0.99] 1.00 [0.98, 1.00] 100 1.00 [0.63, 1.00] 1.00 [0.96, 1.00] TN Sensitivity (95% CI) Specificity (95% CI) 07 1.00 [0.03, 1.00] 1.00 [0.97, 1.00] 07 1.00 [0.03, 1.00] 1.00 [0.98, 1.00] TN Sensitivity (95% CI) Specificity (95% CI) 199 1.00 [0.03, 1.00] 1.00 [0.98, 1.00] TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) Specificity (95% CI) 199 1.00 [0.03, 1.00] 1.00 [0.98, 1.00] 1 TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) 9 1 953 0.90 [0.55, 1.00] 1 11 0 1 174 0.92 [0.62, 1.00] 1 TP FP FN TN Sensitivity (95% CI) Specificity (95% CI) 2 0 184 1.00 [0.16, 1.00] 1 TP FP FN TN Sensitivity (95% CI) Specificity (95% CI)	196 $0.75 [0.19, 0.99]$ $1.00 [0.98, 1.00]$ 100 $1.00 [0.63, 1.00]$ $1.00 [0.96, 1.00]$ TN Sensitivity (95% Cl) Sensitivity (95% Cl) 0 0.2 0.4 0.6 0.8 TN Sensitivity (95% Cl) Sensitivity (95% Cl) Sensitivity (95% Cl) 0 0.2 0.4 0.6 0.8 TN Sensitivity (95% Cl) Sensitivity (95% Cl) Sensitivity (95% Cl) 199 $1.00 [0.03, 1.00]$ $1.00 [0.98, 1.00]$ 0.2 0.4 0.6 0.8 TP FP FN TN Sensitivity (95% Cl) Sensitivity (95% Cl) Sensitivity (95% Cl) 9 1 953 $0.90 [0.55, 1.00]$ $1.00 [0.98, 1.00]$ 0.2 0.4 0.6 0.8 11 0 1.74 $0.92 [0.62, 1.00]$ $1.00 [0.98, 1.00]$ 0.2 0.4 0.6 0.8 TP FP FN TN Sensitivity (95% Cl) 0.2 0.4 0.6 0.2 0.4 0.6 0.8 0.2 0.4

12. Failure rates

Table 7 shows the non-negligible failure rate of gNIPT reported in the studies. gNIPT assay failure rate was reported in 46 out of 65 (71%) studies. The largest failure rate (25%) was observed in a study that used its own developed MPSS assay (Alberti 2015). The main reasons for assay failure included low amount of ccfDNA, low fetal fraction DNA and failure of sample to pass quality control. The failure rate ranged between 0% and 25% for MPSS and between 0.8% and 7.5% for TMPS. The number of aneuploid and euploid cases in failed samples was reported in 23 of 46 (50%) studies. Among these 23 studies, there were 1064 euploid cases and 79 aneuploid cases, ranged between 0% and 50% for MPSS and between 0% and 23% for TMPS. The failure rate among aneuploid cases, ranged between 0% and 50% for MPSS and between 0% and 23% for TMPS. The failure rate among euploid cases ranged between 0% and 6.7% for MPSS and between 1% and 7.6% for TMPS.

Investigation of heterogeneity

We planned to evaluate the effect of potential sources of heterogeneity such as type of reference standard and ethnicity. However, formal investigations using meta-regression were not possible due to limited data and little or no heterogeneity in the sensitivities and specificities. Most studies (55%) used karyotyping while the remaining 29 studies (45%) used multiple reference standards. Ethnicity was not reported by 26 (40%) studies while the population in 21 (32%) studies was more than 50% Asian and in 18 (28%) studies the population was more than 50% Caucasian. In Appendix 8, the number of studies, affected and unaffected pregnancies are shown according to the gNIPT approach and prior risk of fetal aneuploidy. We also planned to assess gNIPT performance according to gestational age and gNIPT approach for autosomes and SCA aneuploidies. The accuracy of gNIPT appears to be high in all gestational age groups.

Sensitivity analyses

We did not perform sensitivity analyses to assess the effect of the interval between blood collection for gNIPT and fluid collection for reference standard because most studies had an acceptable interval between sample collection for index test and reference standard. Due to lack of data or lack of variability in estimates of sensitivity and specificity, analyses of the effect of high or unclear risk of bias according to the QUADAS-2 domains were not done. We performed sensitivity analyses using data from all autosomes combined and all SCA combined in order to have enough studies to assess the impact of study design and number of cases. The results are presented in Table 11. Excluding case-control studies or studies



with less than 10 an euploid cases had little or no impact on our findings.

DISCUSSION

Summary of main results

This review included data from 65 studies of 86,139 pregnant women (including 3141 aneuploids) tested by genomics-based non-invasive prenatal testing (gNIPT) and a reference standard. The gNIPT method used circulating cell-free DNA (ccfDNA) in maternal blood for the detection of common fetal aneuploidies (T21, T18, T13, 45,X, 47,XXY, 47,XXX and 47,XYY). The number of gNIPT studies in unselected populations was limited (five studies), but 42 studies in high-risk cohorts provided data for various metaanalyses. Few (14%) studies included more than 100 aneuploid cases. Importantly, in almost all studies, the risk of bias was generally high with respect to patient selection as well as flow and timing. Some women can spontaneously lose their pregnancy after enrolment into a study. However, none of the studies reported such events. Since women with spontaneous abortions are likely to be lost to follow-up, we believe that any risk of bias has been captured in the quality assessment of studies. Blood samples for gNIPT were mainly taken just before the invasive test (reference standard) and so pregnancies were unlikely to terminate naturally between the gNIPT and the reference standard. Across all studies, applicability concerns were low in the index test and reference standard domains.

These results show that massively parallel shotgun sequencing (MPSS) and targeted massively parallel sequencing (TMPS) perform similarly in terms of clinical sensitivity and specificity for the detection of fetal T21, T18, T13 and sex chromosome aneuploidy (SCA). However, no study compared the two approaches head-to-head in the same cohort of patients.

In high-risk pregnancies, gNIPT methods (MPSS and TMPS) were highly accurate for detection of any of the three major trisomies (T21, T18 and T13) with sensitivities from 95.8% to 99.7% depending on specific trisomies and specificities above 99%. There were no statistically significant differences in accuracy between MPSS and TMPS.

In unselected cohorts of pregnant women, only one study evaluated MPSS. Based on meta-analytic findings for each trisomy, TMPS appeared to be accurate for the detection of T21, with lower accuracy for T18 and T13. When compared to traditional prenatal screening tests, only four studies were identified (three for TMPS and one for MPSS). Genomics-based non-invasive prenatal testing showed greater specificity for T21 and T18 than traditional screening tests, while inconsistent results were observed for sensitivity. The inconsistency may be due to different cutpoints for traditional screening tests though one would expect that to also affect specificity. Given the small number of studies, the differences may be due to chance or there may be other differences between the studies that were not apparent.

With respect to the replacement of invasive tests, the performance of gNIPT observed in this review is not sufficient to replace current invasive diagnostic tests.

We also compared the diagnostic test accuracy of MPSS and TMPS for all three autosomes combined because gNIPT is being clinically

proposed as one test during prenatal follow-up to detect any of the three conditions. Under this scenario, in high-risk pregnancies of fetal aneuploidy, there was no statistically significant difference in diagnostic accuracy between MPSS and TMPS. In unselected cohorts of pregnant women, a test comparison was not possible due to limited data.

There was paucity of data for each SCA. In high-risk cohorts, all SCAs combined gave a pooled sensitivity (95% CI) and specificity (95% CI) of 91.9% (73.8% to 97.9%) and 99.5% (98.8% to 99.8%) from 12 MPSS studies. The pooled sensitivity (95% CI) and specificity (95% CI) were 93.8% (86.8% to 97.2%) and 99.6% (98.1% to 99.9%) from four TMPS studies. SCAs are considered "incidental" findings of current aneuploidy screening programs. It should be noted that SCAs are not of interest for prenatal screening since they do not lead to any intervention prior to birth.

The failure rate associated with gNIPT, which is higher than the current failure rate of traditional screening tests which is close to zero, is worrying and may be a source of bias. Futhermore, the large heterogeneity between laboratory-developed assays in their protocol details and observed failure rates highlight the fact that each laboratory providing gNIPT services should determine its own failure rate and inform healthcare professionals ordering the test about this important test characteristic. Failed samples were excluded from the analyses in the studies. This systematic review found a slightly larger failure rate for TMPS than the MPSS approach. This was also reported by Yaron 2016. We also found that the proportion of failed samples for aneuploid samples was higher than the proportion of failed samples for euploid samples. If these failed samples were included in the summary statistics, the diagnostic performance of gNIPT would be lower.

Comparison with other systematic reviews with meta analysis

At the time of writing, there are four published systematic reviews with meta-analyses of gNIPT (Gil 2015a; HAS 2015; Mackie 2017; Taylor-Phillips 2016). Although these meta-analyses had different criteria for including studies and analyses, they reported similar sensitivities and specificities to our findings.

As reported by Gil 2015a, the detection rate of gNIPT for autosomes was between 91.0% to 99.2% and specificity above 99.9% in singleton pregnancies. The detection rate for 45,X and SCA other than 45,X was 90.3% and 93.0%, respectively with specificity above 99.8% in singleton pregnancies. The results from HAS 2015 group for T21 were respectively 98.0% and 99.9% for sensitivity and specificity. Regarding Mackie 2017, the sensitivity was between 90.6% to 99.4% and specificity above 99.9% for autosomes. For 45,X, the sensitivity and specificity was 92.9% and 99.9%, respectively. They also pointed out that failed results were poorly reported across studies. Finally, Taylor-Phillips 2016 reported sensitivity between 97.4% to 99.3% for autosomes and specificity of 99.9%.

This is the first Cochrane diagnostic test accuracy (DTA) review on gNIPT. There are five published Cochrane DTA reviews on prenatal screening tests (Alldred 2012; Alldred 2015; Alldred 2015a; Alldred 2017a; Alldred 2017b). The suite of reviews addressed traditional biochemical, ultrasound and urine markers for Down syndrome screening (Alldred 2010) and none of the other fetal aneuploidies considered in this review were evaluated in this suite. In the first of the three reviews, Alldred and colleagues

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evaluated second-trimester serum markers and found that double and triple test combinations (involving alpha-fetoprotein, human chorionic gonadotropin (hCG) (free and total) or unconjugated estriol) significantly outperformed individual markers, detecting six to seven out of every 10 Down syndrome pregnancies at a 5% false positive rate (Alldred 2012). The second review evaluated firsttrimester serum markers and found that a test strategy involving maternal age, PAPP-A and free ßhCG significantly outperformed individual markers, detecting about seven out of every 10 Down's syndrome pregnancies at a 5% false positive rate (Alldred 2015a). The third review evaluated urine markers and concluded there was a paucity of evidence to support the use of urine testing for Down syndrome screening (Alldred 2015b). The fourth review evaluated first-trimester ultrasound tests alone or in combination with first-trimester serum tests and found that a combination of ultrasound and serum markers (especially PAPP-A and free ßhCG) and maternal age can detect about nine of 10 T21 affected pregnancies for a fixed 5% false positive rate (Alldred 2017a). The fifth review evaluated first- and second-trimester serum tests with and without first-trimester ultrasound tests and found that a combination of first-trimester ultrasound with first- and secondtrimester serum markers with maternal age are significantly better than those without ultrasound or those evaluating first-trimester ultrasound in combination with second-trimester serum markers, without first-trimester serum markers (the authors cannot make recommendations about a specific strategy) (Alldred 2017b).

Strengths and weaknesses of the review

Strengths

The review methodology was transparent with the full protocol published in the Cochrane Library (1 July 2015) and in PROSPERO (11 November 2015). The review evaluated the screening and diagnostic accuracy of gNIPT by MPSS and TMPS for seven common aneuploidies with no restriction imposed on population characteristics such as maternal age, gestational age, aneuploidy risk, number of fetuses and ethnicity. We performed a comprehensive search with no language restriction and we included studies in the languages used by various authors in the field, including Chinese, Bulgarian, Russian, Polish, Korean and Spanish. Study selection, data extraction and quality assessment were independently performed by two review authors. We contacted authors to clarify data and to avoid duplication of data as a result of overlapping populations.

We evaluated the performance of the two major gNIPT methods (MPSS and TMPS which included digital analysis of selected regions (DANSR) and single nucleotide polymorphism (SNP)-based method) and included data on traditional screening tests when compared to gNIPT.

We collected and reported data on excluded and failed samples and presented the failure rate at first attempt, the number of repeated tests and the final failure rate for each study. When it was possible, we also reported separate failure rates among aneuploid and euploid cases. Where possible, we performed subgroups analyses to investigate heterogeneity, and also performed sensitivity analyses to assess the robustness of these findings.

Weaknesses

Fetal karyotyping is the reference standard for establishing a diagnosis of fetal aneuploidy. This is an invasive procedure

with some risk for the fetus and the pregnant woman. Many pregnant women included in the studies, especially those involving unselected cohorts, were not tested by karyotyping. Rather, clinical examination of the newborn or medical records from birth were used as a secondary reference standard. We are aware that these secondary reference standards are not as accurate as fetal karyotype and some cases may have been missed.

Studies rarely reported the qualification of the person conducting the neonatal clinical examination at birth. Such examination is expected to be more reliable if it was made by a paediatrician or a geneticist. Ideally, this examination should be done a few months after birth because the phenotypic characteristics of aneuploidies are more apparent than at birth (Devlin 2004).

Genomics-based non-invasive prenatal testing assays are laboratory-developed tests that are not standardised in their methods, sequencing platforms, sequencing data manipulation, measures used or cut-offs for interpretation. Each assay was developed and validated by the testing laboratory and each laboratory has a different method. Usually detailed information about the assays were not available. As shown in Table 5, 15, different gNIPT assays were used in the studies included in this review. Thus, they may differ in various aspects and show different analytical and clinical validity. We have grouped them accordingly to the type of assay used (targeted versus shotgun), but there are also differences within each of these two subgroups that we were not able to account for, given the small number of studies published on most of these different assays. Thirteen of the assays were used only in studies of high-risk pregnancies or mixed cohorts. Only a few gNIPT assays were used in a significant number of studies. Thus, caution should be used before generalising the diagnostic accuracy observed in this category of patients to all gNIPT assays. This limits the generalisability of these findings and we cannot infer that all gNIPT assays will show the same performance.

Applicability of findings to the review question

These findings suggest that gNIPT has high sensitivity and specificity for detection of fetal aneuploidies in high-risk pregnancies. Performance varied depending on the type of aneuploidy. There was limited evidence of the performance of gNIPT in unselected cohorts of pregnant women. Most studies involved either high-risk pregnancies or mixed populations where it was not possible to differentiate between high-risk pregnancies and unselected pregnant women. Thus, more studies are needed in the general population of pregnant women before firm conclusions can be made about the sensitivity of gNIPT as a first-tier screening test. The two major types of gNIPT method (MPSS and TMPS) appear to have comparable performance, but there are many different gNIPT assays for each approach. For many of these assays, very little data have been published about their diagnostic accuracy. Additionally, performance in the cohorts studied may not reflect performance in other populations owing to differences in fetal fraction distribution because of, for example, differences in mean body mass index or gestational age. Importantly, summary sensitivities and specificities derived from cohort data can be very different from the probability associated with any particular patient sample to be positive or negative depending on the sample's specific fetal fraction. Thus, summary sensitivity, specificity and associated predictive values of an assay cannot be used as a straightforward measure of the probability of a specific patient's sample to be affected given a positive or negative result. This

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underscores the importance, before clinically offering a laboratory developed gNIPT assay, that it is fully validated according to recognised best practice clinical laboratory molecular diagnostics guidelines. Finally, the methodological quality of studies was generally poor with high risk of bias, especially in terms of patient selection and flow and timing.

AUTHORS' CONCLUSIONS

Implications for practice

Genomics-based non-invasive prenatal testing (gNIPT) appears to be an accurate prenatal screening test, its accuracy having been evaluated as a second-tier screening test to identify pregnancies at very low risk of fetal aneuploidies (T21, T18 and T13) and thus to decrease the false positive rate of traditional screening approaches and avoid invasive procedures in those pregnant women. As a first-tier aneuploidy screening test, based on limited data from comparative studies, gNIPT appears to have significantly better specificity than current screening approaches using maternal serum biochemical markers, ultrasound or both, but evidence about sensitivity is inconsistent. At current gNIPT pricing levels, gNIPT as a second-tier screening test provides the best value for money, especially for publicly-funded screening programs while gNIPT as a first-tier screening test was found not to be cost-effective (Nshimyumukiza 2017). The failure rate of gNIPT is a concern as it is substantially larger than the current failure rate of traditional prenatal screening approaches.

It is worth noting that gNIPT shows good performance for the detection of sex chromosome aneuploidies though data are sparse. The number of studies for sex chromosome aneuploidy (SCA) was small and confidence intervals on sensitivity and specificity estimates are therefore wide. Thus, sex chromosome aneuploidies appear to be more difficult to detect since performances of gNIPT are not as good as for detecting autosomal aneuploidies. SCAs are considered "incidental" findings of current aneuploidy screening programs and they do not lead to any intervention prior to birth.

Maternal serum screening, ultrasound fetal examination, gNIPT and invasive diagnostic tests are thus complementary approaches because in its current state, gNIPT cannot detect all chromosomal abnormalities or adverse obstetrical outcomes. About 44% to 64% of all chromosomal abnormalities found during prenatal diagnostic are common aneuploidies which gNIPT can detect (Kazerouni 2011; Shani 2016). Counselling expectant mothers and their partners is essential for explaining the advantages, limitations and risks of these procedures.

We conclude that given the current data on the performance of gNIPT, invasive fetal karyotyping is still the required diagnostic approach to confirm the presence of a chromosomal abnormality

prior to making irreversible decisions relative to the pregnancy outcome.

Implications for research

This systematic review has highlighted the fact that most published studies on gNIPT have high risk of bias in the patient selection and flow and timing domains. Many different gNIPT assays are in use and for the majority of them, there is insufficient published data to individually assess their clinical performance. Therefore, the results in this systematic review may not be generalisable to all gNIPT assays. Studies are needed that directly compare the accuracy of gNIPT with that of current traditional prenatal screening methods for fetal aneuploidy, especially in unselected populations of pregnant women. Such studies can provide valid evidence of the incremental accuracy of gNIPT if gNIPT is being considered as a first-tier test. Particular attention should be paid to study design in order to minimise patient selection biases as well as biases in flow and timing domain. Further well-designed, independent largescale studies on real life gNIPT's implementation into prenatal care should be performed. Large scale randomised clinical trials of tests and patient outcomes are needed to validate the clinical utility of gNIPT in the various clinical settings. Given the rapid evolution of gNIPT and its capacity to detect other fetal chromosomal anomalies (Benn 2016), future systematic reviews may have to widen the scope of target conditions.

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Badeau M, Lindsay C, Blais J, Takwoingi Y, Langlois S, Légaré F, et al. Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women.

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

Cochrane Database of Systematic Reviews 2015, Issue 7. [DOI: 10.1002/14651858.CD011767]

* Indicates the major publication for the study

Study characteristics	
Patient sampling	Study design: case-control study (1:2) from a prospective cohort. Participants: pregnant women selected from a high-risk population of fetal aneuploidy. Inclusion criteria: pregnant women who had a risk of fetal trisomy 21 (> 1 in 250), based on the combination of maternal age with ultrasound and maternal serum markers during the first or sec- ond trimester and prior invasive testing. Exclusion criteria: multifetal pregnancies, absence of medical coverage by the National Health Sys tem and women declining an invasive procedure.
Patient characteristics and setting	Number enrolled: 976 pregnant women. Number available for 2 x 2 table: 183 pregnant women (subgroup of 19%). 23 euploid samples were used as reference set and 8 samples randomly chosen for pretesting phase. Setting: 3 centres in France. Recruitment period: March 2010 to April 2013. Ethnicity: not reported. Mean gestational age (± SD): 14 (± 2) weeks. Mean maternal age (± SD): 35.2 (± 6.7) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement and biochemical screening. Language of the study: English.
Index tests	gNIPT by MPSS on Illumina HiSeq 2000 without multiplexing. Each library was sequenced using 50 bases-length reads chemistry in a single end-flow cell. Mean fetal fraction DNA: (male only) euploid: 20.11% and T21: 16.86%. Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if Z score > 3. In-house gNIPT.
Target condition and refer- ence standard(s)	Target condition: T21. Reference standard: fetal karyotype of chorionic villi or amniotic fluid.
Flow and timing	 Blood samples were obtained prior to the invasive procedure (reference standard). gNIPT was a second-tier test. 701/976 samples were not selected for the case-control study. 50/275 samples were excluded during DNA extraction (47 for low amount of DNA and 3 for haemolysis) (no gNIPT results). 31/225 samples were excluded from analysis (8 for pretesting phase and 23 for reference set). 11/194 samples were excluded from analysis for insufficient fetal fraction DNA (no gNIPT results). No repeated test reported.

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Alberti 2015 (Continued)			
Aim to study			nto a cytogenetics laboratory in a universi- e on samples collected prospectively.
Funding source or sponsor of the study	Study not funded by indust	ry.	
Informations about the au- thors contacted	Authors were contacted on Last reply received on: 16 M		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard	d		
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard	Yes		

DOMAIN 4: Flow and Timing

results interpreted without knowledge of the results of the

index tests?

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Low

Low



Alberti 2015 (Continued)	
Was there an appropriate in- terval between index test and reference standard?	Yes
Did all analysed patients re- ceive the reference standard?	Yes
Were all patients included in the analysis?	No

High

Ashoor 2012

Study characteristics	
Patient sampling	Study design: nested case-control study (1:3) from a prospective cohort. Participants: pregnant women selected from a high-risk population (archived maternal plas- ma samples) of fetal aneuploidy.
	Inclusion criteria: singleton pregnancies between 11 to 13 weeks' gestation. Exclusion criteria: pregnancies that were conceived by in vitro fertilization.
Patient characteristics and setting	Number enrolled: 400 pregnant women. Number available for 2 x 2 table: 397 archived plasma samples (subgroup of 99%). Setting: 1 centre. Tertiary Referral Centre, King's College Hospital, London, United Kingdom. Recruitment period: March 2006 to August 2011. Ethnicity: Caucasian (88.5%), Afro-Caribbean (5%), South Asian (4%), East Asian (2%) and multiracial (0.5%). Mean gestational age (range): 13.3 (12.1 to 13.7) weeks. Mean maternal age (range): 36.2 (29.9 to 41.2) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measure- ment) and biochemical screening. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex.
	Fetal fraction DNA: not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Usually, Harmony™ prenatal test uses FORTE algorithm; positive if FORTE risk score ≥ 1%. Commercial test: Harmony™ prenatal test by Ariosa Diagnostics, Inc.
Target condition and reference standard(s)	Target conditions: T21 and T18. Reference standard: fetal karyotype of chorionic villi.
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).
	gNIPT was a second-tier test. 3/400 samples failed amplification and sequencing (no gNIPT result).
	25 samples did not meet Ariosa Diagnostics, Inc acceptance criteria but they were replaced with the next available cases.
	No repeated test reported.



Continued)

Ashoor 2012 (Continued) Aim to study	To assess the prenatal detection rate of T21 and T18 and the false-positive rate by chromo- some-selective sequencing of maternal plasma ccfDNA.		
Funding source or sponsor of the study	Study not funded by industry but samples were analysed at Ariosa Diagnostics, Inc.		
Informations about the authors con- tacted	No need for further contact	t.	
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	No		
		High	Low
DOMAIN 2: Index Test TMPS			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		



Ashoor 2012 (Continued)

Did all analysed patients receive the	Yes
reference standard?	

Were all patients included in the No analysis?

High

Ashoor 2013

Study	charact	eristics
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Patient sampling	Study design: blinded prospective cohort (second phase). First phase (case-control study) not shown in the present review. Participants: euploid pregnancies underwent routine first-trimester combined screening and confirmed T13 cases were selected. Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies.
Patient characteristics and set- ting	Number enrolled: 2167 pregnant women. Number available for 2 x 2 table: 1949 pregnant women (subgroup of 90%). Setting: several centres. Euploid pregnancies were from King's College Hospital, London, UK and T13 cases were from the USA. Recruitment period: October 2010 to January 2011 for euploid pregnancies. Not reported for T13 cases. Ethnicity: Caucasian (70.8%), African (20%), Asian (6.8%), mixed (2.6%). Mean gestational age (± SD; range): 12.7 (± 0.62; 13 to 26) weeks. Mean maternal age (± SD): 31.8 (± 5.6) years. Relevant tests carried out prior to index test: not reported. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex. Median fetal fraction DNA (range): euploids: 10.0% (4.1% to 31.0%) and T21: 14.0% (6.1% to 24.0%). Blood samples for gNIPT were collected before reference standard for euploid pregnancies. T13 samples were collected post-confirmation of trisomy by karyotyping (reference standard). Cutpoint: positive if FORTE algorithm risk score ≥ 1%. Commercial test: Harmony [™] Prenatal Test by Ariosa Diagnostics, Inc.
Target condition and reference standard(s)	Target condition: T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or neonatal clinical ex- amination.
Flow and timing	 Blood samples were obtained at the time of screening for euploid pregnancies (before reference standard). Blood samples were obtained after T13 confirmation following invasive procedure (reference standard). gNIPT was a first- or a second-tier test. 165/2167 samples were excluded because they were used in the first phase. 53/2002 samples failed during amplification or sequencing (no gNIPT result). No repeated test reported.

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Comparative			
Aim to study	To assess the performance of chromosome-selective sequencing of maternal plasma cell-free DNA (cfDNA) in non-invasive prenatal testing for trisomy 13.		
Funding source or sponsor of the study	Study not funded by industry but samples were analysed at Ariosa Diagnostics, Inc.		
Informations about the authors contacted	No need for further contact.		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	No		
		High	High
DOMAIN 2: Index Test TMPS			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			



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Benachi 2015 Study characteristics	
	High
Were all patients included in the analysis?	No
Did all analysed patients receive the reference standard?	Yes
Was there an appropriate interval between index test and reference standard?	Yes
Ashoor 2013 (Continued)	

Study design: blinded, retrospective analysis from a prospective cohort. Participants: all pregnant women considered at high risk of fetal aneuploidies who were willing to undergo invasive procedure. Inclusion criteria: at least 18 years old, more than 10 weeks of gestation and singleton or twin preg- nancies. Exclusion criteria: vanishing twin or < 18 years old.
Number enrolled: 900 pregnant women. Number available for 2 x 2 table: 886 pregnant women (subgroup of 98%). Setting: 29 centres. French Fetal Medicine Centres in France. Recruitment period: December 2012 to October 2013. Ethnicity: Caucasian (84.2%), Black or Caribbean (4.6%), Asian (2.0%), mixed (5.7%) and unknown (3.5%). Median gestational age (range): 15.1 (10.2 to 34.6) weeks. Median maternal age (range): 35 (30 to 39) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening. Language of the study: English.
gNIPT by MPSS with Illumina v3 flow-cell on a HiSeq 1500 sequencer in 12-plex. Mean fetal fraction DNA: group 1 (patients without abnormal fetal ultrasound findings, but at high risk of fetal aneuploidy): 10.9% and group 2 (high risk of fetal aneuploidy after ultrasound finding): 11.2%. Blood samples for gNIPT were collected just before reference standard. Cutpoint: positive if Z score > 3 (T21) or > 3.95 (T18 and T13). Commercial test: Laboratoire CERBA's prenatal test.
Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or neonatal clinical examina tion.
Blood samples were obtained prior to the invasive procedure (reference standard). gNIPT was a second-tier test. 8/900 samples without karyotype result were excluded. 42 samples failed the initial MPSS testing for technical issues. 42/42 repeated tests using a second aliquot and 36/42 samples obtained gNIPT results.



Benachi 2015 (Continued)

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Benachi 2015 (Continued)	6/892 samples failed during a gNIPT result).	gNIPT process (low fetal fracti	on DNA or result appeared atypical) (no
Comparative			
Aim to study	To evaluate the performance of the gNIPT (using fetal ccfDNA) for detection of the 3 main autosomal fetal trisomies in a very high-risk population of patients whose fetuses display ultrasonographically identified anomalies by comparing the results with those obtained by conventional fetal karyotyp-ing.		
Funding source or sponsor of the study	Funding source not reported	. 1 author is an employee of L	aboratoire CERBA and also a shareholder.
Informations about the au- thors contacted	Authors were contacted on: 2 Reply received on: 26 May 20		
Notes	Authors are from de Collabo	rative SEquençage a Haut Deb	it et Aneuploidies (SEHDA) Study Group.
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standa	rd		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		



Benachi 2015 (Continued)

		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		

High

Bevilacqua 2015

Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: pregnant women between 10 to 28 weeks' gestation selected at high risk of fetal tri- somy or women who wanted to have the new test as a primary method of screening (unselected population). Inclusion criteria: singleton (not reported in the present review) or twin pregnancies between 10 to 28 weeks' gestation. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 2362 pregnant women including 1847 singleton pregnancies (data not reported in the present review) and 515 twin pregnancies. Number available for 2 x 2 table: 340 twin pregnancies (subgroup of 66%). Setting: multicentre. Recruitment period: May 2013 to September 2014 (twin). Ethnicity: not reported. Median gestational age (range): 13.0 (10 to 28) weeks. Median maternal age (range): 36.8 (19 to 50.3) years. Chorionicity (368/515): 13% monochorionic and 58.4% dichorionic. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening for some women. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex. Mean fetal fraction DNA (range): twins: 8.7% (4.1% to 30.0%) and singleton: 11.7% (4.0% to 38.9%). Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Usually, Harmony [™] prenatal test uses FORTE algorithm; positive if FORTE risk score ≥ 1%. Commercial test: Harmony [™] Prenatal Test by Ariosa Diagnostics, Inc.
Target condition and refer- ence standard(s)	Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or neonatal karyotype.
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard). gNIPT was a first- or second-tier test.



Bevilacqua 2015 (Continued)			
		ollow-up were excluded inclu irths without karyotype of fe	ıding138 lost to follow-up, 19 still pregnant tal tissue.
	29/515 samples failed the i 26/29 samples resequence a gNIPT result.		e first sampling and 13/26 samples obtained
	16/515 samples failed duri	ng sequencing process (no gl	NIPT result).
Comparative			
Aim to study			of maternal blood in screening for T21, T18 ne variables that could influence the failure
Funding source or sponsor of the study	Study not funded by indust	ry but Ariosa Diagnostics, Ind	c made blinded sequencing and analysis.
Informations about the au-	Author was contacted on: 1	June and 27 September 201	6.
thors contacted	No replies received from th	e author.	
Notes	gNIPT results from singletc 2 tables.	n pregnancies were not repo	rted in the present review for incomplete 2 x
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	No		
		High	High
DOMAIN 2: Index Test TMPS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low



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Bevilacqua 2015 (Continued)			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Bianchi 2012

Study characteristics	
Patient sampling	Study design: nested case-control (1:4) study from the MELISSA prospective cohort. Participants: pregnant women randomly selected from a high-risk population (archived maternal plasma samples). Inclusion criteria: singleton pregnancies at high risk of fetal aneuploidy between 8 and 22 weeks of gesta- tion. Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 2882 pregnant women. Number available for 2 x 2 table: 503 pregnant women for T21, 502 for T18, 501 for T13 and 489 for 45,X (subgroup of 17%). Setting: 60 centres. Medical centre in 25 states in USA. Samples from 53 centres were analysed. Recruitment period: June 2010 to August 2011. Ethnicity: Caucasian (72.7%), Afro American (10.9%), Asian (9.9%), Native American or Alaska Native (0.9%) and multiracial (5.6%). Mean gestational age (± SD; range): 15.1 (± 3.16; 10 to 23) weeks. Mean maternal age (± SD; range): 35.2 (± 6.40; 18 to 46) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening. Language of the study: English.
Index tests	gNIPT by MPSS on Illumina HiSeq 2000 sequencer in 6-plex. Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: 1) for T21, T18, and T13: positive if NCV > 4 (aneuploidy suspected if NCV is between 2.5 and 4).

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Bianchi 2012 (Continued)	2) for 45 V: positivo if NOV f	or Chrom. X < -4 and NCV for Chro	Nm V < 2.5
	- -	/ for Chrom. X > 4 and NCV for Ch	
		/ for Chrom. X between -2.5 and 2	
	5) for 47,XYY: positive if NCV greater than expected NCV Commercial test: Verinata's	Chrom. X.	nrom. Y > 4 with NCV for Chrom. Y is 2 times
Target condition and reference standard(s)		, T13, 45,X, 47,XXX, 47,XXY and 47 aryotype of chorionic villi (42.7%	7,XYY.), amniotic fluid (56.4%) or products of con-
Flow and timing	Blood samples were obtain	ed prior to the invasive procedu	re (reference standard).
	gNIPT was a second-tier tes 257/2882 samples were exc nancies).		ithout karyotype and 85 for multifetal preg-
	2091/2625 samples were no	ot selected for this case-control s	tudy.
	2/534 samples were exclud	ed for tracking issue.	
	16/532 samples without fet	al DNA detected were excluded o	during process (no gNIPT result).
	13/516 samples were exclu	ded of T21 2 x 2 table for censore	d complex karyotype.
	14/516 samples were exclu	ded of T18 2 x 2 table for censore	d complex karyotype.
	15/516 samples were exclu	ded of T13 2 x 2 table for censore	d complex karyotype.
	27/516 samples were exclu	ded of 45,X 2 x 2 table for censore	ed complex karyotype.
	No repeated test reported.		
Comparative			
Aim to study	To prospectively determine chromosome fetal aneuplo		ively parallel sequencing to detect whole
Funding source or spon- sor of the study	Study funded by Verinata H	ealth, Inc. (a wholly owned subsi	idiary of Illumina, Inc.).
Informations about the authors contacted	Authors were contacted on Replies received on: 1 Marc	: 1 March and 30 November 2016 h and 8 December 2016.	
Notes	This study is a clinical trial.	MELISSA study. Clinicaltrials.gov	NCT01122524.
	Data for 47,XXY, 47,XYY and view).	47,XXX were incomplete in the p	ublication (data not shown in the present re-
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selec	tion		
Was a consecutive or random sample of pa- tients enrolled?	No		



Bianchi 2012 (Continued)				
Was a case-control de- sign avoided?	No			
Did the study avoid in- appropriate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test M	PSS			
Were the index test re- sults interpreted with- out knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference St	andard			
Is the reference stan- dards likely to correctly classify the target con- dition?	Yes			
Were the reference standard results inter- preted without knowl- edge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Tin	ning			
Was there an appropri- ate interval between in- dex test and reference standard?	Yes			
Did all analysed pa- tients receive the refer- ence standard?	Yes			
Were all patients in- cluded in the analysis?	No			
		High		

Bianchi 2013

Study characteristics



Bianchi 2013 (Continued)	
Patient sampling	Study design: retrospective study (archived maternal plasma samples) from a prospective co- hort. Participants: pregnant women selected from a high-risk population (archived maternal plasma samples). Inclusion criteria: eligible blood samples, singleton pregnancies with karyotype result and nuchal cystic hygroma on fetal ultrasound.
	Exclusion criteria: multifetal pregnancies.
Patient characteristics and set- ting	Number enrolled: 2882 pregnant women. Number available for 2 x 2 table: 113 pregnant women (subgroup of 4%). Setting: 60 centres in USA. Recruitment period: June 2010 to August 2011. Ethnicity: Caucasian (73%), Afro-American (10%), Asian (9%) and multiracial (8%). Mean gestational age (± SD): 13.2 (± 2.0) weeks.
	Median gestational age (range): 12.6 (10 to 21) weeks. Mean maternal age (± SD): 32.2 (± 5.8) years. Median maternal age (range): 32.9 (18 to 44) years. Relevant test carried out prior to index test: ultrasonography (nuchal translucency measure- ment). Language of the study: English.
Index tests	gNIPT by MPSS with the sequencing chemistry Illumina TrueSeq 3.0.
	Fetal fraction DNA: not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint:
	1) for T21, T18 and T13: positive if NCV > 4 (aneuploidy suspected zone between 3 and 4).
	2) for 45,X: positive if NCV Chrom. X < -3 and NCV Chrom. Y < 3. Commercial test: Verinata's prenatal test.
Target condition and reference standard(s)	Target conditions: T21, T18, T13 and 45,X. Reference standard: fetal karyotype of chorionic villi (78%), amniotic fluid (20%) or products of conception (2%).
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).
	gNIPT was a second-tier test. 2769/2882 samples were not selected for this study.
	No failed sample reported.
	No repeated test reported.
Comparative	
Aim to study	To estimate the accuracy and potential clinical effect of using massively parallel sequencing of maternal plasma DNA to detect fetal aneuploidy in a population of pregnant women carrying fe- tuses with nuchal cystic hygroma.
Funding source or sponsor of the study	Study funded by Verinata Health, Inc. (a wholly owned subsidiary of Illumina, Inc.).
Informations about the authors contacted	No need for further contact.
Notes	74/113 samples were previously sequenced during the MELISSA trial. In this study, all 113 sam- ples were newly resequenced (no overlap) with MELISSA study.



Bianchi 2013 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	Yes		
		Low	

Bianchi 2014a **Study characteristics** Study design: blinded, prospective cohort study. Patient sampling Participants: pregnant women who planned to undergo (without prior risk) or had completed (highor low-risk) standard prenatal serum screening for fetal aneuploidy. Inclusion criteria: pregnant women of 18 years or older, gestational age \geq 8 weeks, able to provide consent and pregnancy records accessible and available for data collection. Exclusion criteria: invasive procedure (amniocentesis or CVS) performed within 2 weeks prior enrolment or prenatal screening determination by nuchal translucency measurement only. Patient characteristics and Number enrolled: 2052 pregnant women. Number available for 2 x 2 table: 1952 for T21 and T18 (subgroup of 95%) and 1914 for T13 (subgroup setting of 93%). Setting: 21 centres. In 14 states (USA). Recruitment period: 2 July 2012 to 4 January 2013. Ethnicity: Caucasian (65.4%), Afro-American (22.3%), Asian (7.3%) and other (5%). Mean gestational age (± SD; range): 20.3 (± 8.6; 8 to 39.4) weeks. Mean maternal age (± SD; range): 29.6 (± 5.54; 18 to 48.6) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening. Language of the study: English. Index tests gNIPT by MPSS on Illumina HiSeq 2000 in 8-plex. Mean fetal fraction DNA: more than 35 years old: 11.3%, less than 35 years old: 11.6%, and at third trimester only: 24.6%. Blood samples for gNIPT were collected before or after reference standard. Cutpoint: positive if NCV \geq 4. Resequenced if NCV is between 3 and 4. Commercial test: verifi[®] prenatal test by Verinata Health. The traditional screening tests (first-trimester combined test or a second-trimester result (quadruple, serum integrated, fully integrated or sequential)) were also assessed. Mixed cutpoints used. Target condition and refer-Target conditions: T21, T18 and T13. ence standard(s) Reference standards: fetal karyotype of chorionic villi, amniotic fluid or products of conception, neonatal clinical examination or medical record from birth. Flow and timing Blood samples were obtained prior or after the invasive procedure (reference standard). gNIPT was a first- or second-tier test. 10/2052 samples failed blood quality control before sequencing process. 72/2042 samples without clinical outcome. 38/2042 samples without standard screening result. 17/2042 samples without gNIPT result. 1/2042 samples without standard screening result and without gNIPT result. 12 resequenced samples were in the grey zone (between affected and unaffected) and were successfully resequenced in uniplex. Comparative Aim to study To compare the results of gNIPT with ccfDNA for fetal autosomal aneuploidy with the results of conventional screening for T21 and T18 in a general obstetrical population. To compare false positive rates with the use of each method. To compare false positive rates for T13 in a subset of pregnant women in whom standard screening results included a risk assessment for trisomy 13. To compare fe-



Bianchi 2014a (Continued)

tal ccfDNA fractions in low-risk patients and those in high-risk patients in the CARE study population to assess the potential effects of demographic differences on test performance.

Funding source or sponsor of the study	Study funded by Illumina, Inc.
Informations about the au- thors contacted	Author was contacted on: 10 February, 1 June and 28 June 2016. No replies received from the author.
Notes	This study is a clinical trial (Comparison of Aneuploidy Risk Evaluations; CARE study). ClinicalTrial- s.gov number: NCT0166335.

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns			
DOMAIN 1: Patient Selection	DOMAIN 1: Patient Selection					
Was a consecutive or ran- dom sample of patients en- rolled?	Unclear					
Was a case-control design avoided?	Yes					
Did the study avoid inap- propriate exclusions?	No					
		High	High			
DOMAIN 2: Index Test MPSS						
Were the index test results interpreted without knowl- edge of the results of the reference standard?	Yes					
If a threshold was used, was it pre-specified?	Yes					
		Low	Low			
DOMAIN 2: Index Test Tradit	tional screening tests					
Were the index test results interpreted without knowl- edge of the results of the reference standard?	Yes					
If a threshold was used, was it pre-specified?	Yes					
		Low	Low			
DOMAIN 3: Reference Stand	ard					



Bianchi 2014a (Continued)			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing	3		
Was there an appropriate interval between index test and reference standard?	Yes		
Did all analysed patients receive the reference stan- dard?	Yes		
Were all patients included in the analysis?	No		
		High	

Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: pregnant women selected at high risk of fetal aneuploidy. Inclusion criteria: pregnant women at high risk of fetal aneuploidy with invasive test result. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 10 pregnant women. Number available for 2 x 2 table: 9 pregnant women (subgroup of 90%). Setting: obstetric and gynaecology clinic in Warsaw, Poland. Recruitment period: not reported. Ethnicity: not reported. Median gestational age (range): 16 (13 to 23) weeks. Median maternal age (range): 31 (26 to 36) years. Relevant test carried out prior to index test: ultrasonography (nuchal translucency mea- surement). Language of the study: Polish.
Index tests	gNIPT by MPSS on Illumina Genome Analyzer IIx or HiSeq 2000 sequencer in multiplex with BGI's algorithm. Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Commercial test: NIFTY™ test by Bejing Genomics Institute.
Target condition and reference stan- dard(s)	Target conditions: T21, T18, and T13. Reference standard: fetal karyotype of chorionic villi (30%) or amniotic fluid (70%).

Blood samples were obtained prior to the invasive procedure (reference standard).		
		T result) for low fetal fraction DNA.
No repeated test reported.		
T18, and T13) based on ccfD	NA in maternal plasma in h	
Study not funded by industry but NIFTY™ tests were provided by Beijing Genomics Institute, Shenzen, China.		
Authors were contacted on: 2 May and 4 July 2016. Replies received on: 4 and 16 May 2016.		
Authors' judgement	Risk of bias	Applicability concerns
Unclear		
Yes		
Unclear		
	Unclear	Low
Yes		
Yes		
	Low	Low
Yes		
Yes		
	gNIPT was a second-tier test 1/10 sample failed during set No repeated test reported. To present initial results of r T18, and T13) based on ccfD compare the results with ro Study not funded by industr Shenzen, China. Authors were contacted on: Replies received on: 4 and 1 Muthors' judgement Unclear Yes Unclear Yes Yes Yes Yes	gNIPT was a second-tier test. 1/10 sample failed during sequencing process (no gNIP No repeated test reported. To present initial results of non-invasive prenatal diagr T18, and T13) based on ccfDNA in maternal plasma in f compare the results with routine karyotyping. Study not funded by industry but NIFTY TM tests were pr Shenzen, China. Authors were contacted on: 2 May and 4 July 2016. Replies received on: 4 and 16 May 2016. Muthors' judgement Risk of bias Unclear Yes Unclear Yes Yes Yes Yes Yes



Bijok 2014 (Continued)

Trusted evidence. Informed decisions. Better health.

		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Canick 2012

Patient sampling	Study design: case-control study. Participants: all multifetal pregnant women with T21, T18 or T13 fetus were selected along with all euploid triplet pregnancies and a random selection of euploid twin pregnancies. Inclusion criteria: multifetal pregnant women, at least 18 years old, between about 10 weeks and 21 weeks 6 days of gestation, at high risk of aneuploidies and who undergo an invasive pro cedure. Exclusion criteria: singleton pregnancies or low risk of aneuploidy.
Patient characteristics and set- ing	Number enrolled: 4664 pregnant women. Number available for 2 x 2 table: 27 multifetal pregnancies (25 twin and 2 triplet pregnancies) (subgroup of 0.6%). Setting: 27 centres. Prenatal diagnostic centres (Canada, Italy, Spain, Czech Republic, Argentin Ireland, Hungary, USA, Israel and Australia). Recruitment period: April 2009 to February 2011. Ethnicity: not reported. Mean gestational age (range): 15.0 (10.9 to 19) years. Maternal age: not reported. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measure- ment) and biochemical screening. Language of the study: English.
ndex tests	gNIPT by MPSS on Illumina HiSeq 2000 sequencer in 4-plex. Fetal fraction DNA range: 7% to 55%. Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if Z score ≥ 3. Commercial test: Sequenom's test.
Target condition and reference standard(s)	Target conditions: T21 and T13. T18 was also assessed but no case was found. Reference standard: fetal karyotype of chorionic villi or amniotic fluid.
Flow and timing	Blood samples were obtained immediately prior the invasive procedure (reference standard). gNIPT was a second-tier test. 4637/4664 samples were not selected for this case-control study.



Canick 2012 (Continued)

No repeated test reported.

Comparative	
Aim to study	To study prenatal testing for T21, T18, and T13 by MPSS of fetal ccfDNA in high-risk multifetal pregnant women.
Funding source or sponsor of the study	Study funded by Sequenom, Inc. Some authors are employees and shareholders of Sequenom, Inc. or of Sequenom Center for Molecular Medicine.
Informations about the authors contacted	Author was contacted on: 10 March 2016. Reply received on: 16 March 2016.
Notes	This study is a clinical trial "A New Prenatal Blood Test for Down Syndrome" ClinicalTrials.gov number: NCT00877292.

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoid- ed?	No		
Did the study avoid inappropriate exclusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes		
		Low	Low



DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	Yes
Did all analysed patients receive the reference standard?	Yes
Were all patients included in the analysis?	Yes
	Low

Study characteristics			
Patient sampling	Study design: nested case-control study from a prospective cohort and archived plasma.		
	Participants: pregnant women with clinical indications of fetal aneuploidy (high risk of feta aneuploidy) for invasive procedure.		
	Inclusion criteria: singleton pregnancies with and without trisomy 13, 18 or 21, matched for gestational ages.		
	Exclusion criteria: twin pregnancies.		
Patient characteristics and setting	Number enrolled: 392 pregnant women (252 from the prospective cohort and 140 were archived plasma).		
	Number available for 2 x 2 table: 289 pregnant women (subgroup of 74%).		
	Setting: 10 centres in Hong Kong, the Netherlands, and UK.		
	Recruitment period for the prospective cohort: October 2008 to May 2009.		
	Recruitment period for the archived plasma samples collection: October 2003 to Septembe 2008.		
	Ethnicity: not reported.		
	Gestational age: not reported.		
	Maternal age: not reported.		
	Relevant tests carried out prior to index test: ultrasonography (nuchal translucency mea- surement) and biochemical screening.		
	Language of the study: English.		
ndex tests	gNIPT by MPSS on Illumina Genome Analyzer IIx in 2-plex.		
	Feta fraction DNA: not reported.		
	Blood samples for gNIPT were collected before reference standard.		
	Cutpoint: positive if Z score > 3.		
	Commercial test: Sequenom's test.		

Chen 2011 (Continued)					
Target condition and reference stan- dard(s)	Target conditions: T18 and T13.				
	Reference standard: fetal karyotype of chorionic villi or amniotic fluid.				
Flow and timing	Blood samples were obtaine	ed prior to the invasive proc	cedure (reference standard).		
	gNIPT was a second-tier tes	t.			
	103/392 samples were selected as reference control.				
	No failed sample reported.				
	No repeated test reported.				
Comparative					
Aim to study	To assess the prenatal diagr of pregnant women with T1		S of maternal plasma DNA on a cohort		
Funding source or sponsor of the study	Study co-sponsored by Sequenom, Inc and Life Technologies. Some authors have filed patent on gNIPT (part of this patent has been licensed to Sequenom, Inc).				
Informations about the authors con-	Author was contacted on: 14 December 2015 and 10 May 2016.				
tacted	Reply received on: 12 May 2016.				
Notes					
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	No				
Was a case-control design avoided?	No				
Did the study avoid inappropriate ex- clusions?	No				
		High	Low		
DOMAIN 2: Index Test MPSS					
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Unclear				
If a threshold was used, was it pre- specified?	Yes				
		Unclear	Low		
DOMAIN 3: Reference Standard					

Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review) Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.



Chen 2011 (Continued)			
Is the reference standards likely to correctly classify the target condi-tion?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	Yes		
		Low	

Chiu 2011

Study characteristics	
Patient sampling	Study design: blinded, case-control study (1:5) from a prospective cohort and archived plasma.
	Participants: pregnant women with clinical indications for invasive procedure, mixed risk (mostly high risk (> 1/300 at traditional screening test), intermediate risk (between 1/300 and 1/1000) or oth- er risk factors).
	T21 and non T21 pregnancies matched for gestational ages.
	Inclusion criteria: singleton pregnancies.
	Exclusion criteria: multifetal pregnancies.
Patient characteristics and	Number enrolled: 824 pregnant women.
setting	Number available for 2 x 2 table: 753 (8-plex) (subgroup of 91%).
	Setting: 10 centres in Hong Kong, the Netherlands, and UK.
	Recruitment period for the prospective cohort: October 2008 to May 2009.
	Recruitment period for the archived plasma samples collection: October 2003 to September 2008.
	Ethnicity: not reported.
	Median gestational age: 13.1 weeks.
	Median maternal age: 35.4 years.
	Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening.



Chiu 2011 (Continued)

	Language of the study: Eng	lish.			
Index tests	gNIPT by MPSS on Illumina Genome Analyzer II in 8-plex and 2-plex (not reported in the present re- view).				
	Median fetal fraction DNA (interquartile 1 and 3): male euploid: 15.2% (10.6% and 19.1%), archived samples: 14.7%, and prospective samples: 15.4%.				
	Blood samples for gNIPT w	ere collected before reference st	tandard.		
	Cutpoint: positive if Z score	> 3.			
	Commercial test: Sequenor	n's test.			
Target condition and refer-	Target condition: T21.				
ence standard(s)	Reference standard: fetal k	aryotype of chorionic villi or am	niotic fluid.		
Flow and timing	Blood samples were obtain	ed prior to the invasive procedu	ire (reference standard).		
	gNIPT was a second-tier tes	st.			
	60/824 samples were exclu and 46 failed quality contro		(2 twin pregnancies, 12 without karyotype		
	11/764 samples failed quality control during sequencing process (no gNIPT result).				
	96/753 samples were also used for reference controls (8-plex).				
	No repeated test reported.				
Comparative					
Aim to study	To validate the diagnostic performance and practical feasibility of massively parallel genomic se- quencing for the non-invasive prenatal assessment of trisomy 21 in pregnant women who had un- dergone conventional screening and were clinically indicated for definitive testing.				
Funding source or sponsor of the study	Study sponsored by Seque this patent has been license		ed patent applications on gNIPT (part of		
Informations about the au- thors contacted	No need for further contact				
Notes		ng were excluded from the prese use it is the method most likely t	nt review to avoid double counting. We to be used for routine testing.		
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	No				
	No				
Was a case-control design avoided?					



Chiu 2011 (Continued)			
		High	High
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standa	rd		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Comas	2015
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Study characteristics	
Patient sampling	Study design: blinded, observational prospective cohort study. Participants: all pregnant women who underwent conventional first-trimester combined screening for fetal aneuploidies (without prior risk of fetal aneuploidy). Some pregnant women were referred after their combined test (high risk of fetal aneuploidy). Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies, cases of ultrasound anomalies, nuchal translucency > 99 centile, combined risk at first-trimester screening > 1/10, or women at high risk of other genetic condi- tions.
Patient characteristics and setting	Number enrolled: 333 pregnant women (85.5% without prior risk and 16.5% were at high risk of fetal aneuploidy).



Mean fetal fraction DNA (range): 12.7% (4.2% to 27.9%), Harmony™ prenatal test ma™ prenatal test: 12.7%. Blood samples for gNIPT were collected before reference standard. DANSR assay cutpoint: not reported. Usually, Harmony™ prenatal test uses FOF if FORTE risk score ≥ 1%. SNP-based method cutpoint: not reported. Commercial test: Panorama™ prenatal test by Natera, Inc. or Harmony™ prenat nostics, Inc. Farget condition and ref- erence standard(s) Target conditions: T21, T18, T13, 45,X, 47,XXX, 47, XXY, 47,XYY. SCA data were no present review. T18 and T13 were also assessed but no case was found. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or neon tion. Flow and timing Blood samples were obtained prior to the invasive procedure (reference standard gNIPT was a first- or second-tier test. 17/333 samples failed the initial TMPS testing. 6/9 repeated sampling was performed and results were obtained in 5/6. 3/333 samples without gNIPT result and follow-up were excluded (still pregnan ting. 1/333 samples without gNIPT result and follow-up were excluded (still pregnan ting. Comparative Xim to study To evaluate gNIPT of ccfDNA as a screening method for major chromosomal an ting. Funding source or sponsor of the study Study not funded by industry but Ariosa Diagnostics, Inc and Natera, Inc. made sis.	ability concerns		
Mean gestational age (range): 37 (21 to 46) years. Relevant test carried out prior to index test: biochemical screening for a part of Language of the study: English Index tests gNIPT by TMPS (DANSR assay or SNP-based method). Mean fetal fraction DNA (range): 12.7% (4.2% to 27.9%), Harmony TM prenatal test ma TM prenatal test: 12.7%. Blood samples for gNIPT were collected before reference standard. DANSR assay cutpoint: not reported. Commercial test: Panorama TM prenatal test by Natera, Inc. or Harmony TM prenatal test uses FOF if FORE risk score 2 1%. SNP-based method cutpoint: not reported. Commercial test: Panorama TM prenatal test by Natera, Inc. or Harmony TM prenatal nostics, Inc. Target condition and ref- Target conditions Target conditions and reference standards: fetal karyotype of chorionic villi or amniotic fluid or neon tion. Flow and timing Blood samples were obtained prior to the invasive procedure (reference standards) gNIPT was a first- or second-tier test. 17/333 samples actiled the initial TMPS testing. 6/9 repeated sampling was performed and results were obtained in 5/6. 3/333 samples without gNIPT result and follow-up were excluded (still pregnan ting. 1/333 samples without gNIPT result and follow-up were excluded (still pregnan ting. Comparative To evaluate gNIPT of ccfDNA as a screening method for major chromosomal an ting. Author was contacted on			
Mean gestational age (range): 14 6 (9.5 to 23.5) weeks. Mean maternal age (range): 37 (21 to 46) years. Relevant test carried out prior to index test: biochemical screening for a part of Language of the study: English Index tests gNIPT by TMPS (DANSR assay or SNP-based method). Mean fetal fraction DNA (range): 12.7% (4.2% to 27.9%), Harmony™ prenatal test ma™ prenatal test: 12.7%. Biood samples for gNIPT were collected before reference standard. DANSR assay cutpoint: not reported. Usually, Harmony™ prenatal test uses FOF if FORTE risk score ≥ 1%. SNP-based method cutpoint: not reported. Commercial test: Panorama™ prenatal test by Natera, Inc. or Harmony™ prenat nostics, Inc. Target condition and ref- erence standard(s) Target conditions: T21, T18, T13, 45,X, 47,XXX, 47,XYY, 47,XYY, SCA data were no present review. T18 and T13 were also assessed but no case was found. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or neon tion. Flow and timing Blood samples were obtained prior to the invasive procedure (reference standard gNIPT was a first- or second-tier test. 17/333 samples excluded because still pregnant at the time of publication (no f 9/333 samples failed the initial TMPS testing. 6/9 repeated sampling was performed and results were obtained in 5/6. 3/333 samples without gNIPT result were excluded (unrepeated samples). 1/333 samples without gNIPT result and follow-up were excluded samples). 1/333 samples without gNIPT result and follow-up were excluded (still pregnan ting. Comparative To evaluate gNIPT of ccfDNA as a screening method for major chromosomal an ting. Funding source or sponsor of the study Study not funded by industry but Ariosa Diagnostics, Inc and Natera, Inc. made sis. <td></td>			
Mean gestational age (range): 14.6 (9.5 to 23.5) weeks. Mean maternal age (range): 37 (21 to 46) years. Relevant test carried out prior to index test: biochemical screening for a part of Language of the study: English Index tests gNIPT by TMPS (DANSR assay or SNP-based method). Mean fetal fraction DNA (range): 12.7% (4.2% to 27.9%), Harmony™ prenatal test ma™ prenatal test: 12.7%. Blood samples for gNIPT were collected before reference standard. DANSR assay cutpoint: not reported. Commercial test: Panorama™ prenatal test by Natera, Inc. or Harmony™ prenatal test is score ≥ 1%. SNP-based method cutpoint: not reported. Commercial test: Panorama™ prenatal test by Natera, Inc. or Harmony™ prenat nostics, Inc. Target conditions: T21, T18, T13, 45,X, 47,XXX, 47, XXY, 47,XYY. SCA data were no present review. T18 and T13 were also assessed but no case was found. Reference standard(s) Reference standards: fetal karyotype of chorionic villi or anniotic fluid or neon tion. Flow and timing Blood samples were obtained prior to the invasive procedure (reference standard gNIPT was a first- or second-tier test. 17/333 samples failed the initial TMPS testing. 6/9 repeated sampling was performed and results were obtained in 5/6. 3/333 samples without gNIPT result and follow-up were excluded (still pregnan ting. 1/333 samples without gNIPT result and follow-up were excluded (still pregnan ting. Comparative To			
Mean gestational age (range): 14.6 (9.5 to 23.5) weeks. Mean maternal age (range): 37 (21 to 46) years. Relevant test carried out prior to index test: biochemical screening for a part of Language of the study: English Index tests gNIPT by TMPS (DANSR assay or SNP-based method). Mean fetal fraction DNA (range): 12.7% (4.2% to 27.9%), Harmony™ prenatal test ma™ prenatal test: 12.7%. Blood samples for gNIPT were collected before reference standard. DANSR assay cutpoint: not reported. Usually, Harmony™ prenatal test uses FOF if FORTE risk score ≥ 1%. SNP-based method cutpoint: not reported. Commercial test: Panorama™ prenatal test by Natera, Inc. or Harmony™ prenat nostics, Inc. Target conditions: T21, T18, T13, 45,X, 47,XXX, 47, XXY, 47,XYY. SCA data were no present review. T18 and T13 were also assessed but no case was found. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or neon tion. Flow and timing Blood samples were obtained prior to the invasive procedure (reference standard gNIPT was a first- or second-tier test. 17/333 samples excluded because still pregnant at the time of publication (no f 9/333 samples failed the initial TMPS testing. 6/9 repeated sampling was performed and results were obtained in 5/6. 3/333 samples without gNIPT result and follow-up were excluded (still pregnant at the time of publication (no f 9/333 samples without gNIPT result and follow-up were excluded (still pregnant 2/333 samples without gNIPT result and follow-up were exclud	Study not funded by industry but Ariosa Diagnostics, Inc and Natera, Inc. made sequencing and analy sis.		
Mean gestational age (range): 14.6 (9.5 to 23.5) weeks. Mean maternal age (range): 37 (21 to 46) years. Relevant test carried out prior to index test: biochemical screening for a part of Language of the study: English Index tests gNIPT by TMPS (DANSR assay or SNP-based method). Mean fetal fraction DNA (range): 12.7% (4.2% to 27.9%), Harmony™ prenatal test ma™ prenatal test: 12.7%. Blood samples for gNIPT were collected before reference standard. DANSR assay cutpoint: not reported. Usually, Harmony™ prenatal test uses FOF if FORTE risk score ≥ 1%. SNP-based method cutpoint: not reported. Commercial test: Panorama™ prenatal test by Natera, Inc. or Harmony™ prenat nostics, Inc. Target condition and ref- rerence standard(s) Reference standards: fetal karyotype of chorionic villi or anniotic fluid or neon tion. Flow and timing Blood samples were obtained prior to the invasive procedure (reference standard gNIPT was a first- or second-tier test. 17/333 samples failed the initial TMPS testing. 6/9 repeated sampling was performed and results were obtained in 5/6. 3/333 samples without gNIPT result and follow-up were excluded (still pregnant at follow-up were excluded (still pregnant at the time of publication (no file) result and follow-up were excluded (still pregnant at the time of publication (no file) repeated sampling was performed and results were obtained in 5/6.	nomalies in a clinical set		
Mean gestational age (range): 14.6 (9.5 to 23.5) weeks. Mean maternal age (range): 37 (21 to 46) years. Relevant test carried out prior to index test: biochemical screening for a part of Language of the study: English Index tests gNIPT by TMPS (DANSR assay or SNP-based method). Mean fetal fraction DNA (range): 12.7% (4.2% to 27.9%), Harmony™ prenatal test ma™ prenatal test: 12.7%. Blood samples for gNIPT were collected before reference standard. DANSR assay cutpoint: not reported. Usually, Harmony™ prenatal test uses FOF if FORTE risk score ≥ 1%. SNP-based method cutpoint: not reported. Commercial test: Panorama™ prenatal test by Natera, Inc. or Harmony™ prenat nostics, Inc. Target condition and reference standard(s) Present review. T18 and T13 were also assessed but no case was found. Reference standard(s) Flow and timing Blood samples were obtained prior to the invasive procedure (reference standard gNIPT was a first- or second-tier test. 17/333 samples excluded because still pregnant at the time of publication (no f 9/333 samples failed the initial TMPS testing. 6/9 repeated sampling was performed and results were obtained in 5/6. 3/333 samples without gNIPT result were excluded (unrepeated samples).			
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Mean gestational age (range): 14.6 (9.5 to 23.5) weeks. Mean maternal age (range): 37 (21 to 46) years. Relevant test carried out prior to index test: biochemical screening for a part of			
omas 2015 <i>(Continued)</i> Number available for 2 x 2 table: 312 pregnant women (subgroup of 95%). Setting: 1 private prenatal diagnostics centre in Barcelona, Spain (Hospital Univ Dexeus). Recruitment period: January to December 2013.			



Comas 2015 (Continued) DOMAIN 1: Patient Selection	on		
Was a consecutive or ran- dom sample of patients enrolled?	Yes		
Was a case-control design avoided?	Yes		
Did the study avoid inap- propriate exclusions?	No		
		High	High
DOMAIN 2: Index Test TMP	S		
Were the index test re- sults interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Stan	dard		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timin	Ig		
Was there an appropriate interval between index test and reference stan- dard?	Yes		
Did all analysed patients receive the reference stan- dard?	Yes		
Were all patients included in the analysis?	No		
		High	

del Mar Gil 2014

Study characteristics	
Patient sampling	Study design: retrospective cohort study. Data from prospective cohort were not shown in the present review. Participants: pregnant women without a priori risk who undergo first-trimester screening for trisomies (archived maternal plasma samples). Inclusion criteria: multifetal pregnancies between 11 to 13 weeks' gestation. Exclusion criteria: singleton pregnancies.
Patient characteristics and setting	Number enrolled: 207 pregnant women from the retrospective cohort. Number available for 2 x 2 table: 192 pregnant women (subgroup of 93%).
	Setting: 1 centre at Kings' College Hospital in London, UK. Recruitment period: not reported. Ethnicity: not reported. Median gestational age (range): 13.0 (12.4 to 13.9) weeks. Median maternal age (range): 33.7 (26.7 to 37.9) years. Chorionicity: 41% of pregnancies were monochorionic (85/207) and 59% of pregnancies were di- chorionic (122/207).
	Relevant tests carried out prior to index test: none. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex.
	Mean fetal fraction DNA (range): euploids: 9.8% (7.4% to 12.1%), T21: 10.8% (6.8% to 12.1%), and T13: 7%. Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Usually, Harmony™ prenatal test uses FORTE algorithm; positive if FORTE risk score ≥ 1%. Commercial test: Harmony™ prenatal test by Ariosa Diagnostics, Inc.
Target condition and refer- ence standard(s)	Target conditions: T21 and T13. T18 was also assessed but no case was found. Reference standard: fetal karyotype of chorionic villi or amniotic fluid.
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).
	gNIPT was a first-tier test. 15/207 samples failed during sequencing process (11 for low fetal fraction DNA and 4 for laboratory processing failures) (no gNIPT result).
	No repeated test reported.
Comparative	
Aim to study	To examine the clinical implementation of TMPS of ccfDNA in maternal blood and an algorithm that relies on the lower fetal fraction DNA contribution of the 2 fetuses in the assessment of risk for trisomies in twin pregnancies.
Funding source or sponsor of the study	Study not funded by industry but Ariosa Diagnostics, Inc have performed gNIPT at their own ex- pense. Study funded by a grant from The Fetal Medicine Foundation, UK.
Informations about the au-	Author was contacted on: 27 May and 27 September 2016.
thors contacted	No reply received from the author.
Notes	Data from prospective cohort study were not shown in the present review because patients with gNIPT negative result were without follow-up to confirm gNIPT result.



del Mar Gil 2014 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	Unclear		
		Unclear	Low
DOMAIN 2: Index Test TMPS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard	l		
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	



Ehrich 2011

Study characteristics			
Patient sampling	Study design: blinded, case-control study (1:11) from a prospective cohort. Participants: pregnant women selected from a high-risk population. Inclusion criteria: not reported. Exclusion criteria: not reported.		
Patient characteristics and setting	Number enrolled: 480 pregnant women. Number available for 2 x 2 table: 449 pregnant women (subgroup of 94%). Setting: in clinical practice and pregnancy termination centres. Recruitment period: May 2009 to unknown date. Ethnicity: not reported. Median gestational age (range): 16 (8 to 36) weeks. Mean maternal age (range): 37 (18 to 47) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency mea- surement) and biochemical screening. Language of the study: English.		
Index tests	gNIPT by MPSS on Illumina Genome Analyzer IIx in 4-plex.		
	Minimum fetal fraction DNA as estimated with the fetal quantifier assay: 3.9%. Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if Z score > 2.5. Commercial test: Sequenom's test.		
Target condition and reference stan- dard(s)	Target condition: T21. Reference standard: fetal karyotype of chorionic villi (19%) or amniotic fluid (81%).		
Flow and timing	Blood samples were obtained prior or after the invasive procedure (reference standard).		
	gNIPT was a second-tier test.		
	13/480 samples excluded before sequencing process (9 for plasma volume < 3.5 mL and 4 fo processing errors).		
	20/467 samples failed the initial MPSS testing. 20/20 samples were resequenced using the same library (10 samples in 4-plex and 10 in monoplex) and 2/20 samples obtained a gNIPT results.		
	18/467 samples failed quality control during sequencing process, including 7 samples for low fetal fraction DNA (no gNIPT result).		
Comparative			
Aim to study	To evaluate a multiplexed massively parallel shotgun sequencing assay for noninvasive tri- somy 21 detection using circulating cell-free fetal DNA.		
Funding source or sponsor of the study	Study funded by Sequenom, Inc.		
Informations about the authors con- tacted	Author was been contacted on: 5 May and 28 September 2016. No reply received from the author.		
Notes			
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		



Ehrich 2011 (Continued) DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	No			
Did the study avoid inappropriate ex- clusions?	Unclear			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condi-tion?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		
Fiorentino 2016				
Study characteristics				
Patient sampling St	tudy design: blinded, p	prospective cohort study. Retros	pective cohort (training set) not rep	orted

Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review)

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in the present review.



DOMAIN 1: Patient Selection			
Item	Authors' judgement	Risk of bias	Applicability concerns
Methodological quality			
Notes			
Informations about the authors contacted	Authors were contacted or Reply received on: 6 Septe	: 30 August and 6 September 2 nber.	016.
Funding source or sponsor of the study	Study not funded by indust Italy).	ry but the samples were analys	sed in the GENOMA laboratory (Rome,
Comparative Aim to study	fraction DNA required to de simulate samples at differe	etect common fetal autosomal ent proportions of fetal ccfDNA.	order to define the actual lower fetal trisomies, using a model system to Secondly, to assess the impact of low aternal plasma testing for aneuploi-
Compositive	21/100 unrepeated sample	s failed quality control metrics	(no gNIPT result).
		vith a second blood draw and a	-
Flow and timing	Blood samples for gNIPT w gNIPT was a first- or a seco 100/7103 samples failed th	nd-tier test.	ve procedure (reference standard).
Target condition and reference standard(s)	Target conditions: T21, T18 Reference standards: fetal amination.		mniotic fluid or neonatal clinical ex-
	ploidy) for T21 was determ Blood samples for gNIPT w	ined at 2% fetal fraction level. ere collected before reference 4 (aneuploidy suspected if NCV	
Index tests	gNIPT by MPSS on Illumina	HiSeq 2500 sequencer in 15-pl	ex with SAFeR™ algorithm.
	Mean maternal age (± SD; r	; range): 12.8 (± 2.3; 10 to 30) w ange): 36.4 (± 4.7; 24 to 54) yea prior to index test: ultrasonogra ening or both.	
Patient characteristics and set- ting		gnant women. table: 7082 pregnant women (s	ubgroup of 99.7%).
	Participants: mostly pregn women without prior risk (Inclusion criteria: singletor Exclusion criteria: multifet.	14%). 1 pregnancies.	h-risk population and pregnant

Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review) Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.



Fiorentino 2016 (Continued)			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	No		
		High	High
DOMAIN 2: Index Test MPSS			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	
Gil 2016			

Study characteristics

Patient sampling

Study design: prospective cohort study.

Gil 2016 (Continued)	
	Participants: pregnant women with a first-trimester combined test selected for their risk of fetal aneuploidy (cut-off of 1 in 100 for high risk and 1 in 101 to 1 in 2500 for intermediate risk). Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies, terminations of pregnancy, miscarriages or stillbirths without follow-up.
Patient characteristics and setting	Number enrolled: 11,692 pregnant women. Number available for 2 x 2 table: 3633 pregnant women (subgroup of 31%). Setting: 2 centres. King's College Hospital, London, and Medway Maritime Hospital, Gillingham, Kent in UK. Recruitment period: October 2013 to February 2015. Ethnicity: Caucasian (70%), Afro-Carabbean (20%), Asian (7%) and mixed (3%).
	Gestational age: not reported. Median maternal age (range): 31.6 (25.8 to 39.5) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or both. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay).
	Fetal fraction DNA: not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Usually, Harmony™ prenatal test uses FORTE algorithm; positive if FORTE risk score ≥ 1%. Commercial test: Harmony™ prenatal test by Ariosa Diagnostics, Inc.
	Traditional screening test was also assessed but 2 x 2 tables were incomplete.
Target condition and refer- ence standard(s)	Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi, postnatal karyotype or neonatal clinical ex- amination.
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference standard). gNIPT was a second-tier test.
	7994/11,692 samples did not undergo a gNIPT (no gNIPT result).
	99/3698 samples failed the initial TMPS testing.
	54/99 repeated sampling were processed and 34/54 gNIPT results were obtained.
	65/3698 samples without gNIPT result.
Comparative	
Aim to study	To report the feasibility of implementing gNIPT. To examine the factors affecting patient decisions concerning their options for screening and decisions on the management of affected pregnancies. To report the prenatal diagnosis of fetal trisomies and outcome of affected pregnancies following the introduction of contingent screening.
Funding source or sponsor of the study	Study not funded by industry but the cost of collection and analysis of the blood samples for the cell free DNA test was covered by Ariosa Diagnostics, Inc. These organisations had no role in study design, data collection, data analysis, data interpretation or writing of the report. Study was funded by a grant from The Fetal Medicine Foundation, UK.
Informations about the au- thors contacted	No need for further contact.
Notes	



Gil 2016 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	No		
		High	High
DOMAIN 2: Index Test TMPS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standar	ď		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
		High	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	



Hall 2014

Study characteristics			
Patient sampling	Study design: case-control study (1:3), age-matched randomly selected from a larger cohort. Participants: pregnant women with an affected fetus or considered to be at high risk of fetal aneu- ploidy were recruited.		
	Inclusion criteria: pregnant women at least 18 years of age who had signed an informed consent, and with singleton pregnancy. Exclusion criteria: fetal mosaicism.		
Patient characteristics and setting	Number enrolled: more than 1000 pregnant women. Number available for 2 x 2 table: 64 pregnant women (subgroup of 6%). Setting: 6 centres. Western Institutional (WA, USA), Einstein Institutional (CA and MO, USA), Polish Mother's Memorial Hospital Institutional (Polish), Bio Medical Research Institute of America (CA, USA), and the Mt. Sinai School of Medicine (NY, USA). Recruitment period: March to December of 2012. Ethnicity: not reported. Median gestational age (range): 16.0 (12.1 to 22.7) weeks. Maternal age: not reported. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening. Language of the study: English.		
Index tests	gNIPT by TMPS (SNP-based method) on Illumina Genome Analyzer IIx or HiSeq sequencer. Sam- ples were amplified using 11,000-plex or 19,488-plex targeted polymerase chain reaction (targets included SNPs from chromosomes 13, 18, 21, X, and Y).		
	Mean fetal fraction DNA (median; range): 12.1% (11.1%; 2.2% to 30.4%). Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Commercial test: Natera's prenatal test.		
Target condition and refer- ence standard(s)	Target condition: T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or genetic testing of the cord blood, buccal, saliva or products of conception.		
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).		
	gNIPT was a second-tier test.		
	About 932 samples were not selected for this case-control study. 4/68 samples failed DNA quality threshold for low fetal fraction DNA (no gNIPT result).		
	No repeated test reported.		
Comparative			
Aim to study	To determine how a single nucleotide polymorphism (SNP)- and informatics-based non-invasive prenatal aneuploidy test performs in detecting trisomy 13.		
Funding source or sponsor of the study	Study funded by Natera, Inc. (involved in study design, data collection and analysis, decision to publish, and preparation of the manuscript).		
Informations about the au- thors contacted	Authors were contacted on: 21 April 2016, and 27 May 2016. No reply received from the authors.		
Notes			



Hall 2014 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test TMPS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
		Unclear	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	



Study characteristics	
Patient sampling	Study design: case-control study from archived plasma samples from a prospective cohort. Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive test- ing. Inclusion criteria: pregnant women 18 years and older, with a singleton pregnancy at gestational age 10 weeks or greater, and who were planning to undergo invasive prenatal diagnosis. Exclusion criteria: multifetal pregnancies, pregnant women with a known maternal aneuploidy, active malignancy or a history of metastatic cancer, or those who had already undergone chorionic villus sampling or amniocentesis during the current pregnancy.
Patient characteristics and setting	Number enrolled: not reported. 432 maternal plasma samples were retrieved from the prospective cohort. Number available for 2 x 2 table: 414 samples (subgroup of 96%). Setting: 16 centres. Selected prenatal care centres in the USA, the Netherlands and Sweden. Recruitment period: not reported. Ethnicity: not reported. Mean gestational age (± SD; range): 15.4 (± 3.7; 10 to 34.1) weeks. Mean maternal age (± SD; range): 35.6 (± 5.7; 18.5 to 45.5) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or both. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex. Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Usually, Harmony™ prenatal test uses FORTE algorithm; positive if FORTE risk score ≥ 1%. Commercial test: Harmony™ Prenatal Test by Ariosa Diagnostics, Inc.
Target condition and refer- ence standard(s)	Target conditions: 45,X, 47,XXY and 47,XXX. 47,XYY was also assessed but no case was found. Reference standard: fetal karyotype of chorionic villi or amniotic fluid.
Flow and timing	Blood samples were obtained prior the invasive procedure (reference standard). gNIPT was a second-tier test. 18/432 samples failed during sequencing process (no gNIPT result) for low fetal fraction DNA, un- usually high variation in ccfDNA counts or failure to pass the quality control measures of the DANSR assay. No repeated test reported.
Comparative	
Aim to study	To assess the performance of a directed chromosomal analysis approach in the prenatal evaluation of fetal sex chromosome aneuploidy.
Funding source or sponsor of the study	Study funded by Ariosa Diagnostics, Inc.
Informations about the au- thors contacted	BGI-Shenzhen were contacted on: 19 May 2016. Author was contacted on: 16 June 2016. No reply received from the author.
Notes	



Hooks 2014 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test TMPS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-specified?	Yes		
		Unclear	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Hou 2012

Study characteristics				
Patient sampling	Study design: prospective cohort study. Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive testing. Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies.			
Patient characteristics and setting	Number enrolled: 308 pregnant women. Number available for 2 x 2 table: 205 pregnant women (subgroup of 67%). Setting: 1 centre. Henan Province People's Hospital in China. Recruitment period: October 2010 to January 2012. Ethnicity: Asian. Gestational age range: 14 to 24 weeks. Mean maternal age (range): 31 (21 to 44) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or both. Language of the study: Chinese.			
Index tests	gNIPT by MPSS on Illumina HiSeq 2000 sequencer with BGI's algorithm. Fetal fraction DNA: not reported. Blood samples for gNIPT were collected just before reference standard. Cutpoint: not reported. Commercial test: BGI-Shenzhen's prenatal test.			
Target condition and reference stan- dard(s)	Target conditions: T21, T18, 45,X, 47,XXY and 47,XYY. T13 and 47,XXX were also assessed but no cases were found. Reference standard: fetal karyotype of amniotic fluid.			
Flow and timing	Blood samples were obtained just prior the invasive procedure (reference standard).			
	gNIPT was a second-tier test.			
	103/308 patients did not undergo gNIPT (no gNIPT result). No failed sample reported.			
	No repeated test reported.			
Comparative				
Aim to study	To investigate the clinical	value of gNIPT using ccfDN	NA in maternal blood.	
Funding source or sponsor of the study	Study not funded by indu	stry but BGI-Shenzhen pro	vided the test.	
Informations about the authors contact- ed	Author was contacted on: 11 April 2016 (author) and 19 May 2016 (BGI's contact). No reply received from the author.			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			



Hou 2012 (Continued)				
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclu- sions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-speci- fied?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to cor- rectly classify the target condition?	Yes			
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference standard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Low		

Study characteristics	
Patient sampling	Study design: blinded, prospective cohort study. Participants: pregnant women selected at high risk of fetal aneuploidy (as real clinical samples) Inclusion criteria: twin pregnancies with live fetuses and karyotype result. Exclusion criteria: singleton pregnancies, twins with intrauterine fetal demise at the time of sampling or without fetal karyotype result.
Patient characteristics and set- ting	Number enrolled: 189 pregnant women. Number available for 2 x 2 table: 189 pregnant women (whole cohort included in analyses). Setting: 7 centres. Hospitals in China. Recruitment period: April 2012 to April 2013.

luang 2014 (Continued)				
	Ethnicity: most Asian. Median gestational age (ran Median maternal age (rang Chorionicity: 17% monocho (4/189).	e): 31 (22 to 44) years.	onics (152/189) and 2% unknown	
	Relevant tests carried out p ment) and biochemical scru Language of the study: Eng	eening.	aphy (nuchal translucency measure-	
Index tests	gNIPT by MPSS on Illumina	Genome Analyzer IIx or HiSeq	2000 platform.	
		ere collected 30 minutes befor > 2.5 and L score risk > 1 (warr	re reference standard. ning zone if t score > 2.5 or L score > 1).	
Target condition and reference standard(s)	Target conditions: T21 and T18. Reference standard: fetal karyotype of chorionic villi (2.1%), amniotic fluid (94.2%) or cord blood (3.7%).			
Flow and timing	Blood samples were obtain	ed prior to the invasive proced	dure (reference standard).	
	gNIPT was a second-tier test. No failed sample reported.			
	No repeated test reported.			
Comparative				
Aim to study		of noninvasive prenatal testin rnal plasma in twin pregnancie	g for trisomies 21 and 18 on the basis of es.	
Funding source or sponsor of the study	Funded by the Shenzhen Engineering Laboratory for Clinical Molecular Diagnostic, the China Na- tional GeneBank-Shenzhen, the Medical Centre for Critical Pregnant Women in Guangzhou and Prenatal monitoring, In utero therapy and Follow-up after birth in the complexity of Twin Preg- nancy. Some authors worked for BGI-Shenzhen.			
Informations about the authors	Author was contacted on: 1	0 February 2016.		
contacted	BGI-Shenzhen were contacted on: 19 May 2016. No reply received from author.			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoid- ed?	Yes			
Did the study avoid inappropriate exclusions?	No			



Huang 2014 (Continued)			
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	Yes		
		Low	
Jackson 2014			

Patient sampling	Study design: prospective cohort study. Participants: pregnant women selected at high risk and low risk of fetal aneuploidy present- ing for screening. Inclusion criteria: not reported. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 1228 pregnant women screened at first-trimester, including 1184 pregnan women with normal first-trimester ultrasound and 44 with abnormal ultrasound. Number available for 2 x 2 table: 1161 pregnant women (subgroup of 95%). Setting: 1 centre. South Shore Hospital in USA.
	Recruitment period: June 2012 to January 2013.

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ackson 2014 (Continued)				
	Ethnicity: not reported. Gestational age: not report Median maternal age: 31.5 Relevant tests carried out p surement) or biochemical Language of the study: Eng	years. orior to index test: ultrasonc screening or both.	graphy (nuchal translucency mea-	
Index tests	gNIPT by TMPS (DANSR ass	say).		
		rere collected before referen ually, Harmony™ prenatal te	ice standard. est uses FORTE algorithm; positive if	
Target condition and reference stan- dard(s)	Target conditions: T21, T18 Reference standards: fetal from birth.		or amniotic fluid, or medical record	
Flow and timing	Blood samples were obtair	ned prior to the invasive pro	cedure (reference standard).	
	sound, 14 women opted fo	of 2 x 2 tables, including 7 v	vomen with other abnormal ultra- 2 women declined all testing and 14 cess (no gNIPT result).	
Comparative				
Aim to study			asurement followed by gNIPT in the r-based average-risk population.	
Funding source or sponsor of the study	Funding source not reporte	ed but 1 author is employed	by Ariosa Diagnostics, Inc.	
Informations about the authors con- tacted	Author was contacted on: 22 February 2016 and 15 March 2016. No reply received from the author.			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Unclear			
Did the study avoid inappropriate ex- clusions?	Unclear			
		Unclear	High	

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Jackson 2014 (Continued)				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condi-tion?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	No			

Jeon 2014

Patient sampling	Study design: prospective cohort study.
	Participants: pregnant women selected at high risk of fetal aneuploidy presenting for inva- sive testing.
	Inclusion criteria: women who gave written informed consent participated in the study if they were ≥ 19 years old and had a singleton pregnancy with a gestational age of at least 12 weeks.
	Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 155 pregnant women.
	Number available for 2 x 2 table: 155 pregnant women (whole cohort included in analyses). Setting: 1 centre. Xiamen Maternal & Child Health Care Hospital, Xiamen, Fujian, China. Recruitment period: March 2012 to October 2013. Ethnicity: Asian.
	Gestational age ranges: 12 to 16 weeks (18.1%), 17 to 21 weeks (55.5%), ≥ 22 weeks (26.5%) All between 12 to 24 weeks.
	Mean maternal age (± SD; range): 30.73 (± 4.99; 19 to 43) years.



Jeon 2014 (Continued)	Relevant tests carried out p surement) or biochemical s Language of the study: Engl	creening or both.	graphy (nuchal translucency mea-	
Index tests	gNIPT by MPSS on Ion Torre	nt PGM sequencer with 10	samples per chip.	
	Fetal fraction DNA: not repo Blood samples for gNIPT we Cutpoint: positive if Z score Commercial test: Genome C	ere collected just before ref > 2.566 (T21) or > 2.459 (T1		
Target condition and reference stan- dard(s)	Target conditions: T21 and Reference standard: fetal ka			
Flow and timing	Blood samples were obtain	ed prior to the invasive pro	cedure (reference standard).	
	gNIPT was a second-tier tes No failed sample reported.	t.		
	No repeated test reported.			
Comparative				
Aim to study	To investigated whether fet semiconductor sequencer:		vely and specifically detectable by	
Funding source or sponsor of the study		ext generation bioinformation	evelopment Program, "Bioinformatics tion analysis" funded by the Ministry	
Informations about the authors con- tacted	Author was contacted on: 6 and 11 April 2016. Reply received on: 11 April 2016.			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate ex- clusions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			

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Jeon 2014 (Continued)

If a threshold was used, was it prespecified?

		High	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condi- tion?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Low		

Jiang 2012

Patient sampling	Study design: prospective cohort study. Participants: pregnant women at high risk of fetal aneuploidy presenting for invasive test- ing selected from the cohort. Inclusion criteria: not reported. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 903 pregnant women. Number available for 2 x 2 table: 903 pregnant women (whole cohort included in analyses Setting: 3 centres in Shenzen, China. Recruitment period: June 2009 to August 2010. Ethnicity: Asian. Gestational age range: 10 to 39 weeks. Maternal age range: 20 to 45 years. Relevant tests carried out prior to index test: not reported. Language of the study: English.
Index tests	gNIPT by MPSS on platforms Illumina Genome Analyzer IIx or Illumina HiSeq 2000 by mult plex sequencing. Fetal fraction DNA (range): quality control criteria > 3.5% (1% to 33%).



Jiang 2012 (Continued)	It is not reported if the blood standard. Cutpoint:	d samples for gNIPT were	collected before or after reference
	1) Positive if binary hypothe 3 and if logarithmic LR > 1 (a		s) > 3 and t score (second hypothesis) <
	2) Positive if t score < -2.5 (4	5,X and 47,XXX) without C	hrom. Y representation.
	3) Positive if t score > 2.5 con X and Y independently (47,X Commercial test: NIFTY™ pr	XY and 47,XYY) for male fe	
Target condition and reference stan- dard(s)	Target conditions: T21, T18, Reference standard: fetal ka		
Flow and timing	It is not reported if the blood (reference standard).	d samples were obtained	prior or after the invasive procedure
	gNIPT was a second-tier test No failed sample reported.	t.	
	No repeated test reported.		
Comparative			
Aim to study	To develop an advanced gN	IPT method based on MPS	SS.
Funding source or sponsor of the study	Study funded by industry. B sis and interpretation of res		dy design, conduct of the study, analy-
Informations about the authors con- tacted	Author was contacted on: 19 No reply received from the a		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate ex- clusions?	Unclear		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		



Jiang 2012 (Continued)

If a threshold was used, was it prespecified?

		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi-tion?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	Yes		
		Unclear	

Johansen 2016

Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: pregnant women selected at high risk of fetal aneuploidy presenting for inva- sive testing. Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 375 pregnant women (184 for the validation set). Number available for 2 x 2 table: 173 pregnant women (subgroup of 94%). Setting: Danish public health setting. Recruitment period: not reported. Ethnicity: not reported. Median gestational age (range): 13.4 (10.6 to 31) weeks. Maternal age: not reported. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency mea- surement) or biochemical screening or both. Language of the study: English.
Index tests	gNIPT by MPSS on Ion Proton™ sequencer in 5-plex. Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected just before reference standard.



Johansen 2016 (Continued)			
	Cutpoint: positive if Z score ≥ and 4). In-house test.	4 and WISECONDOR≥1% (un	classified if Z score between 3
Target condition and reference stan- dard(s)	Target conditions: T21, T18 a Reference standard: fetal kar	nd T13. yotype of chorionic villi or am	niotic fluid.
Flow and timing	Blood samples for gNIPT were dard). gNIPT was a second-tier test.	e obtained just prior the invas	ive procedure (reference stan-
		s for the validation set were ex sequencing process for low fe	xcluded. tal fraction DNA (no gNIPT re-
	2/173 samples were resequer 2 results were obtained.	nced because gNIPT results we	ere in the inconclusive zone and
Comparative			
Aim to study	To introduce gNIPT for fetal a ting, using semi-conductor se		er in a Danish public health set- n source scripts for analysis.
Funding source or sponsor of the study	No funding source was report	ed.	
Informations about the authors con- tacted	No need for further contact.		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate ex- clusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			



Johansen 2016 (Continued)				
Is the reference standards likely to cor- rectly classify the target condition?	Yes			
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analy- sis?	No			
		High		

Ke 2015

Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: pregnant women considered at high risk of fetal aneuploidy. Inclusion criteria: singleton pregnancies. Pregnant women at high risk of fetal aneuploidy de- scribe as follows: over age 35, the histories of abnormal pregnancy including children with T21 and repeated spontaneous abortion, stillbirth in pregnancy periods, abnormal serologica screening for T21 at early and mid pregnancy, abnormal screening for fetal nuchal translucen- cy using colour duplex ultrasonography between 11-14 weeks of gestation. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 2340 pregnant women. Number available for 2 x 2 table: 2340 pregnant women (whole cohort included in analyses). Setting: 1 centre. Clinical setting at Shenzhen Second People's Hospital in China. Recruitment period: March 2012 to May 2013. Ethnicity: Asian. Gestational age: positive cases were between 16 to 24 weeks. All cohort: 95% were between 15 to 20 weeks, 3% were between 12 to 14 weeks and 0.9% were ≥ 24 weeks. Maternal age: 88% were less than 35 years old and 12% were 35 years old or more. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measure- ment) or biochemical screening or both. Language of the study: English.
Index tests	gNIPT by MPSS. Fetal fraction DNA: not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if t score > 3. Commercial test: BGI-Shenzhen's prenatal test.



Ke 2015 (Continued)				
Target condition and reference standard(s)	Target conditions: T21, T18, and T13. Reference standards: fetal karyotype or newborn outcome.			
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).			
	gNIPT was a second-tier test. No failed sample reported.			
	No repeated test reported.			
Comparative				
Aim to study	To validate the efficacy of c and 13 in a clinical setting.	letection of fetal cell-free	DNA in maternal plasma of trisomy 21, 18	
Funding source or sponsor of the study		Study not funded by industry but patients had obtained insurance plans on behalf of Shenzhen Huada Genomics Institute.		
Informations about the authors contacted		Author was contacted on: 22 April 2016. No reply received from the author.		
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sam- ple of patients enrolled?	No			
Was a case-control design avoid- ed?	Yes			
Did the study avoid inappropriate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condi- tion?	Yes			



Ke 2015 (Continued)

Were the reference standard results interpreted without knowledge of the results of the index tests?

Unclear

Low

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?	Yes
Did all analysed patients receive the reference standard?	Yes
Were all patients included in the analysis?	Yes

Low

Kim 2016

Study characteristics	
Patient sampling	Study design: blinded, prospective cohort study. Participants: pregnant women selected at high risk of fetal aneuploidy presenting for in- vasive testing. Inclusion criteria: not reported. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 101 pregnant women. Number available for 2 x 2 table: 101 pregnant women (whole cohort included in analy- ses). Setting: 3 centres (Mirae & Heemang, Namujungwon and GN hospitals) in Korea. Recruitment period: December 2014 to April 2015. Ethnicity: Asian. Gestational age range: 11 to 18 weeks. Mean maternal age (± SD; range): 35.45 (± 3.64; 25 to 42) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency mea- surement) and biochemical screening (quadruple test screening). Language of the study: English.
Index tests	gNIPT by MPSS on Ion Torrent PGM (data not shown in the present review) and Ion Pro- ton [™] sequencer in multiplex. Fetal fraction DNA: not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if Z score > 2.10 for Ion Proton [™] . Commercial test: Genome Care's prenatal test.
Target condition and reference stan- dard(s)	Target condition: T21. Reference standard: fetal karyotype of amniotic fluid.
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference stan- dard).



Kim 2016 (Continued)				
	gNIPT was a second-tier te	st.		
	No failed sample reported.			
	No repeated test reported.			
Comparative				
Aim to study	To compare the Ion Torren ly using PGM and Ion Proto		atforms for gNIPT for fetal T21 direct e same set of samples.	
Funding source or sponsor of the study	Study funded by Genome (Genome Care.	Care internal research fun	ding. The first author is employee of	
Informations about the authors contact- ed	No need for further contact.			
Notes	Data from PGM sequencer	Data from PGM sequencer are not shown in the present review to avoid patients overlap.		
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclu- sions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-spec- ified?	No			
		High	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to cor- rectly classify the target condition?	Yes			
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				



Kim 2016 (Continued)	
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes
Did all analysed patients receive the reference standard?	Yes
Were all patients included in the analy- sis?	Yes
	Low

Korostelev 2014

Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: pregnant women selected from a population at high risk or without prior risk of fetal aneuploidy. Inclusion criteria: women who had a singleton pregnancy and more than 9 weeks of gestation. Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 1968 pregnant women. Number available for 2 x 2 table: 685 pregnant women (subgroup of 35%). Setting: private clinics in Moscow, Russia. Recruitment period: 2012 to 2014. Ethnicity: not reported. Median gestational age (range): 14 (9 to 33) weeks. Mean maternal age (range): 34.4 (26 to 45) years. Relevant tests carried out prior to index test: biochemical screening or ultrasonography (nuchal translucency measurement) or both. Language of the study: English.
Index tests	gNIPT by TMPS (SNP-based method) on Illumina Genome Analyzer IIx or HiSeq sequencers with NATUS algorithm. Fetal fraction DNA: not reported (usually NATERA used quality control criteria > 4%). Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Commercial test: Natera's prenatal test.
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. 45,X, 47,XXY, 47,XYY and 47,XXX were also screened but inappropriate reference standard for the present review was used (data not shown in this re- view). Reference standards: fetal karyotype of chorionic villi or amniotic fluid or medical record from birth.
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference standard). gNIPT was a second-tier test. 240/1968 samples did not undergo gNIPT (no gNIPT result). 1043/1728 samples without follow-up were excluded. No repeated test reported.
Comparative	



orenatal diagnostics and th irmation by karyotyping of Study not funded by indust Author was contacted on: 2	neir advantages between cor f CVS or amniocytes. try but gNIPT was carried our 21 June 2016.	d chromosomal microarray analysis for mbined first-trimester screen with con- t by Natera, Inc.		
Author was contacted on: 2	21 June 2016.	t by Natera, Inc.		
		Author was contacted on: 21 June 2016. No reply received from the author.		
Authors' judgement	Risk of bias	Applicability concerns		
No				
/es				
No				
	High	High		
/es				
/es				
	Low	Low		
/es				
Jnclear				
	Unclear	Low		
	No /es No /es /es /es /es /es	No /es /vo /es /es /es /es /uo		



Korostelev 2014 (Continued)

Was there an appropriate interval between index test and reference standard?	Yes	
Did all analysed patients receive the reference standard?	Yes	
Were all patients included in the analysis?	Yes	
		Low

Lau 2012

Study characteristics			
Patient sampling	Study design: blinded, prospective cohort study.		
	Participants: pregnant women mostly at high risk of fetal aneuploidy presenting for invasive testing.		
	Inclusion criteria: not reported.		
	Exclusion criteria: not reported.		
Patient characteristics and setting	Number enrolled: 108 pregnant women.		
	Number available for 2 x 2 table: 108 pregnant women (whole cohort included in analyses).		
	Setting: 1 centre in Japan.		
	Recruitment period: not reported.		
	Ethnicity: Asian.		
	Median gestational age (range): 12.7 (11.6 to 28) weeks, 89.8% < 14 weeks.		
	Mean maternal age (± SD): 37 (± 4.3) years.		
	Relevant tests carried out prior to index test: ultrasonography (nuchal translucency mea- surement) and biochemical screening.		
	Language of the study: English.		
Index tests	gNIPT by MPSS on Illumina HiSeq 2000 sequencer in 12-plex.		
	Fetal fraction DNA: not reported.		
	Blood samples for gNIPT were collected immediately before reference standard.		
	Cutpoint:		
	1) positive if Z score ≥ 3 (T21, T18 and T13).		
	2) for female fetus, positive if Chrom. X Z score ≤ -3 (45,X).		
	3) for female fetus, positive if Chrom. X Z score ≥ 3 (47,XXX).		
	4) for male fetus, positive if Chrom. Y Z score ≥ 3 (47,XXY).		

Lau 2012 (Continued)	Commercial test: NIFTY™ pren	atal test by BGI-Shenzhen.		
Target condition and reference stan- dard(s)	Target conditions: T21, T18, T no case was found.	13, 45,X and 47,XXY. 47,XYY and	47,XXX were also assessed but	
	Reference standard: fetal karyotype of chorionic villi or amniotic fluid.			
Flow and timing	Blood samples for gNIPT were collected immediately before invasive procedure (reference standard).			
	gNIPT was a second-tier test.			
	No failed sample reported.			
	No repeated test reported.			
Comparative				
Aim to study	To investigate the inclusion of an internal reference in the noninvasive prenatal identifica- tion of common fetal aneuploidies using massively parallel sequencing on maternal plasma.			
Funding source or sponsor of the study	Study funded by BGI-Shenzhe	n.		
Informations about the authors con-	BGI-Shenzhen contacted on: 19 May 2016.			
tacted	No reply received from the author.			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate ex- clusions?	Unclear			
		High	High	
DOMAIN 2: Index Test MPSS				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				



Lau 2012 (Continued)				
Is the reference standards likely to correctly classify the target condi-tion?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Low		

Lee 2015

Patient sampling	Study design: blinded, prospective cohort study.
	Participants: pregnant women selected at high risk of fetal aneuploidy presenting for inva- sive testing.
	Inclusion criteria: pregnant women who were > 18 years old and gestational age > 8 weeks, multifetal and singleton pregnancies. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 93 pregnant women. Number available for 2 x 2 table: 92 pregnant women (subgroup of 99%).
	Setting: 1 centre at Asan Medical Centre, Seoul, Korea.
	Recruitment period: August 2014 to February 2015.
	Ethnicity: Asian.
	Median gestational age (range): 21.1 (8.2 to 31.1) weeks.
	Median maternal age (range): 32 (21 to 43) years.
	Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measure ment) or biochemical screening or both.
	Language of the study: English.
Index tests	gNIPT by MPSS on Illumina MiSeq sequencer in 12-plex or on NextSeq 500 sequencer in 96- plex.
	Median fetal fraction DNA (range): male fetus only: 10.2% (3.85% to 25.0%).
	Blood samples for gNIPT were collected before reference standard. Cutpoint:
	1) positive if Z score > 4 (intermediate risk if Z score between 2.5 and 4) for T21 and T18.
	2) positive if Z score > 2.8 (intermediate risk if Z score between 1.9 and 2.8) for T13.



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Lee 2015 (Continued)	Commercial test: MomGuard™ by LabGenomics.			
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. SCA were also assessed but no case was found. Reference standards: fetal karyotype of chorionic villi, amniotic fluid, cord blood or products of conception or neonatal karyotype from peripheral blood.			
Flow and timing	Blood samples for gNIPT were obtained just prior to the invasive procedure (reference stan- dard). gNIPT was a second-tier test. 1/93 samples failed during sequencing process for low fetal fraction DNA (no gNIPT result).			
Comparative	No repeated test reported.			
Aim to study	To evaluate the performance abnormalities recently develo		PT, for detecting T21, T18, T13, and SCA	
Funding source or sponsor of the study	Study funded by a grant from	the LabGenomics Clin	ical Research Institute.	
Informations about the authors con- tacted	No need for further contact.			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sam- ple of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Yes			



Lee 2015 (Continued)

Were the reference standard results Yes interpreted without knowledge of the results of the index tests?

		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Lefkowitz 2016

Study characteristics	
Patient sampling	Study design: Retrospective cohort, blinded case-control study. Participants: pregnant women selected at high risk of fetal aneuploidy from 4 cohorts (archived ma ternal plasma samples). Inclusion criteria: not reported. Exclusion criteria: cases of fetal mosaicism or incomplete karyotype or microarray information.
Patient characteristics and setting	Number enrolled: 5321 pregnant women in all 4 cohorts. 1222 pregnant women selected for this study. Number available for 2 x 2 table: 1166 pregnant women (subgroup of 95%) for autosomes and 1144 pregnant women (subgroup of 94%) for SCA. Setting: multicentre. Recruitment period: not reported. Ethnicity: not reported. Median gestational age (range): 17 (8 to 38) weeks. Median maternal age (range): 36.0 (17.8 to 47) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or both. Language of the study: English.
Index tests	 gNIPT by MPSS on Illumina HiSeq 2000 in 6-plex or uniplex. Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected before (for 1189 pregnant women) or after (for 24 pregnant women) reference standard. Cutpoint: 1) positive for T21 if Z score ≥ 3. 2) positive for T18 or T13 if Z score ≥ 3.95. 3) positive for 45,X if Z score < -3.5 (non-reportable regions between -2.5 and -3.5). 4) positive for 47,XXX if Z score > 3.5 (non-reportable regions between 2.5 and 3.5).

_efkowitz 2016 (Continued)			
	5) positive for 47,XYY if Z score <	-3.5 with Chrom. Y repre	sentation.
	6) positive for 47,XXY if Z score i	s between -3.5 and 3.5 wi	th Chrom. Y representation.
	Commercial test: Sequenom's t	est.	
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T1 also assessed but data not show Reference standard: fetal karyo	vn in the present review.	l 47,XXX. copy number variants ≥ 7 Mb were Imniotic fluid.
Flow and timing	Blood samples for gNIPT were of gNIPT was a second-tier test.	btained prior or after the	invasive procedure (reference standard).
	4099/5321 samples not selected 14/1222 samples were excluded diagnostic information and 3/1	before sequencing proc	ess (11/14 samples excluded for incomplete onfirmed mosaicism).
		NA, 29/42 failed samples	cess (no gNIPT result) including 11/42 failed for technical reasons and 2/42 failed sam-
	22/1166 samples failed SCA sec	uencing process (no gNIF	PT result).
	No repeated test reported.		
Comparative			
Aim to study	To provide a clinical validation genome wide abnormalities.	of the sensitivity and spec	cificity of a novel NIPT for detection of
Funding source or sponsor of the study	Study funded by Sequenom, In	C.	
Informations about the au- thors contacted	No need for further contact.		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge	Yes		



Lefkowitz 2016 (Continued) of the results of the reference standard?				
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standa	rd			
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate in- terval between index test and reference standard?	Yes			
Did all analysed patients re- ceive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		

Liang 2013

Study characteristics	
Patient sampling	Study design: blinded, prospective cohort study. Participants: pregnant women considered at high risk for fetal T21. Inclusion criteria: singleton and twin pregnancies underwent conventional serum screening and ul- trasound scanning, and who invasive prenatal diagnostics were offered. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 435 pregnant women. Number available for 2 x 2 table: 412 pregnant women (subgroup of 94.7%). Setting: 3 hospitals in China. Recruitment period: March 2009 to June 2011. Ethnicity: Asian. Median gestational age (range): 21.4 (11.4 to 39.4) weeks. Most pregnant women (60%) are between 21 to 40 weeks. Only 1 case is in the first trimester (0.23%). Mean maternal age (± SD): 31 (± 5.9) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or multiple screening tests. Language of the study: English.



iang 2013 (Continued)					
Index tests	gNIPT by MPSS on Illumina HiSeq 2000 in 8-plex or 12-plex.				
	Fetal fraction DNA: for a Z score cutoff value of 3 for chromosome 21, fetal DNA was estimated to 5.52%. Blood samples for gNIPT were collected before reference standard. Cutpoint:				
	1) positive if Z score > 3 (T21).				
	2) positive if Z score > 5.91 (T18).				
	3) positive if Z score > 5.72 (T13).				
	4) positive if Z score Chrom. X < -2.91 and Z score Chrom. Y < 3 (45,X).				
	5) positive if Z score Chrom. X range from -2.91 to +2.91 and Z score Chrom. Y > 3 (47,XXY).				
	6) positive if Z score Chrom. X > 2.91 and Z score Chrom. Y < 3 (47,XXX).				
	7) positive if Z score Chrom. X < -2.91 and Z score Chrom. Y > 3 (47,XYY). Commercial test: Berry Genomics's prenatal test.				
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13, 45,X, 47,XXY, 47,XYY, and 47,XXX. Reference standard: fetal karyotype of chorionic villi (1%) or amniotic fluid (77%) or cord blood (22%).				
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).				
	gNIPT was a second-tier test. 12/435 samples failed sequencing process quality control (no gNIPT result).				
	11/423 samples without karyotype were excluded (no reference standard result).				
Comparative					
Aim to study	To determine whether gNIPT by maternal plasma DNA sequencing can uncover all fetal chromosome aneuploidies in 1 simple sequencing event.				
Funding source or sponsor of the study	Study not funded by industry but Berry Genomics Co. Ltd performed the sequencing analysis for free. This study was supported by the grants from the National High Technology Research and Development Program of China (863 Program) (No.2011AA02A112), the National Key Basic Research Program of China (2012CB944600) and the National Key Technology R&D Program of China (2012BAI09B05).				
Informations about the au- thors contacted	No need for further contact.				
Notes					
Methodological quality					
Item	Authors' judgement Risk of bias Applicability concerns				
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	No				
Was a case-control design	Yes				



Liang 2013 (Continued)

Did the study avoid inappro-Yes priate exclusions?

		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standa	rd		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Liu	201	2
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Study characteristics

Patient samplingStudy design: prospective cohort study.
Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive
testing.
Inclusion criteria: women who planned an invasive testing for 1 or more of the following rea-
sons: abnormality in plasma test, older than 35 years old, infant deformity (ultrasound), taken
drugs (teratogen) during early pregnancy or history of malformation caused by virus infection,
history of birth defect caused by abnormal chromosome, history of fetus stopping growth or re-



Liu 2012 (Continued)		ily or either of the couple, to	h for unknown reason, history of chro- oo much or little amniotic fluid.
Patient characteristics and set- ting	Number enrolled: 153 pregn Number available for 2 x 2 ta		vhole cohort included in analyses).
	Setting: Henan Province Pec Recruitment period: Octobe Ethnicity: Asian. Gestational age: more than 1 Mean maternal age (± SD; ra Relevant tests carried out pr ment) or biochemical screer Language of the study: Chine	r to November 2011. 14 weeks. nge): 32.3 (± 1.2; 20 to 44) ye ior to index test: ultrasonog ing or both.	ars. raphy (nuchal translucency measure-
Index tests	gNIPT by MPSS on Illumina H	liSeq sequencer in multiple:	Χ.
	Fetal fraction DNA: not repo Blood samples for gNIPT we Cutpoint: positive if Z score : It is not reported if gNIPT wa	re collected 30 minutes befc ≥ 3.	
Target condition and reference standard(s)	Target conditions: T21, T18, case were found. Reference standard: fetal ka		and 47,XXX were also assess but no
Flow and timing	standard). gNIPT was a second-tier test		r to the invasive procedure (reference
	No failed sample reported. No repeated test reported.		
Comparative			
Aim to study	To determine the feasibility high-flux sequencing analys		umerical chromosomal abnormalities by plasma.
Funding source or sponsor of the study	Study funded by by the Nalio Technology Research Projec		ation of China and a Medical Science and
Informations about the authors contacted	Author was contacted on 11	April 2016 but contact autho	or's email is no longer valid.
Notes			
Methodological quality			
ltem	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoid- ed?	Yes		



Liu 2012 (Continued)

Did the study avoid inappropriate Unclear exclusions?

		Unclear	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards like- ly to correctly classify the target condition?	Yes			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Low		

Study characteristics	
Patient sampling	Study design: blinded, prospective and retrospective cohort study. Participants: pregnant women selected from a high-risk population presenting for invasive testing (prospective cohort) and archived maternal plasma from mixed-risk (high and low risk of fetal aneu ploidy) pregnant women (retrospective cohort). Inclusion criteria: singleton pregnancies with gestational age of 12 weeks or above at the time of sampling.
	Exclusion criteria: women with twin pregnancy or organ donation history or maternal chromosome abnormality.



Ma 2016 (Continued)			
Patient characteristics and setting	cohort. Number available for 2 x 2 tak Setting: 20 centres. Prenatal o Recruitment period: January (prospective). Ethnicity: Asian. Median gestational age: 19 wo Median maternal age (range):	ole: 10,579 pregnant women (so diagnosis clinics in China. 2012 to January 2014 (retrospo eeks. 32 (16 to 53) years. or to index test: ultrasonograph h.	ective cohort and 8159 from retrospective ubgroup of 99.8%). ective) and February to May 2014 ny (nuchal translucency measurement) or
Index tests	gNIPT by MPSS on BGISEQ-10	00 platform in 16 or 24-plex.	
	Fetal fraction DNA: not report Blood samples for gNIPT were Cutpoint: positive if Z score > Commercial test: BGI-Shenzh	e collected before reference sta 3.	andard.
Target condition and refer- ence standard(s)	Target conditions: T21, T18 a Reference standards: fetal ka low-up.		otic fluid or cord blood, or postnatal fol-
Flow and timing	Blood samples were obtained	l prior to the invasive procedu	re (reference standard).
	had incomplete clinical inform	ided from the analysis includir nation and 1 sample failed qua	g 5 from retrospective cohort (4 samples llity control during sequencing) and 14 ll information and 4 samples failed quality
	No repeated test reported.		
Comparative			
Aim to study	throughout gNIPT method ba	sed on combinatorial probe-a	inical performance of a new ultrahigh nchor ligation sequencing (cPAL) of ccfD- s using a centralised testing mode.
Funding source or sponsor of the study	Some authors are employees of BGI-Shenzhen, BGI-Manufacture or BGI-DX. Study funded by Shen- zhen Birth Defect Screening Project Lab, Key Laboratory of Cooperation Project in Guangdong Province, Shenzhen Municipal Government of China, Pilot projects of regional strategic emerging in- dustry cluster development by Hubei provincial development and Reform Commission and Action plan for the development of high-tech industry in biotechnology and new medicine in 2012 by Wuhan Science and Technology Bureau.		
Informations about the au- thors contacted	No need for further contact.		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection	n		



Ma 2016 (Continued)				
Was a consecutive or ran- dom sample of patients en- rolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inap- propriate exclusions?	No			
		High	High	
DOMAIN 2: Index Test MPSS				
Were the index test results interpreted without knowl- edge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Stand	ard			
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all analysed patients receive the reference stan- dard?	Yes			
Were all patients included in the analysis?	No			
		High		

Mazloom 2013

Study characteristics

Mazloom 2013 (Continued)	Chudu dasian blindad museu estive se baut study (validation est)		
Patient sampling	Study design: blinded, prospective cohort study (validation set). Participants: pregnant women selected at high risk of fetal aneuploidy. Inclusion criteria: ≥ 18 years old and singleton pregnancies between 10.5 and 20 weeks of gesta- tion.		
	Exclusion criteria: multifetal pregnancies, mosaic cases for sex chromosomes, or samples without documented karyotype report available.		
Patient characteristics and setting	Number enrolled: 1975 pregnant women including 1564 in the training set (data not shown in the present review) and 411 in the validation set. Number available for 2 x 2 table: 411 pregnant women (subgroup of 95% of validation set). Setting: not reported.		
	Recruitment period: not reported. Ethnicity: Caucasian (58.4%), Asian (18.5%), Afro-American (7.5%), other and not specified (15.6%) Median gestational age (range): 17 (8 to 29) weeks. Median maternal age (range): 36 (19 to 47) years.		
	Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening. Language of the study: English.		
Index tests	gNIPT by MPSS on Illumina v3 flow cells on HiSeq 2000 sequencer in 12-plex.		
	Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint:		
	1) positive for 45,X if Z score < -3.5 (non-reportable regions between -2.5 and -3.5).		
	2) positive for 47,XXX if Z score > 3.5 (non-reportable regions between 2.5 and 3.5).		
	3) positive for 47,XYY if Z score < -3.5 with Chrom. Y representation.		
	4) positive for 47,XXY if Z score is between -3.5 and 3.5 with Chrom. Y representation.		
	Commercial test: Sequenom's prenatal test.		
Target condition and refer- ence standard(s)	Target conditions: 45,X, 47,XXY, 47,XYY and 47,XXX. Reference standard: fetal karyotype of chorionic villi or amniotic fluid.		
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).		
	gNIPT was a second-tier test.		
	1564/1975 excluded samples were used for the training set.		
	21/411 failed samples were in the non reportable region and were considered positive gNIPT result by authors.		
	No repeated test reported.		
Comparative			
Aim to study	To extend the detection of autosomal aneuploidies by MPSS of ccfDNA from maternal plasma clude common sex chromosome aneuploidies.		
Funding source or sponsor of the study	Study funded by Sequenom, Inc. and Sequenom Center for Molecular Medicine (SCMM).		
Informations about the au- thors contacted	Author was contacted on: 26 May 2016. No reply received from the author.		
Notes	Data from the training set were not shown in the present review.		



Mazloom 2013 (Continued)

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard	l		
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	



Study characteristics	
Patient sampling	Study design: retrospective study from a prospective cohort. Participants: selected archived plasma samples from pregnant women without prior risk of fetal aneuploidy (general population) attending for their routine first-trimester combined screening for aneuploidies. Inclusion criteria: singleton pregnancies between 11 to 13.9 weeks' gestation. Archived samples of at least 2 mL. Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 2230 pregnant women. Number available for 2 x 2 table: 1949 pregnant women (subgroup of 87%). Setting: not reported. Recruitment period: October 2010 to January 2011. Ethnicity: Caucasian (69.8%), African (20.6%), South Asian (4%), East Asian (2.8%) and mixed (2.8%). Gestational age range: 11 to 13.9 weeks. Median maternal age (range): 31.8 (27.7 to 35.4) years. Relevant tests carried out prior to index test: none. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay). Median fetal fraction DNA (interquartile range): euploids: 10.0% (7.8% to 13.0%), T21: 12.5% (9.2% to 21.3%), and T18: 9.3% (5.6% to 13.0%). Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if FORTE algorithm risk score ≥ 1%. Commercial test: Harmony [™] Prenatal test by Ariosa Diagnostics, Inc.
	The traditional screening test (combined test at the first trimester) was also assessed. Cutpoint of combined test: 1 in 150.
Target condition and reference stan- dard(s)	Target conditions: T21 and T18. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or neonatal clinical examination.
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard). gNIPT was a first-tier test. 181/2230 samples were ineligible (no fetal karyotype or follow-up, miscarriage, stillbirth, termination of pregnancy or other abnormalities). 100/2049 samples failed during sequencing process including 46 for low fetal DNA and 54 had assay failures (no gNIPT result).
	No repeated test reported.
Comparative	
Aim to study	To assess performance of noninvasive prenatal testing for fetal trisomy in a routinely screened first-trimester pregnancy population.
Funding source or sponsor of the study	The study was supported by a grant from the Fetal Medicine Foundation (UK). The cost of collection and analysis of the samples was covered by Ariosa Diagnostics, Inc.
Informations about the authors con- tacted	No need for further contact.



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Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate ex- clusions?	No		
		High	Low
DOMAIN 2: Index Test TMPS			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 2: Index Test Traditional scr	eening tests		
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi-tion?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			

Was there an appropriate interval be-Yes tween index test and reference standard?

Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review) Copyright @ 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

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No

Nicolaides 2012 (Continued)

Did all analysed patients receive the	Yes
reference standard?	

Were all patients included in the analysis?

High

Nicolaides 2013

Study characteristics	
Patient sampling	Study design: blinded, prospective cohort study. Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive testing. Inclusion criteria: singleton pregnancies at high risk of fetal aneuploidy between 11 to 13 weeks gestation. Exclusion criteria: multifetal pregnancies.
Patient characteristics and set- ting	Number enrolled: 242 pregnant women. Number available for 2 x 2 table: 229 pregnant women (subgroup of 95%). Setting: 1 centre. Fetal Medicine Centre, in UK. Recruitment period: not reported. Ethnicity: not reported. Median gestational age (range): 13.1 (11.3 to 13.9) weeks. Median maternal age (range): 35.7 (18.5 to 46.5) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measure- ment) and biochemical screening. Language of the study: English.
Index tests	gNIPT by TMPS (SNP-based method) on Illumina Genome Analyzer IIx or HiSeq sequencers with NATUS algorithm. Fetal fraction DNA: the lowest fetal fraction DNA on a case that returned a result was 3.95%. Blood samples for gNIPT were collected immediately before reference standard. Cutpoint: not reported. Commercial test: Natera's prenatal test.
Target condition and reference standard(s)	Target conditions: T21, T18, 45,X. 47,XXY, 47,XYY and 47,XXX were also assessed but no case was found. T13 was also assessed but the only 1 case presented in this publication was published thereafter in Hall 2014. T13 case was excluded to avoid double counting. Reference standard: fetal karyotype of chorionic villi.
Flow and timing	Blood samples were obtained just before the invasive procedure (reference standard). gNIPT was a second-tier test. 13/242 samples failed sequencing process quality control (no gNIPT result). No repeated test reported. 1 T13 cases was excluded to avoid double counting because it was published thereafter in Hall 2014.
Comparative	
Aim to study	To assess the performance of ccfDNA testing in maternal blood for detection of fetal aneuploidy of chromosomes 13, 18, 21, X, and Y using TMPS of single-nucleotide polymorphisms.

Nicolaides 2013 (Continued)

Funding source or sponsor of the study	Study funded by a grant from the Fetal Medicine Foundation (UK Charity No: 1037116). Analysis of samples was performed at their own expense by Natera, Inc.
Informations about the authors contacted	No need for further contact.
Notes	T13 cases data are not shown in the present review. They were excluded to avoid double count- ing because they are also published in Hall 2014.

Methodological quality

Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Unclear		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	No		
		High	Low
DOMAIN 2: Index Test TMPS			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		

Nicolaides 2013 (Continued)

Did all analysed patients receive the reference standard?	Yes	
Were all patients included in the analysis?	No	

Nicolaides 2014a

Study characteristics	
Patient sampling	Study design: case-control study.
	Participants: pregnant women selected from a high-risk population (archived maternal plas ma samples).
	Inclusion criteria: singleton pregnancies.
	Exclusion criteria: cases of fetal mosaicism and multifetal pregnancies.
Patient characteristics and setting	Recruited participants: 177 archived maternal plasma.
	Number available for 2 x 2 table: 172 samples (subgroup of 97%).
	Setting: recruitment in London, UK.
	Ethnicity: Caucasian (90%), Afro-Caribbean (4%), Asian (5%) and other (1%).
	Gestational age range: 11.2 to 14.1 weeks.
	Maternal age range: 17.3 to 47.8 years.
	Relevant tests carried out prior to index test: ultrasonography (nuchal translucency mea- surement) and biochemical screening.
	Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex.
	Median fetal fraction DNA (range): euploids: 13.0% (4.8% to 32.0%), 45,X: 10.0% (6.3% to 18.0%), and 47,XXX, 47,XXY, and 47,XYY: 12.0% (6.4% to 16.0%).
	Blood samples for gNIPT were collected just before reference standard.
	Cutpoint: positive if FORTE algorithm risk score \ge 1%.
	Commercial test: Harmony™ Prenatal Test by Ariosa Diagnostics, Inc.
Target condition and reference stan-	Target conditions: 45,X, 47,XXX, 47,XXY, and 47,XYY.
dard(s)	Reference standard: fetal karyotype of chorionic villi or amniotic fluid.
Flow and timing	Blood samples for gNIPT were collected just before invasive procedure (reference standard)
	gNIPT was a second-tier test.
	5/177 samples failed during sequencing process (no gNIPT result), including 1 sample failed laboratory quality control metrics and 4 samples failed for an insufficient fetal ccfDNA frac- tion.

High



Nicolaides 2014a (Continued)

No repeated test reported.

Comparative			
Aim to study			ective sequencing of cfDNA in mater- c of fetal sex chromosome aneuploi-
Funding source or sponsor of the study	No funding source was rep	orted.	
Informations about the authors con-	Author was contacted on: 1	10 February 2016.	
tacted	No reply received from the	author.	
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate ex- clusions?	No		
		High	Low
DOMAIN 2: Index Test TMPS			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi- tion?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			



Nicolaides 2014a (Continued)

Was there an appropriate interval be- tween index test and reference stan- dard?	Yes
Did all analysed patients receive the reference standard?	Yes
Were all patients included in the analysis?	No
	High

Norton 2012

Study characteristics	
Patient sampling	Study design: blinded, prospective cohort study. Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive testing. Inclusion criteria: pregnant women aged ≥ 18 years, at gestational age ≥ 10 weeks, with a sin- gleton pregnancy, who were planning to undergo invasive prenatal diagnosis for any indica- tion. Exclusion criteria: multifetal pregnancies, women with know aneuploidy, had active malignan- cy or a history of metastatic cancer, or had already undergone CVS or amniocentesis during the current pregnancy.
Patient characteristics and setting	Number enrolled: 4002 pregnant women. Number available for 2 x 2 table: 3080 pregnant women (subgroup of 77%). Setting: 48 centres. Selected prenatal care Centres in USA, the Netherlands and Sweden. Recruitment period: not reported. Ethnicity: Caucasian (49.6%), Afro-American (6.4%), Asian (13.4%), Hispanic (22.7%) and other (7.9%). Mean gestational age (± SD; range): 16.9 (± 4.1; 10 to 38.7) weeks. Mean maternal age (± SD; range): 34.3 (± 6.4; 18 to 50) years. Relevant test carried out prior to index test: not reported. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex. Mean fetal fraction DNA (± SD; range): euploids: 11% (± 4.5%; 4.2% to 51.3%), T21: 11.6% (± 4.2%; 5.1% to 23.3%), and T18: 10% (± 3.8%; 4.9% to 20.8%). Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if FORTE algorithm risk score ≥ 1%. Commercial test: Ariosa Diagnostics, Inc's prenatal test.
Target condition and reference standard(s)	Target conditions: T21 and T18. Reference standard: fetal karyotype of chorionic villi (74.7%) or amniotic fluid (25.3%).
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference standard). gNIPT was a second-tier test. 774/4002 samples excluded for ineligible criteria. 148/3228 samples failed during sequencing process (no gNIPT result), including 57 samples failed for low fetal fraction DNA and 91 samples failed sequencing process.
	No repeated test reported.



Norton 2012 (Continued)

Comparative					
Aim to study	To evaluate performance of a gNIPT of fetal T21 and T18.				
Funding source or sponsor of the study	Study funded by Ariosa Diagnostics, Inc.				
Informations about the authors contacted	No need for further contact.				
Notes					
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sam- ple of patients enrolled?	No				
Was a case-control design avoid- ed?	Yes				
Did the study avoid inappropriate exclusions?	No				
		High	Low		
DOMAIN 2: Index Test TMPS					
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes				
If a threshold was used, was it pre- specified?	Yes				
		Low	Low		
DOMAIN 3: Reference Standard					
Is the reference standards likely to correctly classify the target condi- tion?	Yes				
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes				
		Low	Low		
DOMAIN 4: Flow and Timing					

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Norton 2012 (Continued)	
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Was there an appropriate interval between index test and reference standard?	Yes	
Did all analysed patients receive the reference standard?	Yes	
Were all patients included in the analysis?	No	
	н	ligh

Norton 2015

Study characteristics				
Patient sampling	Study design: blinded, prospective cohort study.			
	Participants: unselected population of pregnant women undergoing aneuploidy screening (without prior risk of fetal aneuploidy).			
	Inclusion criteria: singleton pregnancies, at least 18 years of age, and between 10 to 14 weeks of ges- tation.			
	Exclusion criteria: women who had a miscarriage, chose to terminate the pregnancy or had a still- birth without confirmatory genetic testing.			
Patient characteristics and	Number enrolled: 18,955 pregnant women.			
setting	Number available for 2 x 2 table: 15,841 pregnant women (subgroup of 84%).			
	Setting: 35 centres in USA States, Canada, Sweden, the Netherlands, Belgium, and Italy.			
	Recruitment period: March 2012 to April 2013.			
	Ethnicity: Caucasian (70.9%), Afro-American (8.2%), Asian (10.5%), Native American (0.6%), multira- cial (2.7%), other (6.7%) and missing data (0.5%).			
	Mean gestational age (range): 12.5 (10.0 to 14.3) weeks.			
	Mean maternal age (range): 31 (18 to 48) years whose 76% of pregnant women analysed had < 35 years old.			
	Relevant tests carried out prior to index test: none.			
	Language of the study: English.			
Index tests	gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 sequencer in 96-plex.			
	Fetal fraction DNA: amount measured but not reported.			
	Blood samples for gNIPT were collected before reference standard.			
	Cutpoint: not reported. Usually, Harmony™ prenatal test uses FORTE algorithm; positive if FORTE risk score ≥ 1%.			
	Commercial test: Harmony™ Prenatal Test by Ariosa Diagnostics, Inc.			
	The traditional screening tests (combined test at the first trimester) were also assessed.			



lorton 2015 (Continued)	Cutpoint of combined test	: 1 in 270 for T21 or 1 in 150 for	T18 and T13.	
Target condition and refer-	Target conditions: T21, T18 and T13.			
ence standard(s)	Reference standards: fetal karyotype of chorionic villi, amniotic fluid or products of conception or neonatal karyotype, neonatal clinical examination or medical record from birth.			
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).			
	gNIPT was a first-tier test.			
	meet exclusion criteria, 31 withdrawn by investigator sult, 488 failed sequencing	had twins, 121 had unknown o , 384 had sample-handling erro , and have no gNIPT result (192	amples did not meet inclusion criteria or ovum-donor status, 64 withdrew or were ors, 308 without standard screening test re- for low fetal fraction DNA, 83 for non fetal ures) and 1489 were lost to follow-up.	
Comparative				
Aim to study	(with measurement of nuc		than standard first-trimester screening ical analytes) in risk assessment for trisomy g for aneuploidy screening.	
	To also evaluate the perfor somies 18 and 13.	rmance of gNIPT and standard :	screening in the assessment of risk for tri-	
Funding source or sponsor of the study	Study funded by Ariosa Diagnostics, Inc and Perinatal Quality Foundation.			
Informations about the au-	Author was contacted on: 10 February 2016.			
thors contacted	Reply received on: 11 February 2016.			
Notes	This study is a clinical trial (Noninvasive Examination of Trisomy (NEXT) ClinicalTrials.gov number, NCT01511458).			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoided?	Yes			
Did the study avoid inappro- priate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test TMPS				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes			



Norton 2015 (Continued)

If a threshold was used, was Yes it pre-specified?

		Low	Low
DOMAIN 2: Index Test Traditio	onal screening tests		
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standa	rd		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	No		
Were all patients included in the analysis?	No		
		High	

Palomaki 2012

Study characteristics	
Patient sampling	Study design: nested case-control (1:3) study. Participants: pregnant women at high risk of fetal aneuploidy presenting for invasive testing were selected. Inclusion criteria: ≥ 18 years old, between about 10 weeks and 21 weeks 6 days of gestation, at high risk of aneuploidies and who underwent a diagnostic procedure. Exclusion criteria: multifetal pregnancies or low risk of fetal aneuploidy.

Palomaki 2012 (Continued)			
Patient characteristics and setting	analysed samples from Palo Number available for 2 x 2 ta 2011) (subgroup of 42%). Setting: 27 centres. Prenatal Ireland, Hungary, USA, Israel Recruitment period: April 200 Ethnicity (only for 293 pregn and unknown (5.5%). Mean gestational age (range) Mean maternal age (± SD): 37	maki 2011. ble: 1971 pregnant women diagnostic centres (Canada and Australia). 09 to February 2011. ant women): Caucasian (84): 14.7 (9 to 22) weeks. 7.2 (± 5) years. for to index test: ultrasonog both.	t women selected for this study and 212 re- (1759 from this study + 212 from Palomaki a, Italy, Spain, Czech Republic, Argentina, .9%), Afro-american (4.1%), Asian (5.5%) graphy (nuchal translucency measurement)
Index tests	gNIPT by MPSS on Illumina H	liSeq 2000 sequencer in 4-p	olex.
	Mean (geometric) fetal fracti Blood samples for gNIPT wer Cutpoint: positive if Z score > Commercial test: Sequenom	re collected before reference > 3 (T21), > 3.88 (T18) or > 7.	e standard.
Target condition and refer- ence standard(s)	Target conditions: T21, T18 a Reference standard: fetal ka		nniotic fluid or products of conception.
Flow and timing	Blood samples were obtaine	d immediately prior the inv	vasive procedure (reference standard).
	gNIPT was a second-tier test. 2888/4664 samples were not		
	110/1776 samples failed the	initial MPSS testing.	
	105/110 samples required re quenced with the same libra		l aliquot and 5/110 samples were rese- d a gNIPT results.
	17/1776 samples failed durin sult).	g sequencing process, mos	t for low fetal fraction DNA (no gNIPT re-
Comparative			
Aim to study	To determine whether mater	nal plasma ccfDNA sequen	cing can identify T18 and T13 as well as T21.
Funding source or sponsor of the study	Study fully funded by Sequer	nom, Inc.	
Informations about the au- thors contacted	No need for further contact.		
Notes	This study is a clinical trial "A ber: NCT00877292.	New Prenatal Blood Test f	or Down Syndrome" ClinicalTrials.gov num-
Methodological quality			
ltem	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		



Palomaki 2012 (Continued)				
Was a case-control design avoided?	No			
Did the study avoid inappro- priate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard	d			
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate in- terval between index test and reference standard?	Yes			
Did all analysed patients re- ceive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		

Papageorghiou 2016a

 Study characteristics

 Patient sampling
 Study design: blinded, case-control study (1:9). Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive testing. Inclusion criteria: at least 18 years of age, singleton or twin pregnancies of at least 10 weeks' gestation and a clinical indication for an invasive procedure.

Papageorghiou 2016a (Continued)

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			riplets or more), known mosaicism, par- ıg twin, malignancy or known aneuploidy	
Patient characteristics and set- ting	 Number enrolled: 442 pregnant women. Number available for 2 x 2 table: 426 singleton pregnancies (subgroup of 96%). Setting: 6 hospital centres in England, UK. Recruitment period: April 2008 to November 2014. Ethnicity: not reported. Median gestational age (range): 15.4 (11 to 36.6) weeks. Median maternal age (range): 35 (18 to 55) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measu ment) or biochemical screening or both. Language of the study: English. 			
Index tests	gNIPT by MPSS on Ion Proto	n™ sequencer in 8-plex.		
	Fetal fraction DNA: amount Blood samples for gNIPT we Cutpoint: positive if likeliho Commercial test: IONA® test	re collected before reference od ratio > 1 and maternal ag	ce standard. ge-adjusted probability risk score.	
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or medical record from birth.			
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference standard). gNIPT was a second-tier test. 5/442 samples failed during sequencing process including 3 samples for low fetal fraction DNA and 2 samples did not have sufficient DNA fragment counts (no gNIPT result).			
	11/437 twin pregnancies were not selected.			
	No repeated test reported.			
Comparative				
Aim to study	To investigate the accuracy and those affected by fetal t		rimination between euploid pregnancies	
Funding source or sponsor of the study	Study funded by Premaitha Health (public limited company). Some authors are employees of Pre- maitha Health plc.			
Informations about the authors contacted	Author was contacted on: 19 September 2016. Reply received on: 20 September 2016.			
Notes	Data from singleton pregna pregnancies.	ncies only reported here. Se	ee Papageorghiou 2016b for data on twin	
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			



Papageorghiou 2016a (Continued)				
Was a case-control design avoided?	No			
Did the study avoid inappropri- ate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards like- ly to correctly classify the target condition?	Yes			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate inter- val between index test and ref- erence standard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		

Papageorghiou 2016b

 Study characteristics

 Patient sampling
 Study design: blinded, case-control study (1:9). Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive testing. Inclusion criteria: at least 18 years of age, a singleton or twin pregnancies of at least 10 weeks' gestation and a clinical indication for an invasive procedure.

Papageorghiou 2016b (Continued)

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Was a consecutive or random sample of patients enrolled?	No			
Item DOMAIN 1: Patient Selection	Authors' judgement	Risk of bias	Applicability concerns	
Methodological quality				
Notes	Data from twin pregnancies only reported here. Data from singleton pregnancies reported in Pa- pageorghiou 2016a.			
Informations about the authors contacted	Author was contacted on: 19 September 2016. Reply received on: 20 September 2016.			
Funding source or sponsor of the study	Study funded by Premaitha Health (public limited company). Some authors are employees of Pre- maitha Health plc.			
Aim to study	To investigate the accuracy and those affected by fetal t		mination between euploid pregnancies	
Comparative				
	No repeated test reported.			
	426/437 singleton pregnanc	-	nts (no gNIPT result).	
	5/442 samples failed during sequencing process including 3 samples for low fetal fraction DNA and 2 samples did not have sufficient DNA fragment counts (no gNIPT result).			
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference standard). gNIPT was a second-tier test.			
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or medical record from birth.			
	Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint: positive if likelihood ratio > 1 and maternal age-adjusted probability risk score. Commercial test: IONA® test by Premaitha Health (public limited company).			
Index tests	gNIPT by MPSS on Ion Proto	n™ sequencer in 8-plex.		
Patient characteristics and set- ting	Number enrolled: 442 pregnant women. Number available for 2 x 2 table: 11 twin pregnancies (subgroup of 2%). Setting: 6 hospital centres in England, UK. Recruitment period: April 2008 to November 2014. Ethnicity: not reported. Median gestational age (range): 15.4 (11 to 36.6) weeks. Median maternal age (range): 35 (18 to 55) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measure- ment) or biochemical screening or both. Language of the study: English.			
Detient de metricities en dest	tial trisomy or translocation in the pregnancy.	s, fetal demise, disappearing	plets or more), known mosaicism, par- g twin, malignancy or known aneuploidy	



Papageorghiou 2016b (Continued)				
Was a case-control design avoided?	No			
Did the study avoid inappropri- ate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards like- ly to correctly classify the target condition?	Yes			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate inter- val between index test and ref- erence standard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		

Pergament 2014

Study characteristics Patient sampling Study design: blinded, prospective cohort study. Participants: pregnant women from a population with mixed risk of fetal aneuploidy presenting for aneuploidy screening (51% high risk and 49% low risk). Inclusion criteria: women were 18 years of age or older with a singleton pregnancy of at least 7 weeks of gestation and signed an informed consent.

Pergament 2014 (Continued)	Exclusion criteria: women with confirmed sex chromosome abnormality (47,XXX, XXY, XYY), confirmed triploidy, confirmed fetal mosaicism or multifetal pregnancy or egg donor.
Patient characteristics and setting	Number enrolled: 1064 pregnant women. Number available for 2 x 2 table: 963 pregnant women for T21, 964 for T18 and 45,X and 965 for T13 (subgroup of 91%). Setting: 35 centres. Prenatal care centres worldwide in Czech Republic, Japan, USA, Ireland and Spain. Recruitment period: not reported. Ethnicity: not reported. Mean gestational age (± SD; range): 17.0 (± 4.1; 7.6 to 40.6) weeks. Median gestational age: 14.3 weeks. Mean maternal age (± SD; range): 30.3 (± 7.4; 18 to 47) years. Median maternal age: 30.0 years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or both. Language of the study: English.
Index tests	gNIPT by TMPS (SNP-based method) on Illumina Genome Analyzer IIx or HiSeq sequencers, 19,488-plex targeted PCR with NATUS algorithm. Range fetal fraction DNA: 2% to 50%. Blood samples for gNIPT were collected before (93%) or 4 days or later after (7%) reference standard. Cutpoint: not reported. Commercial test: Natera's prenatal test.
Target condition and ref- erence standard(s)	Target conditions: T21, T18, T13 and 45,X. Reference standards: fetal karyotype with confirmatory fluorescence in situ hybridisation or cytoge- netic karyotype analysis or by genetic testing of cord blood, buccal sample, saliva, or products of con- ception, post-natal or post-live birth follow-up.
Flow and timing	Blood samples for gNIPT were obtained prior (93%) or after (7%) to the invasive procedure (reference standard). gNIPT was a second-tier test.
	13/1064 samples excluded for other aneuploidies, including 6 cases with triploidy, 3 fetal mosaics, 2 cases with 47,XXY, 1 case with 47,XXX and 1 case with 47,XYY.
	85/1051 samples failed quality control (no gNIPT result) including 64 low fetal fraction DNA, 12 low DNA, 6 contaminations, 2 loss of heterozygosity and 1 poor model fit.
	Between 1 to 3 samples did not passed quality control for all 5 chromosomes.
	No repeated test reported.
Comparative	
Aim to study	To estimate performance of a single nucleotide polymorphism–based gNIPT (TMPS) for fetal aneu- ploidy in high-risk and low-risk populations on single venipuncture.
Funding source or sponsor of the study	Study funded by Natera, Inc. and a grant from the National Institute of Health, National Institute of Child Health and Human Development (4R44HD062114-02). The majority of the authors are employees of Natera, Inc. and hold stock or options to hold stock in the company.
Informations about the authors contacted	Author was contacted on: 22 June 2016. No reply received from the author.
Notes	
Methodological quality	



Pergament 2014 (Continued) Item **Authors' judgement Risk of bias Applicability concerns DOMAIN 1: Patient Selection** Was a consecutive or ran-Unclear dom sample of patients enrolled? Was a case-control design Yes avoided? Did the study avoid inap-No propriate exclusions? High High **DOMAIN 2: Index Test TMPS** Were the index test re-Yes sults interpreted without knowledge of the results of the reference standard? If a threshold was used, Yes was it pre-specified? Low Low **DOMAIN 3: Reference Standard** Is the reference standards Yes likely to correctly classify the target condition? Were the reference stan-Yes dard results interpreted without knowledge of the results of the index tests? Low Low **DOMAIN 4: Flow and Timing** Was there an appropriate Yes interval between index test and reference standard? Did all analysed patients Yes receive the reference standard? Were all patients included No in the analysis? High



Persico 2016

Study characteristics			
Patient sampling	Study design: blinded, pro Participants: pregnant wo Inclusion criteria: singleto Exclusion criteria: multifet	men selected from a high-ri n pregnancies.	sk population.
Patient characteristics and setting	Setting: 4 fetal medicine c Recruitment period: March Ethnicity: not reported. Gestational age: not repor Median maternal age (rang	table: 249 pregnant womer entres in Italy. n to December 2014. ted. ge): 36 (20 to 46) years. prior to index test: ultrason screening or both.	n (subgroup of 96%). ography (nuchal translucency mea-
Index tests	gNIPT by TMPS (SNP-base 19,488-plex targeted PCR v		ome Analyzer IIx or HiSeq sequencers,
	trol criteria > 4%).	vere collected just before re sk score > 1%.	ed (usually NATERA used quality con- ference standard.
Target condition and reference stan- dard(s)	case was found.	8, T13, 45,X, 47,XXY and 47,) karyotype of chorionic villi (XXX. 47,XYY was also assessed but no or amniotic fluid.
Flow and timing	dard). gNIPT was a second-tier te 10/259 samples failed duri	est.	wasive procedure (reference stan- gNIPT result) including 2 samples fetal fraction DNA.
	No repeated test reported		
Comparative			
Aim to study	To investigate a strategy fo cies after first-trimester co		of ccfDNA testing in high-risk pregnan-
Funding source or sponsor of the study	Study not funded by indus	try but the cost of ccfDNA to	esting were covered by Natera, Inc.
Informations about the authors con- tacted	No need for further contac	:t.	
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns

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Persico 2016 (Continued)				
Was a consecutive or random sample of patients enrolled?	Yes			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate ex- clusions?	No			
		High	Low	
DOMAIN 2: Index Test TMPS				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to cor- rectly classify the target condition?	Yes			
Were the reference standard results in- terpreted without knowledge of the re- sults of the index tests?				
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analy- sis?	- No			
		High		
Poon 2016				
Study characteristics				
Patient sampling		or invasive testing (CVS).	case-control study. ant women selected at high risk of f	etal

Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies.



Poon 2016 (Continued)			
Patient characteristics and setting	Number enrolled: 242 pregn Number available for 2 x 2 ta Setting: 1 centre at King's Co Recruitment period: April 20 Ethnicity: Caucasian (75%), <i>J</i> Median gestational age (range Relevant tests carried out pr ment) or biochemical screer Language of the study: Engli	ble: 241 pregnant women ollege Hospital, London, Uk 07 to June 2012. Afro-Caribbean (17%), Asia ge): 12.7 (11.4 to 13.6) week): 33.7 (29.2 to 40.5) years. ior to index test: ultrasono ning or both.	r. (5%) and mixed (3%).
Index tests	gNIPT by MPSS on Ion Proto	n™ sequencer.	
		re collected just before refe uthors used the same prer od ratio > 1 and maternal a	erence standard. natal test than Papageorghiou 2016a ge-adjusted probability risk score).
			ic innited company).
Target condition and reference standard(s)	Target conditions: T21, T18 a Reference standard: fetal ka		
Flow and timing	Blood samples for gNIPT we dard). gNIPT was a second-tier test 1/242 samples failed for low No repeated test reported.		invasive procedure (reference stan- PT result).
Comparative			
Aim to study	To assess the potential perfo sis of maternal blood using t		tal T21, T18 and T13 by ccfDNA analy-
Funding source or sponsor of the study	Study not funded by industr chester, UK. Study supporte		ovided by Premaitha Health plc, Man- Medicine Foundation.
Informations about the authors con- tacted	Author was contacted on: 19 No reply received from the a	•	
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	No		
Was a case-control design avoided?	No		
Did the study avoid inappropriate exclusions?	No		
		High	Low



Poon 2016 (Continued)

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DOMAIN 2: Index Test MPSS			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Porreco 2014

Study characteristics Study design: blinded, prospective cohort, observational study. Patient sampling Participants: pregnant women selected at high risk of fetal aneuploidy presenting for invasive testing when research personnel have been available. Inclusion criteria: singleton pregnancy in a patient 18 years of age or older who had provided written informed consent and who had made the decision to pursue invasive prenatal diagnosis by CVS or amniocentesis. Exclusion criteria: inability to give written informed consent, multifetal pregnancies, or fetal demise of an additional embryo during the current pregnancy at 8 weeks or more of gestation. Patient characteris-Number enrolled: 4170 pregnant women. Number available for 2 x 2 table: 3322 for autosomes (subgroup of 80%), 3278 for 45,X and 47,XXX (subgroup tics and setting of 79%) and 3201 for 47,XXY and 47,XYY (subgroup of 77%). Setting: 31 centres in USA. Recruitment period: September 2009 to April 2011. Ethnicity: Caucasian (60,1%), Asian (18,7%), Afro-American (4,5%) and other (16.7%).



Porreco 2014 (Continued)	Mean gestational age (± SD; range): 16.3 (± 3.5; 9.0 to 37.0). Mean maternal age (± SD; range): 35.1 (± 5.6; 18.0 to 50.0). Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and bio- chemical screening. Language of the study: English.
Index tests	gNIPT by MPSS on Illumina HiSeq 2000 in 12-plex.
	Range fetal fraction DNA: 4% to 50%. Blood samples for gNIPT were collected before reference standard. Cutpoint:
	1) for T21, positive if Z score \geq 3.
	2) for T18 and T13, positive if Z score ≥ 3.95.
	3) positive for 45,X if Z score < -3.5 (non-reportable regions between -2.5 and -3.5).
	4) positive for 47,XXX if Z score > 3.5 (non-reportable regions between 2.5 and 3.5).
	5) positive for 47,XYY if Z score risk < -3.5 with Chrom. Y representation.
	6) positive for 47,XXY if Z score risk is between -3.5 and 3.5 with Chrom. Y representation.
	Commercial test: Sequenom's prenatal test.
Target condition and reference stan- dard(s)	Target conditions: T21, T18, T13, 45,X, 47,XXX, 47,XXY and 47,XYY. Reference standards: fetal karyotype of chorionic villi or amniotic fluid, or medical record from birth.
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).
	gNIPT was a second-tier test. 740/4170 samples excluded before sequencing process including 320 samples for insufficient sample vol- ume,120 samples processed outside of the 6 hours laboratory process window, 270 failed laboratory quality control set, 24 for incomplete case report form and 6 without invasive procedure performed).
	For autosomes: 54/3430 autosomes samples excluded for quality control deviation (low fetal DNA fraction, library concentration, total counts, and amplification bias).
	For autosomes: 54/3376 samples excluded for complex autosome karyotypes (mosaic, triploidies, unbal- anced rearrangements with missing or duplicated genetic material).
	For 45,X and 47,XXX: 102/3430 samples excluded for low fetal fraction DNA or copy number variation of the Chrom. X is confounded by maternal component and cannot be determined.
	For 45,X and 47,XXX: 50/3328 samples excluded for complex SCA karyotype.
	For 47,XXY and 47,XYY: 182/3430 samples excluded for low fetal fraction DNA or copy number variation of the Chrom. X is confounded by maternal component and cannot be determined.
	For 47,XXY and 47,XYY: 47/3248 samples excluded for complex SCA karyotype.
	No repeated test reported.
Comparative	
Aim to study	To validate the clinical performance of MPSS of ccfDNA contained in specimens from pregnant women at high risk of fetal aneuploidy to test fetuses for T21, T18, T13, 45,X, 47,XXX, 47,XXY and 47,XYY.
Funding source or sponsor of the study	Study funded by Sequenom, Inc.



Porreco 2014 (Continued)

Informations about	Author was contacted on: 30 May 2016.
the authors contact-	Reply received on: 31 May 2016.
ed	

Notes

This study is a clinical trial (Non-Invasive Screening for Fetal Aneuploidy) ClinicalTrials.gov number, NCT00847990.

Methodological quality

ltem	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Se	election		
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclu- sions?	No		
		High	Low
DOMAIN 2: Index Test	t MPSS		
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 3: Reference	e Standard		
Is the reference stan- dards likely to cor- rectly classify the tar- get condition?	Yes		
Were the reference standard results in- terpreted without knowledge of the results of the index tests?	Yes		



Porreco 2014 (Continued)	
Was there an appro- priate interval be- tween index test and reference standard?	Yes
Did all analysed pa- tients receive the ref- erence standard?	Yes
Were all patients in- cluded in the analy- sis?	Νο
	High

Quezada 2015

Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: self-selected pregnant women from the general population presenting for aneu- ploidy screening (without prior risk of fetal aneuploidy). Inclusion criteria: pregnant women between 10 to 11 weeks' gestation with singleton pregnan cy who underwent the combined test. Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 2905 pregnant women. Number available for 2 x 2 table: 2785 pregnant women (subgroup of 96%). Setting: 1 centre. Fetal Medicine Centre in London, UK. Recruitment period: October 2012 to January 2014. Ethnicity: Caucasian (88.5%), South Asian (6.0%), East Asian (3.3%), Afro-Caribbean (0.7%) and mixed (1.5%). Median gestational age (range): 10.6 (10 to 11.9) weeks. Median maternal age (range): 36.9 (20.4 to 51.9) years. Relevant tests carried out prior to index test: none. Language of the study: English.
Index tests	gNIPT by TMPS (DANSR assay). Median fetal fraction DNA (range): 11% (4% to 40%). Blood samples for gNIPT were collected before reference standard. Cutpoint: not reported. Usually, Harmony [™] prenatal test uses FORTE algorithm; positive if FORTE risk score ≥ 1%. Commercial test: Harmony [™] Prenatal test by Ariosa Diagnostics, Inc. The traditional screening tests (combined test at the first trimester) was also assessed. Cutpoint of combined test: 1 in 100 for T21.
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi, amniotic fluid or products of concep- tion, neonatal karyotype, neonatal clinical examination or medical record from birth.
Flow and timing	Blood samples were obtained prior to invasive procedure (reference standard). gNIPT was a first-tier test. 122/2905 failed the initial TMPS testing (122 = 123 - 1 sample lost in mail).

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Quezada 2015 (Continued)	66/2851 samples without fo	ollow-up were excluded.		
	 110/122 required repeat testing using a second blood sample and results were obtained in 69/110 samples. 53/2905 samples failed during sequencing process (41 samples failed second sequencing and 12 unrepeated tests) (no gNIPT result). 			
Comparative				
Aim to study		testing for T21, T18 and T1	without prior risk of fetal aneuploidy), 3 at 10 to 11 weeks' gestation and com- gestation.	
Funding source or sponsor of the study	Study not funded by indust	ry but Ariosa Diagnostics, Iı	nc made sequencing and analyses.	
Informations about the authors contacted	Author was contacted on: 2 No reply received from the		16.	
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sam- ple of patients enrolled?	No			
Was a case-control design avoid- ed?	Yes			
Did the study avoid inappropriate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test TMPS				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 2: Index Test Traditional s	creening tests			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			



Quezada 2015 (Continued)			
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi- tion?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	

Study characteristics	
Patient sampling	Study design: blinded, prospective cohort study. Participants: pregnant women at high or low risk of fetal aneuploidy (known sex chromosome ane uploidy and euploid pregnancies). Inclusion criteria: women were at least 18 years of age, had singleton pregnancy, or with known sex chromosome aneuploidy. Exclusion criteria: pregnant women with known mosaicism, autosomal trisomy, or triploidy.
Patient characteristics and setting	Number enrolled: 201 pregnant women. Number available for 2 x 2 table: 186 pregnant women (subgroup of 93%). Setting: 8 prenatal care centres in UK, USA, Poland, and Czech Republic. Recruitment period: not reported. Ethnicity: not reported. Mean gestational age: euploid pregnancies 13.2 weeks, and aneuploid pregnancies 15.3 weeks. Gestational age range: overall 9.4 to 36.4 weeks. Maternal age: not reported. Relevant test carried out prior to index test: not reported. Language of the study: English.
Index tests	gNIPT by TMPS (SNP-based method) on Illumina HiSeq 2000 sequencer with NATUS algorithm. Mean fetal fraction DNA: euploids: 10.9% and aneuploids: 12.1%. Overall range: 2.9% to 37.7%. Blood samples for gNIPT were collected just before or at least 4 days after reference standard. Cutpoint: not reported.

amango-Sprouse 2013 (Continue	^{d)} Commercial test: Natera's p	orenatal test.		
Target condition and refer- ence standard(s)	Target conditions: 45,X, 47,XXX, 47,XXY, and 47,XYY. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or genetic testing of cord blood, buccal sample, saliva, or products of conception.			
Flow and timing	Blood samples were collected just before or at least 4 days after invasive procedure (reference standard). gNIPT was a first- or second-tier test. 14/201 samples failed sequencing process quality control (no gNIPT result) including 12 for low fe- tal fraction or poor DNA quality and 2 samples did not return a result for SCA.			
	1/187 sample excluded for a	conflicting algorithm metric	cs (no meaningful gNIPT result).	
	No repeated test reported.			
Comparative				
Aim to study	To develop a SNP-based and informatics-based gNIPT that detects sex chromosome aneuploidies early in pregnancy.			
Funding source or sponsor of the study	It is unclear if the study was funded by industry but all authors are employees of Natera, Inc. except the first author (Carole Samango-Sprouse). This study was supported in part by a grant from the National Institute of Health, National Institute of Child Health and Human Development.			
Informations about the au-	Author was contacted on: 22 April, 4 July and 29 September 2016.			
thors contacted Replies received on: 29 and 30 September 2016.				
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappro- priate exclusions?	No			
		High	High	
DOMAIN 2: Index Test TMPS				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes			

If a threshold was used, was it Yes pre-specified?



Samango-Sprouse 2013 (Continued)					
		Low	Low		
DOMAIN 3: Reference Standard	DOMAIN 3: Reference Standard				
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes				
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes				
		Low	Low		
DOMAIN 4: Flow and Timing					
Was there an appropriate in- terval between index test and reference standard?	Yes				
Did all analysed patients re- ceive the reference standard?	Yes				
Were all patients included in the analysis?	No				
		High			

Study characteristics	
Patient sampling	Study design: blinded retrospective study (archived maternal plasma samples). Participants: pregnant women selected from a high risk of fetal aneuploidy population. Inclusion criteria: pregnant women age 18 years or older with singleton or multifetal pregnancy. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: overall: 1014 pregnant women including 71 women selected on 435 for the train- ing set (not shown in the present review) and 48 women selected on 575 for the test set. Number available for 2 x 2 table: 47 (subgroup of 8%). Setting: 13 centres in USA. Recruitment period: January 2010 to June 2010. Ethnicity: Caucasian (62.7%), Hispanic (16.5%), Asian (6.2%), multiethnic (5.2%), Afro-American (4.0%), Native American (0.9%) and other or not specified (1.8%). Mean gestational age (range): 15.4 (10.6 to 28.4) weeks. Mean maternal age (± SD; range): 34.2 (± 8.22; 18 to 46) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or both. Language of the study: English.
Index tests	gNIPT by MPSS on Illumina Genome Analyzer IIx sequencer in uniplex. Fetal fraction DNA: not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint prespecified with the training set:

ehnert 2011 (Continued)	1) positive if NCV > 4 for autos	somes. There is a "no call zon	e" between 2.5 and 4 considering as gNIPT	
	1) positive if NCV > 4 for autosomes. There is a "no call zone" between 2.5 and 4 considering as gNIP positive result for the present review.			
	2) positive if NCV for Chrom. \ -3.0 SDs from the mean of fen Commercial test: Verinata's p	nale samples for 45,X.	male samples and if NCV for Chrom. X <	
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13 and 45,X. Reference standard: fetal karyotype of chorionic villi (58,3%) or amniotic fluid (41.7%).			
Flow and timing	Blood samples for gNIPT were gNIPT was a second-tier test.	e obtained prior to the invasiv	e procedure (reference standard).	
	895/1014 samples were not s	elected for sequencing.		
	71/119 samples were selected	d for the training set (not show	vn in the present review).	
	1/48 sample from twin gestat	ion in the test set was remove	ed from the final analysis.	
	No repeated test reported.			
Comparative				
Aim to study		mapping and chromosome q	ta and demonstrated the potential uni- uantification method for the detection of	
Funding source or sponsor of the study		choice of enrolled patients,	unding organizations played a direct role he review and interpretation of data, and	
Informations about the au- thors contacted	No need for further contact.			
Notes				
Methodological quality				
ltem	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappro- priate exclusions?	No			
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results in- terpreted without knowledge	Yes			



Sehnert 2011 (Continued)

If a threshold was used, was Yes it pre-specified?

		Low	Low	
DOMAIN 3: Reference Standa	DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate in- terval between index test and reference standard?	Yes			
Did all analysed patients re- ceive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Low		

Shaw 2014 Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: consecutive pregnant women were selected from a mixed-risk population. They were classified in extremely high-risk group for T21 with a screening T21 risk > 1:30 or nuchal translucer cy > 3.0 mm and low-risk group with a screening T21 risk < 1:1500. Inclusion criteria: pregnant women at > 12 weeks' gestation, singleton or multifetal pregnancies. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 201 pregnant women. Number available for 2 x 2 table: 200 pregnant women (subgroup of 99.5%). Setting: 11 medical centres in Taiwan. Recruitment period: June to December 2012. Ethnicity: Asian. Mean gestational age (± SD): high-risk pregnant women 17.3 (± 2.1) weeks, and low-risk pregnant women 16.1 (± 3.0) weeks. Gestional age range: overall 12 to 20 weeks.
	Mean maternal age (± SD): high-risk pregnant women 35.1 (± 3.2) years, and low-risk pregnant women 34.6 (± 2.6) years. Chorionicity: all dichorionic (4/4).



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Shaw 2014 (Continued)	Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) and biochemical screening. Language of the study: English.				
Index tests	gNIPT by MPSS on Illumina v2	2 HiSeq 2000 sequencer in 1	2-plex.		
	Fetal fraction DNA: not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint:				
	1) positive if Z score > 3 (T21,	T18, and T13).			
	2) positive if Z score Chrom. A	< -3 and Z score Chrom. Y <	< 3 (45,X).		
	3) positive if Z score Chrom. X Commercial test: Berry Geno		• 3 (47,XYY).		
	The traditional screening test data for 2 x 2 tables were una		trimester) was also assessed but complete		
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T case was found. Reference standards: fetal ka		(and 47,XXY were also screened but no medical record from birth.		
Flow and timing	Blood samples for gNIPT wer gNIPT was a second-tier test. 1 sample excluded for early g		sive procedure (reference standard).		
	No repeated test reported.				
Comparative					
Aim to study	To evaluate the performance of gNIPT for all fetal chromosomal aneuploidies in an extremely high- risk group undergoing first-trimester combined T21 screening.				
Funding source or sponsor of the study	Funding sources were not reported but 2 authors are affiliated to Berry Genomics Co. Ltd., Beijing, PR China.				
Informations about the au- thors contacted	Author was contacted on: 10 No reply received from the au				
Notes					
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	Unclear				
Was a case-control design avoided?	Yes				
Did the study avoid inappro- priate exclusions?	No				
		High	High		



Shaw 2014 (Continued)				
DOMAIN 2: Index Test MPSS				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard	1			
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate in- terval between index test and reference standard?	Yes			
Did all analysed patients re- ceive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Low		

Song 2013

Study characteristics	
Patient sampling	Study design: blinded, prospective cohort study. Participants: pregnant women without a priori risk of fetal aneuploidy who undergo routine prenatal screening. Inclusion criteria: singleton pregnancies and pregnant women younger than 35 years old. Exclusion criteria: not reported.
Patient characteristics and set- ting	Number enrolled: 1916 pregnant women. Number available for 2 x 2 table: 1741 pregnant women (subgroup of 91%). Setting: 2 clinical centres in Beijing, China. Recruitment period: April 2011 to December 2011. Ethnicity: Asian. Mean gestational age (± SD; range): 16.57 (± 1.56; 11 to 21.9) weeks. Mean maternal age (± SD; range): 29.03 (± 2.70; 20 to 34) years.



ong 2013 (Continued)	Relevant tests carried out p Language of the study: Eng			
Index tests	gNIPT by MPSS on Illumina	v2 HiSeq 2000 sequencer in 1	2-plex.	
	Fetal fraction DNA: not repo Blood samples for gNIPT we Cutpoint: positive if Z score Commercial test: Berry Gen	ere collected before reference ≥3.	standard.	
	The traditional screening te	est (second-trimester triple tes	st) was also assessed.	
	Cutpoint of triple test: 1 in 2	270 for T21 and T18.		
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. 45,X, 47,XXX, 47, XXY, 47,XYY were also screened but inap propriate reference standard for the present review was used. Reference standards: fetal karyotype of chorionic villi, amniotic fluid or cord blood or medica record from birth.			
Flow and timing	It is not reported if the blood samples were collected before or after invasive procedure (refer- ence standard). It is not reported if the gNIPT was a first- or second-tier test. 64/1916 samples failed sequencing process (failed DNA quality control criteria or sequencing quality control) (no gNIPT result).			
	102/1916 samples without follow-up were excluded.			
	9/1916 samples were without follow-up and failed sequencing process (no gNIPT result).			
	No repeated test reported.			
Comparative				
Aim to study	women younger than 35 ye	ars old in a prospective clinica	al aneuploidies in a Chinese cohort of al setting. Also, to compare the perfor- ond-trimester combined test).	
Funding source or sponsor of the study	Study not funded by industry. This study was supported by a grant (2006BAI05A10) from the Na- tional Key Technology Research and Development Program of China during the '11th Five-Year Plan'.			
Informations about the authors contacted	No need for further contact			
Notes	SCA were also screened but gNIPT data from SCA were r		dard for the present review was used.	
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Was a case-control design avoid- ed?	Yes			



Song 2013 (Continued)

Did the study avoid inappropriate No exclusions?

		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 2: Index Test Traditional	screening tests		
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards like- ly to correctly classify the target condition?	Yes		
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all analysed patients receive the reference standard?	No		
Were all patients included in the analysis?	No		
		High	

ong 2015			
Study characteristics			
Patient sampling	Study design: blinded, prosp Participants: pregnant wome for aneuploidy screening by g Inclusion criteria: advanced r Exclusion criteria: multifetal	n selected arbitrarily at high NIPT. naternal age (≥ 35 years) and	n risk of fetal aneuploid presenting d singleton pregnancies.
Patient characteristics and setting	Number enrolled: 213 pregna Number available for 2 x 2 tal Setting: 1 centre. Peking Unio Recruitment period: May 201 Ethnicity: Asian. Median gestational age (rang Mean maternal age (range): 3 Relevant tests carried out pri Language of the study: Englis	ole: 204 pregnant women (su on Medical College Hospital (2 to August 2013. e): 9.9 (8 to 12.9) weeks. 7.25 (35 to 45) years. or to index test: none.	
Index tests	gNIPT by MPSS on Illumina v2 Median fetal fraction DNA (ra Blood samples for gNIPT wer Cutpoint: positive if Z score ≥ Commercial test: Berry Geno	nge): only male fetus: 8.54% e collected before reference 3.	(2.69% to 18.75%).
Target condition and reference standard(s)	Target conditions: T21, T18, T case was found. Reference standards: fetal ka women (178/178) and neona	ryotype of chorionic villi or a	
Flow and timing	Blood samples for gNIPT wer gNIPT was a first-tier test. 1/213 sample failed quality co		e procedure (reference standard).
	8/212 samples without refere trauterine fetal deaths and 1		l including 5 miscarriages, 2 in-
	No repeated test reported.		
Comparative			
Aim to study	To evaluate the feasibility of a nese women in early gestation		amples collected from pregnant Chi- s' gestation.
Funding source or sponsor of the study	Study not funded by industry Study funded by a grant from		analysed at Berry Genomics Co. Ltd. Se Foundation of China.
Informations about the authors con- tacted	No need for further contact.		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			

Cochrane Library

Song 2015 (Continued)					
Was a consecutive or random sam- ple of patients enrolled?	Yes				
Was a case-control design avoided?	Yes				
Did the study avoid inappropriate exclusions?	No				
		High		Low	
DOMAIN 2: Index Test MPSS					
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes				
If a threshold was used, was it pre- specified?	Yes				
		Low		Low	
DOMAIN 3: Reference Standard					
Is the reference standards likely to correctly classify the target condition?	Yes				
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes				
		Low		Low	
DOMAIN 4: Flow and Timing					
Was there an appropriate interval between index test and reference standard?	Yes				
Did all analysed patients receive the reference standard?	Yes				
Were all patients included in the analysis?	No				
		High			
Sparks 2012a					
Study characteristics					
Patient sampling	Study design: case-control s Participants: pregnant wom Inclusion criteria: women at gleton pregnancy.	en selected from	a high risk of fet		

parks 2012a (Continued)	Exclusion criteria: multife	al pregnancies.		
Patient characteristics and setting	the training set (data not s were selected for this stud Number available for 2 x 2 Setting: not reported. Recruitment period: not re Ethnicity: not reported. Mean gestational age (± SD; Mean maternal age (± SD;	hown in the present review) y. table: 167 pregnant women ported. D; range): 18.6 (± 4.0; 11 to 36 range): 33.5 (± 7.1; 18 to 51) y prior to index test: not repor	.1) weeks. /ears.	
Index tests	gNIPT by TMPS (DANSR as gorithm.	say) on Illumina HiSeq 2000 :	sequencer in multiplex with FORTE al	
	standard. Cutpoint: not reported.		ollected before or after reference t.	
Target condition and reference stan- dard(s)	Target conditions: T21 and T18. Reference standards: fetal karyotype or chromosome analysis by FISH or both.			
Flow and timing	It is not reported if the blo erence standard). gNIPT was a second-tier te 171/338 samples were exc No failed sample reported	st. luded for the training set.	efore or after invasive procedure (ref	
	No repeated test reported.			
Comparative				
Aim to study		mical assay and algorithm fo DNA obtained from materna	or the prenatal evaluation of risk for al blood.	
Funding source or sponsor of the study	Study funded by Ariosa Diagnostics, Inc. All authors are employees of Aria Dx Inc. (now Ar- iosa Diagnostics). K Sparks is a member of the board of the company.			
Informations about the authors con- tacted	Author was contacted on: 23 June 2016. No reply received from the author.			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	No			



Sparks 2012a (Continued)

Did the study avoid inappropriate ex-No clusions?

		High	Low	
DOMAIN 2: Index Test TMPS				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Unclear			
		Unclear	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condi-tion?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Unclear			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Unclear		

Study characteristics	
Patient sampling	Study design: prospective cohort study. Blinded for T21 and unblinded for T18 and T13. Participants: all consecutively enrolled pregnant women selected at high risk of fetal aneu- ploidy. Inclusion criteria: pregnant women at least 18 years old, at high risk for chromosomal aberra- tions, signed informed consent, planned a conventional karyotyping procedure (invasive diag- nostic), had singleton pregnancy and blood drawn before the invasive procedure. Exclusion criteria: multifetal pregnancies.
Patient characteristics and set- ting	Number enrolled: 522 pregnant women. Number available for 2 x 2 table: 472 pregnant women (subgroup of 90%).

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Authors' judgement	nd GATC Biotech AG. ruary 2016, 24 February and 19		
Germany and Switzerland. Study funded by LifeCodexx AG a Author was contacted on: 22 Feb Reply received on: 24 February 20	nd GATC Biotech AG. ruary 2016, 24 February and 19 016.		
Germany and Switzerland. Study funded by LifeCodexx AG a Author was contacted on: 22 Feb	nd GATC Biotech AG. ruary 2016, 24 February and 19		
Germany and Switzerland. Study funded by LifeCodexx AG a Author was contacted on: 22 Feb	nd GATC Biotech AG. ruary 2016, 24 February and 19		
Germany and Switzerland. Study funded by LifeCodexx AG a Author was contacted on: 22 Feb	nd GATC Biotech AG. ruary 2016, 24 February and 19		
Germany and Switzerland.		, T18 and T13 for a population in	
-	cy of a given infor acceeding 121	, T18 and T13 for a population i	
	cy of a gNIPT for detecting T21		
No repeated test reported.			
32/504 samples failed during sequencing process (no gNIPT result), including 14 samples failed sequencing quality criteria and 18 samples failed libraries.			
gNIPT was a second-tier test.			
Target conditions: T21, T18 and T13. Reference standard: fetal karyotype of chorionic villi (30.3%), amniotic fluid (69.1%) or cord blood (0.6%).			
2) positive if MAD-based Z score ≥	: 3.2 for T18.		
1) positive if MAD-based Z -score	≥ 3 for T21.		
gNIPT by MPSS on Illumina HiSec correction.	2000 sequencer in 12-plex wit	h DAP.21 algorithm without CG	
Recruitment period: not reported Ethnicity: not reported. Mean gestational age (range): 15. Mean maternal age (range): 36.0 Relevant tests carried out prior to	l. 6 (11.0 to 32.1) weeks. (19 to 47) years. p index test: ultrasonography (I	nuchal translucency measure-	
	 Recruitment period: not reported. Ethnicity: not reported. Mean gestational age (range): 15. Mean maternal age (range): 36.00 Relevant tests carried out prior to to ment) or biochemical screening of Language of the study: English. gNIPT by MPSS on Illumina HiSec correction. Mean fetal fraction DNA (range): n Blood samples for gNIPT were correction: 1) positive if MAD-based Z score and the study: English if MAD-based Z score and the study: Score and the study if MAD-based Z score and the study is previously analysed. 32/504 samples failed during seq sequencing quality criteria and 1. 	 Mean gestational age (range): 15.6 (11.0 to 32.1) weeks. Mean maternal age (range): 36.0 (19 to 47) years. Relevant tests carried out prior to index test: ultrasonography (rement) or biochemical screening or both. Language of the study: English. gNIPT by MPSS on Illumina HiSeq 2000 sequencer in 12-plex wit correction. Mean fetal fraction DNA (range): male fetus only: 12.3% (3.7% to Blood samples for gNIPT were collected just before reference st Cutpoint: positive if MAD-based Z -score ≥ 3 for T21. positive if MAD-based Z score ≥ 3.2 for T18. positive if MAD-based Z score ≥ 3.9 for T13. Commercial test: LifeCodexx's prenatal test. Target conditions: T21, T18 and T13. Reference standard: fetal karyotype of chorionic villi (30.3%), ar blood (0.6%). Blood samples for gNIPT were obtained just prior the invasive p gNIPT was a second-tier test. 18/522 samples excluded, including 8 without reference standard 1 was previously analysed. 32/504 samples failed during sequencing process (no gNIPT rest sequencing quality criteria and 18 samples failed libraries. 	



Stumm 2014 (Continued)					
Was a case-control design avoid- ed?	Yes				
Did the study avoid inappropriate exclusions?	No				
			High	Low	
DOMAIN 2: Index Test MPSS					
Were the index test results inter- preted without knowledge of the results of the reference standard?	No				
If a threshold was used, was it pre-specified?	No				
			High	Low	
DOMAIN 3: Reference Standard					
Is the reference standards like- ly to correctly classify the target condition?	Yes				
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes				
			Low	Low	
DOMAIN 4: Flow and Timing					
Was there an appropriate interval between index test and reference standard?	Yes				
Did all analysed patients receive the reference standard?	Yes				
Were all patients included in the analysis?	No				
			High		
Sukhikh 2015					
Study characteristics					
Patient sampling		Study design: prospec Participants: pregnan invasive testing.		al aneuploidy p	presenting for

Genomics-based non-invasive prenatal testing for detection of fetal chromosomal aneuploidy in pregnant women (Review) Copyright © 2017 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd.

Inclusion criteria: not reported. Exclusion criteria: not reported.

ukhikh 2015 (Continued) Patient characteristics and setting	Number enrolled: 200 pre	gnant women			
Fatient characteristics and setting	Number enrolled: 200 pregnant women. Number available for 2 x 2 table: 200 pregnant women (whole cohort included in analy-				
	ses). Setting: not reported. Recruitment period: not reported. Ethnicity: not reported. Median gestational age (range): 14 (10 to 20) weeks.				
	Maternal age: not reporte				
	Relevant tests carried out measurement) or biocher	prior to index test: ultrasor	nography (nuchal translucency		
	Language of the study: Ru				
Index tests	gNIPT by MPSS on Ion Pro	oton™ sequencer.			
	Fetal fraction DNA: not re	ported.			
	Blood samples for gNIPT	were collected before refere	ence standard.		
	Cutpoint:				
	1) Positive for T21 and T18 if T score > 5.				
	2) Positive for T13 if T score > 4.				
	3) Positive for 45,X if T score for chrom. X > 0.04 and for chrom. Y < 0.04.				
	In-house test.				
Target condition and reference stan-	Target conditions: T21, T18, T13 and 45,X.				
dard(s)	Reference standard: fetal	karyotype of chorionic villi,	amniotic fluid or placenta.		
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference stan-				
	dard). gNIPT was a second-tier test.				
	No failed sample reported.				
	No repeated test reported.				
Comparative					
Aim to study	To estimate the feasibility	of using a next-generation	sequencing technique for the non		
,	invasive prenatal diagnos				
Funding source or sponsor of the study	Funding source not repor	ted.			
Informations about the authors contact-		9 September and 4 Octobe	r 2016.		
ed	No reply received from th	e author.			
Notes					
Methodological quality					
Item	Authors' judgement	Risk of bias	Applicability concerns		
DOMAIN 1: Patient Selection					
Was a consecutive or random sample of patients enrolled?	No				
	Yes				



Sukhikh 2015 (Continued)

Did the study avoid inappropriate exclusions?

		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results interpreted without knowledge of the results of the reference standard?	Unclear		
If a threshold was used, was it pre-speci- fied?	Yes		
		Unclear	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to cor- rectly classify the target condition?	Yes		
Were the reference standard results inter- preted without knowledge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval be- tween index test and reference standard?	Yes		
Did all analysed patients receive the ref- erence standard?	Yes		
Were all patients included in the analysis?	Yes		
		Low	

Sung-Hee 2015

Study characteristics	
Patient sampling	Study design: retrospective study from a prospective cohort. Participants: pregnant women selected from a high risk of fetal aneuploidy population. Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 918 pregnant women. Number available for 2 x 2 table: 901 pregnant women (subgroup of 99%). Setting: various medical sites in Korea. Recruitment period: May 2012 to December 2013. Ethnicity: Asian. Mean gestational age (± SD; range): 16.6 (± 2.2; 11 to 25) weeks. Mean maternal age (± SD; range): 35.3 (± 4.1; 22 to 46) years.



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ung-Hee 2015 (Continued)			
	Relevant tests carried out pu biochemical screening (59% Language of the study: Engl) or both.	bhy (nuchal translucency measurement) or
Index tests	gNIPT by MPSS on Illumina	Genome Analyzer IIx or HiSeq 2	000 sequencer in 12-plex.
		re collected before reference s > 1 and t score > 2.5 (warning zo	tandard. one if t score risk > 2.5 or L score risk > 1).
	Biochemical serum-screenir	ng results were reported in the	study but 2 x 2 tables could not be derived.
Target condition and ref- erence standard(s)	SCA were also screened but	inappropriate reference standa	t the only case found was without follow-up ard for the present review was used. ses and medical record from birth for gNIPT
Flow and timing	Blood samples for gNIPT we gNIPT was a first- or a secon 8/918 samples were ineligib	d-tier test.	e procedure (reference standard).
	9/910 samples without follo had abortion and 7/9 wome		es had positive gNIPT result). 2/9 women
			rst blood samples including 1 haemolysed o cell-free DNA extraction failures and 9 sam
		d with new sampling. 14/16 sa formative results and were clas	mples obtained a gNIPT results and 2/16 sified as test failures.
		sequencing process (no gNIPT enced and 2/7 samples failed t	result). 5/7 samples failed the initial MPSS he second MPSS testing.
Comparative			
Aim to study		performance of gNIPT in detect ngleton pregnancies in Korea.	ing fetal chromosomal aneuploidies, espe-
Funding source or sponsor of the study		y but BGI performed sequencir rant (2015, President: Kyoung-I	ng and analysis. Study funded by Seoul Clini- Ryul Lee).
Informations about the authors contacted		3, 19 and 26 September 2016. 25 September and 11 October 2	2016.
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selectio	n		
Was a consecutive or ran- dom sample of patients enrolled?	No		
enrolled?			



Sung-Hee 2015 (Continued)

Did the study avoid inap-	No	
propriate exclusions?		

		High	Low	
DOMAIN 2: Index Test MPS	S			
Were the index test re- sults interpreted without knowledge of the results of the reference standard?	Yes			
f a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Stan	dard			
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference stan- dard results interpreted without knowledge of the results of the index tests?	No			
		High	Low	
DOMAIN 4: Flow and Timin	g			
Was there an appropriate interval between index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference stan- dard?	Yes			
Were all patients included in the analysis?	No			
		High		
ynan 2016				
Study characteristics				

ynan 2016 (Continued)			
Patient characteristics and set- ting	Number enrolled: 1100 preg Number available for 2 x 2 ta Setting: multicentre. Recruitment period: beginni Ethnicity: not reported. Gestational age (range): 9 to Maternal age (range): 18 to 4 Relevant tests carried out pr ment) or biochemical screer Language of the study: Engli	ble: 1048 pregnant women (ng in November 2009. 38.1 weeks. 5 years. ior to index test: ultrasonogr ing or both.	subgroup of 95%). raphy (nuchal translucency measure-
Index tests	gNIPT by MPSS on Illumina H	liSeq 2000 or HiSeq 2500 sec	quencer in multiplex.
	Mean fetal fraction DNA (± S 4.8%), and high-risk group (a		1%), high-risk group (< 35 years): 11.9% (±
	Median fetal fraction DNA (ra years): 10.7% (4.9% to 28.3% Blood samples for gNIPTwer Cutpoint: positive if risk scon Commercial test: VisibiliT™ t)), and high-risk group (≥ 35 y e collected before reference e ≥ 1%.	
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or medical record from birth.		
Flow and timing	gNIPT was a first- or second- 52/1100 samples failed durin	tier test. ng sequencing process (no gl or low aligned reads counts)	sive procedure (reference standard). NIPT result) including 28 for technical and 24 for discretionary non-reporting
	No repeated test reported.		
Comparative			
Aim to study	ibiliT™) that combines a mat	ernal age-based risk for T21 presentation of chromosome	low coverage, low cost MPSS assay (Vis- , T18, and T13, the fractional concentra- es 21, 18, and 13 in the sample to provide tal sex result.
Funding source or sponsor of the study	Study funded by Sequenom	Inc.	
Informations about the authors contacted	No need for further contact.		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		

	Cochrane
マノ	Library

ynan 2016 (Continued)				
Was a case-control design avoided?	Yes			
Did the study avoid inappropri- ate exclusions?	No			
		High	High	
DOMAIN 2: Index Test MPSS				
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre-specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards like- ly to correctly classify the target condition?	Yes			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Yes			
		Low	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate inter- val between index test and ref- erence standard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		

Verweij 2013

Study characteristics

Patient sampling	Study design: blinded, prospective cohort study. Participants: consecutive pregnant women selected at high risk of fetal aneuploidy presenting for invasive testing. Inclusion criteria: women who sign informed consent, ≥ 18 years old and carrying a singleton preg- nancy with a gestational age of at least 10 weeks.

sampling, history or active sign	ificant malignancy requiring majo	r surgery or systemic chemothera-
Number enrolled: 595 pregnan	t women.	
		of 85%).
		fro-European (1.3%), and other
(4.6%).		
		ichal translucency measurement)
Language of the study: English		
gNIPT by TMPS (DANSR assay)	on Illumina HiSeq 2000 in 96-plex v	with FORTE algorithm.
Blood samples for gNIPT were	collected just before reference star score > 1%	ndard.
	atype of chorionic villi (54%) or am	niotic fluid (46%)
	obtained just prior the invasive pro	ocedure (reference standard).
-		
-	-	
51/51 samples were repeated v tained a gNIPT results.	vith a second aliquot of the first sa	mpling and 35/51 samples ob-
Study funded by Ariosa Diagno	stics, Inc. 2 authors are paid emplo	oyees of Ariosa Daignostics. 1 au-
		,
Author was contacted on: 22 Au	aril 2016	
Authors' judgement	Risk of bias	Applicability concerns
Yes		
	sampling, history or active sign py, or language restriction with Number enrolled: 595 pregnam Number available for 2 x 2 table Setting: multicentres in the Net Recruitment period: May 2011 t Ethnicity: Caucasian (84.8%), M (4.6%). Mean gestational age (± SD; rang Relevant tests carried out prior or biochemical screening or bo Language of the study: English. gNIPT by TMPS (DANSR assay) of Mean fetal fraction DNA (± SD; r Blood samples for gNIPT were of Cutpoint: positive if FORTE risk Commercial test: Ariosa Diagno Target condition: T21. Reference standard: fetal karyo Blood samples for gNIPT were of gNIPT was a second-tier test. 75/595 samples were ineligible 51/520 samples failed the initia 51/51 samples were repeated w tained a gNIPT results. 16/520 samples failed during so fetal DNA fraction and 9 samples Study funded by Ariosa Diagno thor is a board member of Arios Author was contacted on: 22 Ag Reply received on: 25 April 2010 Authors' judgement	Mean gestational age (± SD; range): 14.0 (± 2.1; 10 to 28) weeks. Mean maternal age (± SD; range): 36.4 (± 4.6; 20 to 47) years. Relevant tests carried out prior to index test: ultrasonography (nu or biochemical screening or both. Language of the study: English. gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex to the study: English. gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex to the study: provide the study: English. gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex to the study: provide the study: English. gNIPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex to the study: English. gNode the study: English. gNiPT by TMPS (DANSR assay) on Illumina HiSeq 2000 in 96-plex to the study: English. gNode the study: English. gNode the study: English. gNode the study: English. Commercial test: Ariosa Diagnostics, Inc's test. Target condition: T21. Reference standard: fetal karyotype of chorionic villi (54%) or am Blood samples for gNIPT were obtained just prior the invasive program gNIPT was a second-tier test. 75/595 samples were ineligible. 51/51 samples were repeated with a second aliquot of the first sa tained a gNIPT results. 16/520 samples failed during sequencing process (no gNIPT result fetal DNA fraction and 9 samples from Europe to a laboratory in the Study fund



erweij 2013 (Continued)	N.		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test TMPS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard	1		
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	
/ang 2014			
Study characteristics			
Patient sampling		ign: prospective cohort study. hts: pregnant women in the first trimes	ter of pregnancy with advanced maternal

ages or ultrasound abnormality (high risk of fetal aneuploidy).

Inclusion criteria: singleton pregnancies between 11 to 14 weeks' gestation.

Nang 2014 (Continued)	Evolution critoria, multifut	al programation	
	Exclusion criteria: multifet	ai pregnancies.	
Patient characteristics and setting	 Number enrolled: 136 pregnant women. Number available for 2 x 2 table: 136 pregnant women (whole cohort included in analyse Setting: 1 centre. General Hospital of PLA, Beijing, China. Recruitment period: March 2011 to August 2013. Ethnicity: Asian. Gestational age range: 11 to 13.9 weeks. Maternal age range: 35 to 44 years. Relevant test carried out prior to index test: ultrasonography for some women. Language of the study: Chinese. 		
Index tests	gNIPT by MPSS on Illumin	a HiSeq 2000 sequencer with N	IIFTY™ algorithm.
	Fetal fraction DNA: not rep Blood samples for gNIPT v Cutpoint: not reported. Commercial test: BGI-Sher	vere collected before reference	e standard.
Target condition and reference standard(s)	Target conditions: T21 and T18. T13 was also assessed but no case was found. 45,X was also screened but inappropriate reference standard for the present review was used for pregnant women with gNIPT negative result. gNIPT data from 45,X were not shown in this review. Reference standards: fetal karyotype of amniotic fluid or cord blood or neonatal clinical examination at 42 days after birth or both.		
Flow and timing Blood samples for gNIPT were o gNIPT was a first- or second-tier No failed sample reported.		d-tier test.	sive procedure (reference standard).
	No repeated test reported		
Comparative			
Aim to study	To investigate the value of maternal plasma ccfDNA examination in detection of fetal chromo somal aneuploidy in pregnant women at advanced maternal age during the first trimester of pregnancy.		
Funding source or sponsor of the study	Study not funded by industry. Study funded by National Science & Technology Pillar Program during the Twelfth Five-year Plan Period (2012BA131B06).		
Informations about the authors con- tacted	Author was contacted on: 19 May and 27 June 2016. No replies received from the author.		
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	Unclear		
Was a case-control design avoided?	Yes		
Did the study avoid inappropriate exclusions?	No		



Wang 2014 (Continued)			
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre- specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condi-tion?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all analysed patients receive the reference standard?	Yes		
Were all patients included in the analysis?	Yes		
		Unclear	

Wang 2015a

Study characteristics	
Patient sampling	Study design: prospective cohort study. Participants: pregnant women selected from a high risk of fetal aneuploidy population. Inclusion criteria: pregnant women at high risk of fetal aneuploidy between 14 and 26 weeks o gestation. Exclusion criteria: not reported.
Patient characteristics and setting	Number enrolled: 917 pregnant women. Number available for 2 x 2 table: 917 pregnant women (whole cohort included in analyses). Setting: 1 centre at prenatal clinic, Lianyungang Maternal and Child Health Hospital, Lianyun- gang, Jiangsu 222001, China. Recruitment period: January 2012 to December 2013. Ethnicity: Asian. Gestational age range: 14 to 26 weeks.

Vang 2015a (Continued)			
	Maternal age range: 18 to 4 Relevant tests carried out J ment) or biochemical scree Language of the study: Eng	prior to index test: ultrasonog ening or both.	graphy (nuchal translucency measure-
Index tests	gNIPT by MPSS on Illumina	v2 HiSeq 2000 flow cell on a	HiSeq sequencer.
	Fetal fraction DNA: not rep Blood samples for gNIPT w Cutpoint:	orted. ere collected before referenc	ce standard.
	1) for T21, T18 and T13, po	sitive if Z score > 3.	
	2) for 47,XXY and 47,XYY, po	ositive if Z score Chrom. X > -3	3 and Z score Chrom. Y < 3.
	3) for 45,X and 47,XXX, pos tation. Commercial test: Berry Ge		een -3 and 3 without Chrom. Y represen-
Target condition and reference standard(s)	sessed but inappropriate r	eference standard for the pre	but no case was found. SCA was also as- esent review was used. or clinical follow-up (once per month)
Flow and timing	Blood samples for gNIPT w gNIPT was a second-tier te		asive procedure (reference standard).
	No failed sample reported.		
	No repeated test reported.		
Comparative			
Aim to study	To investigate the clinical e	fficiency of gNIPT identifying	g fetal chromosomal aneuploidies.
Funding source or sponsor of the study		lopment Fund, granted by th	td give technical support. Study fund- ne Department of Family Planning and
Informations about the authors contacted	No need for further contac	t.	
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sam- ple of patients enrolled?	No		
Was a case-control design avoid- ed?	Yes		
Did the study avoid inappropriate exclusions?	No		



Wang 2015a (Continued)				
		High	Low	
DOMAIN 2: Index Test MPSS				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear			
		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	Yes			
		Low		

Yao 2014

Study characteristics	
Patient sampling	Study design: retrospective study. Participants: pregnant women presenting with low-, high- or without prior risk factors of fetal aneu- ploidy (gNIPT was offered routinely as a prenatal screening test). Inclusion criteria: singleton pregnancies. Exclusion criteria: multifetal pregnancies.
Patient characteristics and setting	Number enrolled: 5950 pregnant women. Number available for 2 x 2 table: 5530 pregnant women (subgroup of 93%). Setting: 1 centre. The Prenatal Diagnosis Centre, Southwest Hospital, Chongqing, China. Recruitment period: June 2011 to December 2012. Ethnicity: Asian. Mean gestational age (range): 19.6 weeks (65% of the cohort were between 16 to 20.9 weeks).

Yao 2014 (Continued)	
	Mean maternal age (± SD): 30 (± 5) years. Relevant tests carried out prior to index test: ultrasonography (nuchal translucency measurement) or biochemical screening or both for some women. Language of the study: English.
Index tests	gNIPT by MPSS on Illumina Genome Analyzer IIx or HiSeq 2000 sequencer in 12-plex with NIFTY™ al- gorithm.
	Fetal fraction DNA: amount measured but not reported. Blood samples for gNIPT were collected before reference standard. Cutpoint:
	1) positive if t score \geq 2.5 for autosomes.
	2) positive if t score for Chrom. X < -2.5 for female fetuses for 45,X.
	3) positive if t score for Chrom. X > 2.5 for female fetuses for 47,XXX.
	4) positive if t score for Chrom. X > 2.5 combined with estimation of fetal ccfDNA concentration by Chrom. X (expected value of zero) for 47,XXY.
	5) positive if t score for Chrom. X > 2.5 and R-value (the ratio of the fetal DNA fraction estimated by chromosome X) between 1.8 and 2.2 for 47,XYY.
	Commercial test: BGI-Shenzhen's prenatal test.
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13. 45,X, 47,XXY, 47,XYY and 47,XXX were also screened but inappropri- ate reference standard for the present review was used. gNIPT data from SCA were not shown in this review. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or follow-up by telephone in- terview with the clinician after the expected delivery date.
Flow and timing	Blood samples for gNIPT were obtained prior to the invasive procedure (reference standard). gNIPT was a first- or second-tier test. 420/5950 samples without follow-up were excluded.
	No failed sample reported.
	No repeated test reported.
Comparative	
Aim to study	To evaluate the performance of a MPSS in detecting fetal sex chromosome aneuploidy (SCA) and to present a comprehensive clinical counselling protocol for SCA-positive patients. Author also assessed autosomes aneuploidies.
Funding source or sponsor of the study	Funding source not reported but many authors are employees of the Clinical Laboratory of BGI Health, BGI-Shenzen or of the Shenzen Birth Defect Screening Projet Lab.
Informations about the au- thors contacted	No need for further contact.
Notes	
Methodological quality	
Item	Authors' judgement Risk of bias Applicability concerns
DOMAIN 1: Patient Selection	



Yao 2014 (Continued)			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	No		
		High	High
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standar	rd		
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear		
		Unclear	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Yes		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	No		
		High	
Zhang 2016			
Study characteristics			
Patient sampling	Study design: blinded, prospect	ive cohort study.	



Zhang 2016 (Continued)			
	a soft marker in the genetic scan genetic sonography, not suitable munodeficiency virus infection, p tive blood type, a history of abor Exclusion criteria: multifetal pres	35 years at the time of delivery median, elevated fetal nuchal tr , or cardiac structural abnormal e for invasive prenatal diagnosis placenta previa, low-set place tion, threatened abortion or pre gnancies, maternal with chrome	r, single birth, high risk of T21 or ranslucency in the early pregnancy, lities in the second-trimester s, such as those with human im- centa, oligohydramnios, Rh-nega-
Patient characteristics and setting	Recruitment period: January 201 Ethnicity: Asian. Median gestational age (range): 1 Mean maternal age (± SD): 37.48	87 pregnant women (whole col is and Gynecology Hospital of Fu 12 to December 2013. 19 (12.4 to 32.5) weeks. (± 2.17) years. o index test: ultrasonography (n	hort included in analyses). udan University (Shanghai, China). nuchal translucency measurement)
Index tests	gNIPT by MPSS on Illummina His Fetal fraction DNA: not reported. It is not reported if the blood san Cutpoint for T21: positive if Z sco No other cutpoint reported. Commercial test: Berry Genomic	nples for gNIPT were collected b ore ≥ 3.	before or after reference standard.
Target condition and refer- ence standard(s)	Target conditions: T21, T18. 45,X dard for the present review was u	and 47,XXX were also screened used.	l but inappropriate reference stan- lood or neonatal clinical examina-
Flow and timing	It is not reported if the blood san standard). gNIPT was a second-tier test. No failed sample reported. No repeated test reported.	nples were collected before or a	after invasive procedure (reference
Comparative			
Aim to study	To evaluate the efficacy of using maternal age and to provide evic		
Funding source or sponsor of the study	Funding source not reported.		
Informations about the au- thors contacted	Author was contacted on: 7 Sept No reply received from the autho		
Notes			
Methodological quality			
ltem	Authors' judgement	Risk of bias	Applicability concerns



Zhang 2016 (Continued)

DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	No		
Was a case-control design avoided?	Yes		
Did the study avoid inappro- priate exclusions?	No		
		High	Low
DOMAIN 2: Index Test MPSS			
Were the index test results in- terpreted without knowledge of the results of the reference standard?	Yes		
If a threshold was used, was it pre-specified?	Yes		
		Low	Low
DOMAIN 3: Reference Standard	1		
Is the reference standards like- ly to correctly classify the tar- get condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
		Low	Low
DOMAIN 4: Flow and Timing			
Was there an appropriate in- terval between index test and reference standard?	Unclear		
Did all analysed patients re- ceive the reference standard?	Yes		
Were all patients included in the analysis?	Yes		
		Unclear	

Zhou 2014a

Study characteristics

hou 2014a (Continued)			
Patient sampling	Study design: blinded, prospect Participants: pregnant women s or without a priori risk (46.1%). Inclusion criteria: singleton preg Exclusion criteria: multifetal pre	elected at high risk for T gNIPT was integrated in o gnancies.	21 (51.3%), low risk for T21 (2.6%) Llinical workflow.
Patient characteristics and setting	Number enrolled: 306 pregnant	women.	
	of 98%). See Zhou 2014b for the Setting: 1 centre. Women's Hosp China. Recruitment period: September Ethnicity: Asian. Gestational age range: 12 to 24 w Maternal age: not reported.	integration set. bital, Zhejiang University 2011 to October 2011. weeks. to index test: ultrasonogr	n the pilot validation set (subgroup School of Medicine, Hangzhouin, raphy (nuchal translucency mea- phort.
Index tests	gNIPT by MPSS on Illumina Gen	ome Analyzer IIx or HiSec	2000 sequencer in 12-plex.
	Fetal fraction DNA: amount mea Blood samples for gNIPT were c Cutpoint: positive if T score > 2. Commercial test: NIFTY™ prena	ollected before reference 5 and L score > 1 (warning	g zone if t score > 2.5 or L score > 1)
Target condition and reference stan- dard(s)	Target conditions: T21, T18 and Reference standards: fetal karyo come.		neonatal karyotype or birth out-
Flow and timing	Blood samples were obtained p	rior to the invasive proce	dure (reference standard).
	gNIPT was a first- or second-tier For the pilot validation set: 5/30		v-up were excluded.
	No failed sample reported.		
	No repeated test reported.		
Comparative			
Aim to study	To report the clinical application of gNIPT to detect chromosomal aneuploidies, especially T21, T18 and T13 in Chinese singleton pregnancies.		
Funding source or sponsor of the study	Study not funded by industry but BGI-Shenzhen made sequencing and analysis. Some au- thors are employees of BGI-Shenzhen.		
Informations about the authors con- tacted	Author was contacted on: 31 Ma No reply received from author.	y 2016.	
Notes			
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			



Zhou 2014a (Continued)				
Was a consecutive or random sample of patients enrolled?	No			
Was a case-control design avoided?	Yes			
Did the study avoid inappropriate ex- clusions?	No			
		High	High	
DOMAIN 2: Index Test MPSS				
Were the index test results interpret- ed without knowledge of the results of the reference standard?	Yes			
If a threshold was used, was it pre- specified?	Yes			
		Low	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	Unclear			
		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval be- tween index test and reference stan- dard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		
Zhou 2014b				
Study characteristics				
Patient sampling	Study design: blinded, pro Participants: pregnant wo gNIPT was integrated in cl	men selected at high risk, lov	risk for T21 or without a pric	ri risk.

Inclusion criteria: singleton pregnancies.



hou 2014b (Continued)	Exclusion criteria: multifeta	l pregnancies.		
Patient characteristics and setting	Number enrolled: 7705 pre	gnant women.		
		ables: 3950 pregnant women	in the integration set (subgroup of	
	-	Hospital, Zhejiang University	School of Medicine, Hangzhouin, Chi-	
	na. Recruitment period: Septer	nber 2011 to July 2013.		
	Ethnicity: Asian.	24 wooks		
	Gestational age range: 12 to Maternal age: not reported.			
			raphy (nuchal translucency measure-	
	Language of the study: Eng	eening for a part of this cohor ish.	τ.	
Index tests	gNIPT by MPSS on Illumina	Genome Analyzer IIx or HiSed	q 2000 sequencer in 12-plex.	
		measured but not reported.		
		ere collected before reference		
		enatal test by BGI-Shenzhen	g zone if t score > 2.5 or L score > 1).	
Target condition and reference	Target conditions: T21, T18			
standard(s)	Reference standards: fetal l	caryotype of amniotic fluid or	r neonatal karyotype or birth outcome.	
Flow and timing	Blood samples were obtained prior to the invasive procedure (reference standard).			
	gNIPT was a first- or second-tier test.			
	141/7705 samples failed the initial MPSS testing. 141/141 samples were repeated with a new sampling and 137/141 samples obtained a gNIPT results.			
	4/7705 samples failed the s	econd MPSS testing for low fe	etal fraction DNA (no gNIPT result).	
	3751/7701 samples without	birth outcome were exclude	ed (no reference standard).	
Comparative				
Aim to study	To report the clinical applic T18 and T13 in Chinese sing		mosomal aneuploidies, especially T21	
Funding source or sponsor of the study	Study not funded by indust are employees of BGI-Shen		equencing and analysis. Some authors	
Informations about the authors contacted	Author was contacted on: 3 No reply received from the			
Notes				
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sam- ple of patients enrolled?	No			



Zhou 2014b (Continued)				
Was a case-control design avoid- ed?	Yes			
Did the study avoid inappropriate exclusions?	No			
		High	High	
DOMAIN 2: Index Test MPSS				
Were the index test results inter- preted without knowledge of the results of the reference standard?	Unclear			
If a threshold was used, was it pre- specified?	Yes			
		Unclear	Low	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard re- sults interpreted without knowl- edge of the results of the index tests?	Unclear			
		Unclear	Low	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all analysed patients receive the reference standard?	Yes			
Were all patients included in the analysis?	No			
		High		
CVS: chorionic villi sampling DANSR: digital analysis of selected reg FISH: fluorescence in situ hybridisatio gNIPT: genomics-based non-invasive MAD: Median absolute deviation MPSS: massively parallel shotgun seq NCV: normalised chromosome value SD: standard deviation SNP: single nucleotide polymorphism	n prenatal testing uencing			

TMPS: targeted massively parallel sequencing



Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Anderson 2015	Not a diagnostic test accuracy study. Poster abstract.
Anselem 2016	Decision making study. Observational study. Not a diagnostic test accuracy study.
Bayindir 2015	Samples overlap with Brady 2016. Most gNIPT results unconfirmed by a reference standard test. In- sufficient information to derive 2 x 2 tables.
Beamon 2013	Poster abstract of the 33 rd Annual Meeting of the Society for Maternal-Fetal Medicine: The Pregnan- cy Meeting. Observational study with incomplete follow-up. Samples overlap with Beamon 2014.
Beamon 2014	Observational study with incomplete follow-up. Not a diagnostic test accuracy study. Some gNIPT results unconfirmed by a reference standard test.
Belloin 2016	Most women (95%) completed a questionnaire to report their birth outcome (inappropriate reference standard for this review).
Benachi 2015b	Tribune. Not a diagnostic test accuracy study.
Benachi 2016	All samples overlap with Benachi 2015.
Benn 2015	Letter to the editor on Zhang 2015 without data. Not a diagnostic test accuracy study.
Bhatt 2014	Poster abstract of the 18 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2014. Patients with gNIPT negative result were without follow-up (no reference standard). Incomplete 2 x 2 tables.
Bianchi 2012a	Samples overlap with Bianchi 2012. Data excluded to avoid double counting.
Bianchi 2014b	Editorial. Not a diagnostic test accuracy study.
Bianchi 2014c	Data excluded to avoid double counting. Samples overlap with Bianchi 2015b.
Bianchi 2015a	Not a diagnostic test accuracy study. Author presented some false positive cases in women with malignancy.
Bianchi 2015b	Incomplete 2 x 2 table. In this observational study, most women (98.9%) had no follow-up (no refer- ence standard).
Bianchi 2015c	Not a diagnostic test accuracy study. Author presented some false positive cases in women with malignancy. Samples overlap with Bianchi 2015a.
Bianchi 2015d	Poster abstract. Not a diagnostic test accuracy study. Author presented some false positive cases in women with malignancy. Samples overlap with Bianchi 2015a.
BlueCross BlueShield Asssoci- ation 2014	Technology Evaluation Center Assessment. Review.
Brady 2016	Review with new data but most gNIPT results unconfirmed by a reference standard test. Insuffi- cient information to derive 2 x 2 tables.
Chen 2013	Poster abstract of the ISPD 17th International Conference on Prenatal Diagnosis and Therapy. Samples overlap with Huang 2014.



Study	Reason for exclusion
Chen 2014	Poster abstract. Samples overlap with Yeang 2014.
Cherry 2014	Poster abstract. Samples overlap with Meck 2015.
Cheung 2015	Incomplete 2 x 2 table. This letter presented women who had positive results after screening and were referred for invasive procedure to confirm the presence of fetal aneuploidy. Only, the true positive and false positive gNIPT results were reported.
Chiu 2008	Proof-of-concept. Not a diagnostic test accuracy study.
Chiu 2010	Proof-of-concept. Not a diagnostic test accuracy study.
Christina 2012	Proof-of-concept. Not a diagnostic test accuracy study.
Cinnioglu 2012	Poster abstract. Samples overlap with Rabinowitz 2012a.
Cirigliano 2013	Full poster from the 17 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2013 received. Samples overlap with Cirigliano 2014 and Ordoñez 2015. Insufficient information to derive 2 x 2 tables.
Cirigliano 2014	Full poster from the 18 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2014 received. Samples overlap with Cirigliano 2013 and Ordoñez 2015. Insufficient information to derive 2 x 2 tables.
Cuckle 2015	Review with simulation model for gNIPT.
Curnow 2014	Poster abstract of the 18 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2014. Samples overlap with Dar 2014.
Dan 2012	Incomplete 2 x 2 table. Women with gNIPT negative results completed a questionnaire to report their birth outcome (inappropriate reference standard for this review).
Dar 2014	Implementation study. Incomplete 2 x 2 table. Most patients with gNIPT negative result were with- out follow-up (no reference standard). Some women had follow-up by telephone (inappropriate reference standard for this review).
De Ligt 2013	Case report (deletion).
Denona 2016	Poster abstract. Retrospective observational study. Insufficient information to derive 2 x 2 tables.
Discenza 2015	Poster abstract. Some gNIPT results unconfirmed by a reference standard test. Insufficient informa- tion to derive 2 x 2 tables.
Dobson 2015	Poster abstract. Insufficient information to derive 2 x 2 tables (gNIPT positive results only). Decision making. Samples overlap with Dobson 2016.
Dobson 2016	Insufficient information to derive 2 x 2 tables (gNIPT positive results only). Decision making.
Dong 2016	Sequencing not based on maternal plasma ccfDNA.
Duenwald 2016	Method development. Analytical accuracy. Not a diagnostic test accuracy study.
Ehrich 2011a	Editorial comment without new data.
Eiben 2014	Review. Not a diagnostic test accuracy study.

Study	Reason for exclusion
Ellison 2015	Poster abstract. All gNIPT results (TMPS) were confirmed with a previous gNIPT result (MPSS) (inap- propriate reference standard for this review). Insufficient information to derive 2 x 2 tables.
Faas 2011	Poster abstract of the 8 th European Cytogenetics Conference. Samples overlap with Faas 2012.
Faas 2012	Proof-of-concept. Not a diagnostic test accuracy study.
Fairbrother 2013a	Observational study. Incomplete 2 x 2 data. Most patients were without follow-up (no reference standard).
Fairbrother 2013b	Conference abstract of the 17 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2013. Samples overlap with Fairbrother 2013a.
Fan 2008	Proof-of-concept. Not a diagnostic test accuracy study.
Fang 2015	Insufficient information to derive 2 x 2 tables.
Ferres 2013	Not a diagnostic test accuracy study (implementation study).
Fiorentino 2015	Poster abstract. All samples overlap with Fiorentino 2016.
Fosler 2015	Poster abstract of the 35 th Annual Meeting of the Society for Maternal-Fetal Medicine: The Pregnan- cy Meeting. Observational study. Incomplete 2 x 2 table. Most patients with gNIPT negative result were without follow-up (no reference standard).
Futch 2013	Observational study with incomplete follow-up. Incomplete 2 x 2 table. Many gNIPT results uncon- firmed by a reference standard test.
Gabriel 2014	Conference abstract. Proof-of-concept.
Galea 2014	Full poster from the 18 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2014. Incomplete 2 x 2 table. Most patients with gNIPT negative result were without follow-up (no refer- ence standard).
Gao 2014	News, comment on Liao 2014 and Yuan 2013 without new data.
Gao 2015	Poster abstract. Insufficient information to derive 2 x 2 tables.
Geifman-Holtzman 2013	Poster abstract of the 33 rd Annual Meeting of the Society for Maternal-Fetal Medicine: The Pregnan- cy Meeting. Samples overlap with Xiong 2015.
Geifman-Holtzman 2014	Poster abstract of the 34 th Annual Meeting of the Society for Maternal-Fetal Medicine: The Pregnan- cy Meeting. Samples overlap with Xiong 2015.
Gerundino 2017	Women were asked to complete a questionnaire to report their birth outcome (inappropriate refer- ence standard for this review). Insufficient information to derive 2 x 2 tables.
Gil 2013	Most patients with gNITP negative result were without follow-up (no reference standard) because 962 women had not yet delivered at the time of writing the publication. Insufficient information to derive 2 x 2 tables. Some patients overlap with del Mar Gil 2014.
Gil 2015	Decision making including gNIPT accuracy data. All samples overlap with Gil 2016.
Gnetetskaya 2015	Poster abstract. Samples overlap with Kurtser 2015.



Study	Reason for exclusion
Grati 2014	Not a diagnostic test accuracy study. No sequencing data.
Gray 2013	Observational study. Not a diagnostic test accuracy study. Full poster received from the authors. Poster of the 17 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2013.
Gromminger 2014	Data excluded to avoid double counting. Blinded DNA sequencing libraries were provided by Se- quenom from their clinical trial cohort (NCT00877292) and were resequenced by LifeCodexx.
Guex 2013	Research letter. Samples overlap with Pescia 2017.
Halks-Miller 2015	In reply to Bianchi 2015a. Not a diagnostic test accuracy study.
Harasim 2016	Poster abstract. Insufficient information to derive 2 x 2 tables.
Hernandez-Gomez 2015	Implementation study. Not a diagnostic test accuracy study.
Hofmann 2013	Poster abstract. Samples overlap with Stumm 2014.
Hofmann 2014	Conference abstract. Insufficient information to derive 2 x 2 tables.
Hofmann 2015	Method development. Data were reanalysed by a new algorithmic approach of PraenaTest®. Not a diagnostic test accuracy study.
Hu 2014	Not a next generation sequencing publication. NIPT was ultrasound and serum biomarkers.
Hu 2015	Incomplete 2 x 2 tables. Only gNIPT positive results presented.
Hui 2015a	Poster abstract. All samples overlap with Hui 2015b.
Hui 2015b	Implementation study. Incomplete 2 x 2 table. Most patients with gNIPT negative result were with- out follow-up (no reference standard).
Jackson 2013	Poster abstract. All samples overlap with Jackson 2014.
Jensen 2013	Proof-of-concept. Not a diagnostic test accuracy study. Samples overlap with Palomaki 2012.
Jensen 2015	Proof-of-concept study with unblinded samples. Not a diagnostic test accuracy study.
Jin 2014	Incomplete 2 x 2 table. Women with gNIPT negative results were followed-up by telephone (inap- propriate reference standard for this review).
Johnson 2013	Not a next generation sequencing method.
Juneau 2014	Method development. Incomplete 2 x 2 table. Most patients were without follow-up (no reference standard).
Kagan 2015	Not a diagnostic test accuracy study. Simulation model.
Kalantar 2014	Not next generation sequencing method.
Karlsson 2015	Methodological publication. Not a diagnostic test accuracy study.
Kershberg 2015	Poster abstract. Some gNIPT results unconfirmed by a reference standard test. Insufficient information to derive 2 x 2 tables.



Study	Reason for exclusion
Kinde 2012	Methodological publication. Not a diagnostic test accuracy study.
Korabecna 2012	Bioinformatic simulation with Palomaki 2011 data.
Koumbaris 2016	Method development (proof-of-concept study). Development of an advanced fetal fraction estima- tion method and aneuploidy determination algorithm. Not a diagnostic test accuracy study.
Kurtser 2015	Most patients with gNIPT negative result were without follow-up (no reference standard). Incom- plete 2 x 2 tables.
Lambert-Messerlian 2014	Samples overlap with Palomaki 2011 and Palomaki 2012. Data excluded to avoid double counting.
Larion 2015	Poster abstract. Implementation study.
Lau 2012a	Incomplete 2 x 2 table. Women with gNIPT negative results were followed up by telephone or by email (inappropriate reference standard for this review). All samples overlap with Lau 2014.
Lau 2013	Sample overlap with Lau 2014.
Lau 2014	Incomplete 2 x 2 table. Women with gNIPT negative results were followed up by telephone or by email (inappropriate reference standard for this review).
Lebo 2015	Incomplete 2 x 2 table.
Leung 2013	Proof-of-concept. Not a diagnostic test accuracy study.
Levandoski 2015	Poster abstract. Observational study about discordant gNIPT results. Insufficient information to de- rive 2 x 2 tables.
Levy 2013	Poster abstract. Incomplete 2 x 2 table.
Levy 2013a	Poster abstract. Incomplete 2 x 2 table.
Levy 2013b	Proof-of-concept. Not a diagnostic test accuracy study. Samples overlap with Zimmermann 2013.
Li 2012	Methodological publication about relation between fetal fraction and multiple clinical factors.
Li 2015	Observational study. Unavailable information about gNIPT approach used. It is unclear if patients with gNIPT negative result were followed up (no reference standard).
Liao 2011	Not a diagnostic test accuracy study. No aneuploid case.
Liao 2012	Proof-of-concept. Not a diagnostic test accuracy study.
Liao 2013	This is a poster abstract. The full publication was also excluded. See Liao 2014 for reasons of exclusion.
Liao 2014	Incomplete 2 x 2 table for the retrospective and the prospective cohort. In prospective cohort, most patients were without follow-up (no reference standard). For the retrospective cohort, number of gNIPT results was not reported. Sensitivity and specificity were presented for the retrospective cohort but 2 x 2 tables could not be derived.
Liao 2014a	Letter to the editor about Bianchi 2014b without new data.
Liu 2015	Incomplete 2 x 2 table. Women with gNIPT negative results were followed up by telephone (inap- propriate reference standard for this review).

Study	Reason for exclusion
Lo 2014	Bioinformatic development. Comparison of sensitivity and specificity using 3 different count nor- malisation methods.
Lo 2014a	Poster abstract. Bioinformatic development. Comparison of sensitivity and specificity using 3 different count normalisation methods. Samples overlap with Lo 2014.
Loucký 2013	Samples overlap with Palomaki 2012. Data excluded to avoid double counting.
Louis-Jacques 2014	Full poster. Observational study. Not a diagnostic test accuracy study.
Ma 2015	Samples overlap with Ma 2016.
Ma 2015a	Poster abstract. All samples overlap with Ma 2016.
Manotaya 2016	Insufficient information to derive 2 x 2 tables. Women without invasive testing results were encour- aged to report birth outcomes through the insurance policy reimbursed (inappropriate reference standard for this review).
Marchili 2015	Poster abstract. Implementation study. Not a diagnostic test accuracy study. Insufficient informa- tion to derive 2 x 2 tables.
Mayen 2015	Observational study. Not a diagnostic test accuracy study.
Mazloom 2013a	Poster abstract. Samples overlap with Mazloom 2013.
McCullough 2014	Incomplete 2 x 2 table. The clinician of women who passed gNIPT was encouraged to send ad hoc feedback to the lab (inappropriate reference standard for this review).
McCullough 2014a	Poster abstract. Incomplete 2 x 2 table. Samples overlap with McCullough 2014.
McCullough 2015	Full poster of the 19 th International Conference on Prenatal Diagnosis and Therapy, ISPD 2015. Most gNIPT results unconfirmed by a reference standard test. Insufficient information to derive 2 × 2 tables. Some patients overlap with McCullough 2014.
McLennan 2016	Most patients with gNIPT negative result were without follow-up (no reference standard). Insuffi- cient information to derive 2 x 2 tables.
Meck 2014	Poster abstract. Samples overlap with Meck 2015.
Meck 2015	Not a diagnostic test accuracy study. Observational study.
Meck 2015a	Poster abstract. Samples overlap with Meck 2015.
Mennuti 2015	Review without original data.
Minarik 2015	gNIPT negative results unconfirmed by a reference standard test. Not a diagnostic test accuracy study.
Miron 2011	Not a diagnostic test accuracy study. This these explore traditional screening tests.
Mundy 2008	Health Technology Assessment. Not a diagnostic test accuracy study.
Mundy 2009	Health Technology Assessment. Not a diagnostic test accuracy study.
Musci 2014	Poster abstract. Samples overlap with Norton 2015.



Study	Reason for exclusion
Musci 2014a	Poster abstract. Samples overlap with Hooks 2014 and Nicolaides 2014a.
NCT00770458	Not a gNIPT method (other method).
NCT00877292	Not with ccfDNA (other sampling).
NCT00891852	Not a gNIPT method (other method).
NCT00971334	Completed clinical trial but no published data.
NCT01052688	Incomplete 2 x 2 data (ongoing study with cases only).
NCT01256606	Not a gNIPT method (other method).
NCT01451671	Incomplete 2 x 2 data (ongoing study with cases only).
NCT01451684	Observational study on gNIPT without fetal karyotype.
NCT01555346	Completed clinical trial but no published data.
NCT01574781	Completed clinical trial but no published data.
NCT01597063	Completed clinical trial but no published data.
NCT01661010	Not a diagnostic test accuracy study.
NCT01663675	Adult with T21. Not with pregnant women (other population).
NCT01668251	Not a diagnostic test accuracy study.
NCT01725438	Not with ccfDNA (other sampling).
NCT01837979	Incomplete 2 x 2 data.
NCT01966991	Completed clinical trial but no published data.
NCT02127515	Not a diagnostic test accuracy study. Pregnant women with gNIPT have not a reference standard.
NCT02226315	Inappropriate reference standard for this review (pregnancy outcome data obtained from the pa- tient).
NCT02872948	Not a gNIPT method (other method).
Neufeld-Kaiser 2015	Observational study. Not a diagnostic test accuracy study. Incomplete 2 x 2 tables. Most gNIPT re- sults unconfirmed by a reference standard test.
Neveling 2015	Method validation for the NextSeq 500 platform. Not a diagnostic test accuracy study.
Nickolich 2016	Not a diagnostic test accuracy study.
Nicolaides 2013a	Poster abstract. All samples overlap with Nicolaides 2012.
Nicolaides 2014	Simulation model on gNIPT implantation in first- or second-tier test.
Nicolaides 2014b	Note on Nicolaides 2014a without new data.



Study	Reason for exclusion
Nicolaides 2014c	Target condition presented in this publication is not the focus of this review. Publication of next generation sequencing with ccfDNA for fetal triploidy.
Norem 2015	Full poster received from authors. Most patients were without follow-up (no reference standard). Incomplete 2 x 2 tables.
Norton 2014	Bioinformatic simulation.
Norton 2014a	Poster abstract. Samples overlap with Norton 2015.
Norton 2015a	Bioinformatic simulation.
Norton 2015b	Editorial on Norton 2015 without new data.
Norton 2015c	Author reply to comments from Sentilhes 2015 and Smith-Bindman 2015 about Norton 2015 with- out new data.
Norton 2016	Simulation model to compare sequential and ccfDNA screening with data published in the litera- ture. Not a diagnostic test accuracy study.
O'Leary 2014	Bioinformatic simulation.
Oepkes 2015	Most patients were without follow-up (no reference standard). Insufficient information to derive 2 x 2 tables.
Oneda 2016	Poster abstract. Insufficient information to derive 2 x 2 tables.
Ordoñez 2015	Full poster received. Some gNIPT results unconfirmed by a reference standard test. Insufficient in- formation to derive 2 x 2 tables.
Palomaki 2011	Samples overlap with Palomaki 2012 (samples in Palomaki 2011 have been reanalysed in Palomaki 2012). Study excluded to avoid double counting.
Palomaki 2012a	Samples overlap with Palomaki 2012. Conference abstract about Palomaki 2012 data.
Palomaki 2012b	Editorial on Palomaki 2011 without new data.
Palomaki 2015	Not a diagnostic test accuracy study.
Palomaki 2015a	Note about Palomaki 2015. Not a diagnostic test accuracy study.
Perez-Pedregosa 2015	Incomplete 2 x 2 tables. Some women with gNIPT negative results were followed up by telephone (inappropriate reference standard for this review).
Pescia 2017	Follow-up for gNIPT negative results was ensured by an inquiry of two sets of randomly selected samples (inappropriate reference standard for this review).
Petersen 2014	Not a next generation sequencing publication. NIPT was ultrasound measurement and serum bio- markers.
Pettit 2014	Most patients with gNIPT negative result were without follow-up (no reference standard). Insufficient information to derive 2 x 2 tables.
Porreco 2014a	Reply to Grati 2014 without sequencing data.
Rabinowitz 2012	Poster abstract. Proof-of-concept. Not a diagnostic test accuracy study.



Study	Reason for exclusion
Rabinowitz 2012a	Poster abstract with incomplete 2 x 2 tables.
Rabinowitz 2012b	Poster abstract. Samples overlap with Rabinowitz 2012a.
Rabinowitz 2013	Poster abstract. Samples overlap with Pergament 2014.
Rabinowitz 2014	Poster abstract. Sample overlap with Pergament 2014.
Rad 2014	Implementation study without sequencing data presented.
Radoi 2015	Incomplete 2 x 2 tables. Most patients were without follow-up (no reference standard).
Rava 2012	Poster abstract. Samples overlap with Bianchi 2012.
Rava 2014	Methodological publication about fetal DNA fraction with MELISSA samples.
Reiff 2015	Insufficient information to derive 2 x 2 tables.
Reiff 2016	Insufficient information to derive 2 x 2 tables.
Reimers 2015	Conference abstract from the 19 th International Conference on Prenatal Diagnosis and Theraphy, ISPD 2015. Simulation model. Not a diagnostic test accuracy study.
Revello 2016	Not a diagnostic test accuracy study. All samples overlap with Gil 2016 and Quezada 2015.
Ryan 2016	Method development of version 2 to SNP-based gNIPT. Not a diagnostic test accuracy study.
Sachse 2015	Proof-of-concept of fetal fraction quantification by qPCR.
Samura 2015	Most patients were without follow-up (no reference standard). Insufficient information to derive 2 x 2 tables. Samples overlap with Sago 2015.
Sarno 2016	Some women reported their birth outcome (inappropriate reference standard for this review). In- formation about false positive results were insufficient to derive all 2 x 2 tables.
Schöck 2015	Poster abstract. Bioinformatics development with unblinded samples.
Sehnert 2013	Poster abstract. Incomplete 2 x 2 table.
Sehnert 2014	Poster abstract. Samples overlap with Bianchi 2014b.
Sentilhes 2015	Comment about Norton 2015 without new data.
Seo 2015	Women with gNIPT result were without follow-up at birth.
Settler 2015	Full poster received. Insufficient information to derive 2 x 2 tables. Some gNIPT results uncon- firmed by a reference standard test.
Shani 2016	Simulation model. Not a diagnostic test accuracy study.
Shaohua 2012	Poster abstract. Full poster not received. Incomplete 2 x 2 table.
Sharma 2015	Poster abstract about patient perceptions of gNIPT from the multi-centered Canadian PEGASUS tri- al. gNIPT results compared with first trimester combined test (inappropriate reference standard for this review).



Study	Reason for exclusion
Shaw 2013	Poster abstract. Samples overlap with Shaw 2014.
Shen 2016	Method development. Not a diagnostic test accuracy study.
Shi 2015	Incomplete 2 x 2 table. gNIPT negative result unconfirmed by a reference standard test.
Shulman 2014	Poster abstract. Incomplete 2 x 2 table. Most patients with gNIPT negative result were without a reference standard test.
Sistermans 2015a	Letter to the editor on Bianchi 2015a without data.
Smith-Bindman 2015	Comment about Norton 2015 without new data.
Song 2012	Poster abstract. Some samples overlap with Sparks 2012a.
Sparks 2012	Method development (all unblinded samples). Incomplete 2 x 2 table. Most patients with gNIPT negative result were unconfirmed by a reference standard test.
Srinivasan 2013	Poster abstract. Samples from MELISSA study (potentially overlap).
Stokowski 2015	Not a next generation sequencing method.
Strah 2015	Women were followed up by telephone interview to find out their birth outcome (inappropriate reference standard for this review).
Straver 2014	Proof-of-concept.
Strom 2015	Incomplete 2 x 2 table. Only women with gNIPT positive result were reported.
Stumm 2011	Proof-of-concept. Not a diagnostic test accuracy study.
Stumm 2012	Proof-of-concept. Not a diagnostic test accuracy study.
Stumm 2012a	Poster abstract. Samples overlap with Stumm 2014.
Stumm 2013	Poster abstract. Samples overlap with Stumm 2014.
Stumm 2016	Not a diagnostic test accuracy study.
Swanson 2012	Publication about Bianchi 2012 without new data.
Syngelaki 2014	Not a diagnostic test accuracy study. Simulation model.
Tan 2016	Women with gNIPT negative results were followed up by telephone interview (inappropriate refer- ence standard for this review). Insufficient information to derive 2 x 2 tables.
Taneja 2016	Incomplete follow-up. Incomplete 2 x 2 table. Many gNIPT results unconfirmed by a reference stan- dard test.
Taneja 2017	Most patients with gNIPT negative result were without reference standard test. Providers were en- couraged to report discordant clinical outcomes. Insufficient information to derive 2 x 2 tables.
Tarrier 2015	gNIPT results unconfirmed by a reference standard test. Their reference method is verifi® results (inappropriate reference standard for this review).

Study	Reason for exclusion
Taylor 2014	Not a diagnostic test accuracy study. Observational study and decision making about gNIPT uptake in their center.
Togneri 2016	Full poster received. Internal verification set and implantation in their centre. Not a diagnostic test accuracy study.
Tong 2016	Not a next-generation sequencing method with ccfDNA.
Valderramos 2016a	Poster abstract. Insufficient information to derive 2 x 2 tables.
Valderramos 2016b	Poster abstract. Samples overlap with Valderramos 2016c.
Valderramos 2016c	Insufficient information to derive 2 x 2 tables. Retrospective cohort of patients with gNIPT positive results.
van den Oever 2012a	Proof-of-concept. Not a diagnostic test accuracy study.
van den Oever 2012b	Proof-of-concept. Not a diagnostic test accuracy study.
van den Oever 2013	Proof-of-concept. Not a diagnostic test accuracy study.
Van Opstal 2016	Simulation model. Not a diagnostic test accuracy study.
Verweij 2013a	Poster abstract. All samples overlap with Verweij 2013.
Wald 2015a	Not a diagnostic test accuracy study. Prenatal screening workflow proposed.
Wald 2015b	Not a diagnostic test accuracy study. Prenatal screening workflow proposed.
Wang 2012	Incomplete 2 x 2 table. Women with gNIPT negative results were followed up by telephone (inap- propriate reference standard for this review).
Wang 2015b	Not a diagnostic test accuracy study.
Wang 2015c	Proof-of-concept. Not a diagnostic test accuracy study.
Wang 2015d	Editorial on Wang 2015b without new data.
Wang 2015e	Not a diagnostic test accuracy study.
Xiong 2015	Full poster received. Observational study and incomplete follow-up.
Yankova 2015	Simulation model for gNIPT implantation. Not a diagnostic test accuracy study.
Yaron 2015	Commentary about gNIPT for microdeletion syndromes and rare autosomal trisomies. Not a diag- nostic test accuracy study.
Yeang 2014	Proof-of-concept. Not a diagnostic test accuracy study.
Yu 2014	Proof-of-concept. Not a diagnostic test accuracy study.
Yuan 2013	Proof-of-concept. Not a diagnostic test accuracy study.
Zhang 2015	Incomplete 2 x 2 table. Women with gNIPT negative results were followed-up by telephone (inap- propriate reference standard for this review).

Study	Reason for exclusion
Zhou 2013	Poster abstract. Incomplete 2 x 2 table.
Zimmermann 2012	Proof-of-concept. Not a diagnostic test accuracy study.
Zimmermann 2013	Proof-of-concept. Not a diagnostic test accuracy study.
Zwiefelhofer 2013	Implementation assessment of 2 sequencing platforms for gNIPT in a routine clinical environment. Not a diagnostic test accuracy study.

ccfDNA: circulating cell-free DNA gNIPT: genomics-based non-invasive prenatal testing MPSS: massively parallel shotgun sequencing TMPS: targeted massively parallel sequencing

Characteristics of ongoing studies [author-defined order]

Basaran 2015 Trial name or title Publication's title: False positive and false negative results of cell free DNA testing. Target condition and refer-Target conditions: T21, T18, T13, 45,X, 47,XXY, 47,XYY and 47,XXX. ence standard(s) Reference standard: fetal karyotype of chorionic villi or amniotic fluid. Index and comparator tests gNIPT by TMPS or MPSS by commercial company providing gNIPT in Turkey (Ariosa Diagnostics, Inc., BGI-Shenzhen, Illumina, Inc, Natera, Inc. and Sequenom, Inc). Blood samples for gNIPT were collected before reference standard. Starting date Not reported. Contact information Dr Seher Basaran **Department of Medical Genetics** Istanbul University, Istanbul Medical Faculty TURKEY 90 (212) 4142000 basarabs@istanbul.edu.tr Aim to study To demonstrate the importance of confirmation of fetus genotype by invasive testing after gNIPT. Funding source or sponsor of The genetic centre is not affiliated with any commercial company providing gNIPT. the study Information about the authors Author was contacted on: 12, 14 and 18 January 2016. contacted Last reply received on: 19 January 2016. Notes At the time of this writing, the authors plan to publish a full publication soon.

Buresch 2016

Trial name or title	Poster's title: Actual rates of recommended diagnostic testing after first-trimester screening vs same-day screening by cell free DNA.
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13, 45,X, 47,XXY, 47,XYY and 47,XXX. Reference standard: not reported.
Index and comparator tests	MPS.
Starting date	January to June 2015.
Contact information	Susan Klugman
	Department of Obstetrics and Gynecology and Women's Health,
	Albert Einstein College of Medicine,
	Montefiore Medical Center 1695 Eastchester Road,
	Bronx, NY 10461, United States.
	sklugman@montefiore.org
Aim to study	To compare actual patient referrals for post-screen diagnostic tests following first-trimester screen- ing vs same day ccfDNA.
Funding source or sponsor of the study	Not reported.
Information about the authors contacted	Author was contacted on: 1 and 23 September 2016. Reply received on: 23 September 2016.
Notes	Authors are working on data at the time of writing and they plan to submit for publication.

Chen 2011a

chen zorra	
Trial name or title	Oral presentation's title: Noninvasive prenatal diagnosis of fetal aneuploidy by massively parallel sequencing of maternal plasma DNA.
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13 and SCA. Reference standard: fetal karyotype.
Index and comparator tests	gNIPT by MPSS on Illumina GAIIx/HiSeq 2000 sequencer. Cutpoint: positive if t score < -4. Commercial test: BGI's test.
Starting date	Not reported.
Contact information	Fang Chen, Beijing Genomics Institute, Shenzhen, China
Aim to study	To assess gNIPT with ccfDNA performance on fetal aneuploidies.
Funding source or sponsor of the study	Not reported.
Information about the authors contacted	BGI was contacted on: 19 May 2016. No reply received from the author.



Chen 2011a (Continued)

Notes

Cohort of 5268 pregnant women. They successfully identified 62 cases of T21, 40 cases of T18, 3 cases of T13, 13 cases of SCA. In a cohort of karyotyping cases, the sensitivity and specificity of the aneuploidy fetus detection was 100% and 100%, respectively.

Da Fonseca 2015	
Trial name or title	Abstract's title: Non-Invasive prenatal testing for the most common aneuploidies (trisomies 21, 18, and 13) using a semiconductor-sequencing platform: a French multicenter pilot study.
Target condition and refer- ence standard(s)	Target conditions: T21, T18 and T13. Reference standard: fetal karyotype.
Index and comparator tests	gNIPT on semiconductor sequencing platform (MPSS). Blood sample collection not reported. Cutpoint: not reported.
Starting date	Not reported.
Contact information	J.P. Da Fonseca, Inserm U1016 Plateforme Génomique, Paris, France.
Aim to study	To validate a common protocol and to evaluate the efficiency and reliability of gNIPT of the most common chromosomal aneuploidies using a semiconductor sequencing platform.
Funding source or sponsor of the study	Not reported.
Information about the authors contacted	Author was contacted on: 19 January 2016 and 23 March 2016. Reply received on: 16 February 2016.
Notes	Conference Abstract of the 10 th European Cytogenetics Conference of the European Cytogenetics Association, ECA 2015. Prospective study of 500 pregnant women at high risk of fetal aneuploidy who undergo fetal karyotyping. The NIPT results matched the fetal karyotyping results in all of the cases: all trisomies were detected.

ISRCTN11174071

Trial name or title	Comparison of false positive rates in prenatal combined screening and cell free DNA screening for trisomy 21 (ReFaPo study).
Target condition and refer- ence standard(s)	Target condition: T21. Reference standard: prenatal or postnatal karyotype.
Index and comparator tests	gNIPT.
Starting date	July 2016.
Contact information	Karl Oliver Kagan University of Tuebingen Department of Obstetrics and Gynaecology Calwerstrasse 7 Tuebingen 72076

ISRCTN11174071 (Continued)

	Germany
Aim to study	To compare the false positive rate of cell-free DNA and traditional screening methods in a ran- domised controlled trial in a cohort without prior risk of fetal aneuploidy.
Funding source or sponsor of the study	Study funded by CENATA GmbH who does the analysis.
Information about the authors contacted	No need for further contact.
Notes	Target number of participants: 1400.
	Recruitment end date: March 2017.
	Intention to publish date: October 2018.
	DOI 10.1186/ISRCTN11174071

Lin 2014

Trial name or title	Clinical implementation of noninvasive prenatal testing in twin pregnancies with assisted repro- ductive technique treatment.
Target condition and reference standard(s)	Target conditions: T21, T18 and T13.
	Reference standards: fetal karyotype or clinical outcomes.
Index and comparator tests	gNIPT by MPSS.
	Commercial test: BGI Shenzhen's prenatal test.
Starting date	Not reported.
Contact information	BGI-Shenzhen
	Shenzhen, China
Aim to study	To assess the clinical implementation of MPS-based NIPT in twin pregnancies with assisted repro- ductive technique treatment.
Funding source or sponsor of the study	Not reported but BGI-Shenzhen made sequencing and analyses.
Information about the authors contacted	No need for further contact.
Notes	Some women were still pregnant at the time of writing this poster abstract.

Mu 2014

 Trial name or title
 Maternal non-invasive fetal DNA test used in prenatal diagnosis.

 Target condition and reference standard(s)
 Target conditions: T21, T18, T13 and 45,X.



Mu 2014 (Continued)	
	Reference standard: fetal karyotype of amniotic fluid.
Index and comparator tests	gNIPT by MPSS.
	NIFTY™ prenatal test by BGI-Shenzhen.
Starting date	In 2012.
Contact information	Mu Y.
	Beijing United Family Hospital
	Beijing, China.
Aim to study	Not reported.
Funding source or sponsor of the study	Not reported.
Information about the authors con- tacted	Author was contacted on: 19 April and 19 May 2016.
	No reply received from the author.
Notes	Poster abstract. Some women were still pregnant at the time of writing this poster ab- stract.

NCT01429389

Trial name or title Target condition and reference standard(s)	Specimen collection from pregnant women at increased risk for fetal aneuploidy. Target condition: T21.
Target condition and reference standard(s)	Target condition: T21.
0	с С
	Reference standard: fetal karyotype.
Index and comparator tests	gNIPT.
Starting date	May 2011.
Contact information	Sequenom, Inc.
Aim to study	To develop a prenatal aneuploidy test using ccfDNA from blood samples from preg- nant women who have an increased risk indicator/s for fetal chromosomal aneu- ploidy detection (T21).
Funding source or sponsor of the study	Study funded by Sequenom, Inc.
Information about the authors contacted	No need for further contact.
Notes	

NCT01472523

Trial name or title	A safer pre-natal diagnosis using free DNA in maternal blood (IONA®).
Target condition and reference standard(s)	Target conditions: T21, T18, T13 and other chromosomal abnormalities yet to be determined.

NCT01472523 (Continued)

Reference standards: prenatal karyotype and follow-up for 1 year.

Index and comparator tests	gNIPT by TMPS (selective amplification of fetal DNA) by Premaitha Health.
Starting date	April 2007.
Contact information	Brenda Kelly
	National Health Service, United Kingdom
Aim to study	To validate a novel gNIPT method that could increase the titre of fetal DNA within a given sample.
Funding source or sponsor of the study	Study funded by Premaitha Health.
Information about the authors con- tacted	No need for further contact.
Notes	

NCT01545674

Trial name or title	Prenatal Non-invasive Aneuploidy Test Utilizing SNPs trial (PreNATUS).
Target condition and reference standard(s)	Target conditions: aneuploidy in a fetus at chromosomes 13, 18, 21, X and Y.
	Reference standard: fetal karyotype.
Index and comparator tests	gNIPT by TMPS (SNP based technology by Natera, Inc.).
Starting date	January 2012.
Contact information	Ronald Wapner, MD, Columbia University
Aim to study	To assess the diagnostic capability of an informatics enhanced SNP based technology (Parental Support) to identify pregnant women who are carrying a fetus with an aneuploidy from free floating DNA in the maternal blood.
Funding source or sponsor of the study	Study funded by Natera, Inc.
Information about the authors contacted	No need for further contact.
Notes	

NCT01925742

Trial name or title	Study of the efficacy of new non-invasive prenatal tests for screening for fetal trisomies using ma- ternal blood (PEGASUS).
Target condition and refer- ence standard(s)	Target conditions: T21, T18 and T13.

NCT01925742 (Continued)	Reference standards: prenatal or neonatal karyotype or medical record from birth.
Index and comparator tests	gNIPT by Semiconductor MPSS (Ion Torrent Proton™) or optical-based MPSS (Illumina) or by TMPS with Harmony™ prenatal test by Ariosa Diagnostics, Inc.
Starting date	August 2013.
Contact information	François Rousseau CHU de Québec Québec, Canada
Aim to study	To perform a pan-Canadian large-scale validation study comparing the relative effectiveness and clinical performances of 2 index gNIPT methods using fetal ccfDNA in maternal blood in Canadian clinical laboratories between themselves and with that of fetal karyotype for detecting fetal aneuploidy of chromosomes 13, 18 and 21 and to compare the accuracy of this new gNIPT method with traditional prenatal screening methods.
Funding source or sponsor of the study	Study funded by Centre Hospitalier Universitaire de Québec, Laval University, Genome Canada, Genome Quebec, Genome British Columbia and Canadian Institutes of Health Research (CIHR).
Information about the authors contacted	No need for further contact.
Notes	Recruitment of patients completed (near 5000 pregnant women enrolled). at the time of writ- ing, they are sequencing 3600 pregnant women with the 2 gNIPT MPSS platforms. A subsample of about 2300 blood samples was analysed by Ariosa Diagnostics, Inc (TMPS).
	Estimated study completion date: June 2017.

NCT02201862	
Trial name or title	Non-invasive Chromosomal Evaluation of Trisomy study (NICHE).
Target condition and reference	Target conditions: T21, T18 and T13.
standard(s)	Reference standard: fetal karyotype.
Index and comparator tests	gNIPT by TMPS by Ariosa Diagnostics, Inc.
Starting date	April 2014.
Contact information	Romielle Aquino
	408-209-9098
	raquino@ariosadx.com
	Or
	Thomas Musci
	408-229-7500
	tmusci@ariosadx.com



NCT02201862 (Continued)

Aim to study	To provide clinically annotated samples to support continued improvements in the Ariosa Di- agnostics, Inc Test content, methodology, specimen processing and quality control.
Funding source or sponsor of the study	Study funded by Ariosa Diagnostics, Inc.
Information about the authors con- tacted	No need for further contact.
Notes	

NCT02278536

Trial name or title	Multiple gestation study.
	Multiple gestation study.
Target condition and reference standard(s)	Target conditions: T21, T18, T13 and SCA.
	Reference standards: fetal karyotype (amniocentesis or CVS) or genetic testing from cheek swab or saliva from live-born children.
Index and comparator tests	gNIPT by TMPS by Natera, Inc.
Starting date	March 2013.
Contact information	Brian Kirshon
	Houston Perinatal Associates
	Or
	Zach Demko
	Natera, Inc.
Aim to study	To demonstrate the accuracy of our new NATUS diagnostic method to determine the genetic health of the developing fetuses in a multiple gestation pregnancy from a maternal blood sam ple.
Funding source or sponsor of the study	Study funded by Natera, Inc.
Information about the authors contacted	No need for further contact.
Notes	

NCT02278874

Trial name or title	High risk multiple gestation study.
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13 and SCA.
	Reference standards: fetal karyotype (amniocentesis or CVS) or genetic testing from cheek swab or saliva from live-born children.



NCT02278874 (Continued)

Index and comparator tests	gNIPT by TMPS by Natera, Inc.
Starting date	August 2014.
Contact information	Joanne Stone
	Mt. Sinai Hospital, New York
Aim to study	To demonstrate the accuracy of our proprietary algorithm method to determine the genetic health of the developing fetuses in a multiple gestation pregnancy from a maternal blood sample.
Funding source or sponsor of the study	Study funded by Natera, Inc., Mount Sinai Hospital New York, Montefiore Medical Center, Long Is- land Jewish Medical Center and Tufts Medical Center.
Information about the authors contacted	No need for further contact.
Notes	

NCT02317965	
Trial name or title	Non-invasive screening for fetal aneuploidy.
Target condition and reference stan-	Target conditions: T21 and T18.
dard(s)	Reference standard: fetal karyotype.
Index and comparator tests	gNIPT by MPSS by Progenity, Inc.
Starting date	March 2015.
Contact information	Richard Porreco
	Obstetrix Medical Group of Colorado
Aim to study	To detect whole chromosome abnormalities on all chromosomes 13, 16, 18, 21, X and Y, in the fetus through analysis of ccfDNA and compound sample DNA in maternal blood.
Funding source or sponsor of the study	Study funded by Progenity, Inc.
Information about the authors contacted	No need for further contact.
Notes	

NCT02424474

Trial name or title	T21,18 and 13 screening by cell free fetal DNA in low risk patients (DEPOSA).
Target condition and reference standard(s)	Target conditions: T21, T18 and T13.
	Reference standard: fetal karyotype.
Index and comparator tests	gNIPT by MPSS.



NCT02424474 (Continued)

Starting date	June 2015.
Contact information	Alexandra Benachi
	Antoine Béclère Hospital
Aim to study	To evaluate the performance of gNIPT in a population of pregnant women with and without in vit- ro fertilisation (IVF) concomitantly to regular first-trimester trisomy 21 (T21) screening using ma- ternal age, nuchal fold measurement and serum screening.
Funding source or sponsor of the study	Study funded by Assistance Publique - Hôpitaux de Paris.
Information about the authors contacted	No need for further contact.
Notes	Recruitment of patients completed (933 pregnant women enrolled).

NCT02787486

Trial name or title	Expanded Noninvasive Genomic Medical Assessment: the Enigma study.
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13, microdeletion syndromes, sex chromosome abnormalities, infec- tious and other diseases, and blood group typing.
	Reference standard: fetal karyotype or medical records.
Index and comparator tests	gNIPT by MPSS provided by Progenity, Inc.
Starting date	October 2015.
Contact information	Paul Bien
	760-494-1743
	paul.bien@progenity.com
Aim to study	To evaluate the relative clinical sensitivity, specificity, and performance of the laboratory-devel- oped test as a screening test for fetal chromosomal aneuploidy, infectious and other diseases, and RhD genotyping in the general population of pregnant women.
Funding source or sponsor of the study	Study funded by Progenity, Inc.
Information about the authors contacted	No need for further contact.
Notes	

Sago 2015

Trial name or title

Nationwide demonstration project of next-generation sequencing of cell-free DNA in maternal plasma in Japan: 1-year experience



Sago 2015 (Continued)	
Target condition and refer- ence standard(s)	Target conditions: T21, T18 and T13. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or tissues of the miscar- riage or medical record from birth.
Index and comparator tests	gNIPT by MPSS. Blood samples were collected before invasive procedure. Commercial test: MaterniT21™ prenatal test from Sequenom, Inc.
Starting date	15 November 2012. Recruitment period between April 2013 to March 2014.
Contact information	Haruhiko Sago
	National center for Child-health and development
	Perinatal Center
	2-10-1Ookura, Setagaya-ku, Tokyo
	03-3416-0181
	sagou-h@ncchd.go.jp
Aim to study	To evaluate the quality of the genetic counselling in Japan. Sago 2015 reported the 1-year experi- ence of a nationwide demonstration project to introduce gNIPT of fetal aneuploidy from maternal plasma and discuss how to implement this program in Japan.
Funding source or sponsor of the study	Study supported by the Grant of the National Center for Child Health and Development 24-3, Japan. Sequenom, Inc made gNIPT.
Information about the authors contacted	Author was contacted on: 6 April and 14 June 2016. No reply received from the author.
Notes	Authors continue collecting follow-up data in the study population.

Trial name or title	Clinical implementation of non-invasive prenatal study for detecting aneuploidies by fetal DNA
	based on single nucleotide polymorphisms: 2 years in Mexico.
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13, 45,X, 47,XXY, 47,XYY and 47,XXX.
	Reference standards: fetal karyotype of chorionic villi or amniotic fluid or medical record from birth.
Index and comparator tests	gNIPT by TPMS.
	Commercial test: Natera's prenatal test.
Starting date	Recruitment period: March 2013 to February 2015.
Contact information	Dr. Rafael Sánchez Usabiaga
	rsanchez@medicafertil.com.mx
Aim to study	To describe our experience of 2 years integrating gNIPT by ccfDNA in its variant of single nucleotide polymorphism (SNPs) as a screening method for the detection of common aneuploidies, since 9 weeks of gestation.



Sanchez-Usabiaga 2015 (Continued)

Funding source or sponsor of the study	Not reported but Natera, Inc. made gNIPT sequencing and analyses.
Information about the authors contacted	No need for further contact.
Notes	There are 270 pregnant women included in this study.

Sistermans 2015	
Trial name or title	TRIDENT: or monitored NIPT implementation in the Netherlands.
Target condition and refer- ence standard(s)	Target conditions: T21, T18 and T13. Reference standard: fetal karyotype of chorionic villi or amniotic fluid is recommended in case of abnormal gNIPT test results. Neonatal clinical examination not mentioned.
Index and comparator tests	gNIPT by MPSS.
Starting date	01 April 2014.
Contact information	Dr. Erik Sistermans.
	VU University Medical Center Dept. of Clinical and Human Genetics Van der Boechorststraat 7 1081 BT Amsterdam NETHERLANDS
	+31-20-020-4448346 Email: e.sistermans@vumc.nl
Aim to study	To investigate and evaluate all relevant aspects of the introduction of NIPT in the Dutch prenatal screening program.
Funding source or sponsor of the study	The TRIDENT study was designed and proposed by the national multidisciplinary NIPT consortium.
Information about the authors contacted	Author have been contacted on: 9 December 2015 and 15 March 2016. Reply received on: 16 March 2016.
Notes	Conference abstract presented at the Annual conference of the European Society of Hu- man Genetics at Glasgow, Scotland, UK. http://www.emgo.nl/research/quality-of-care/re- search-projects/1451/trident-study-trial-by-dutch-laboratories-for-evaluation-of-non-invasive-pre- natal-testing-nipt/background/
	The authors plan to publish a full publication soon.

Torres 2015

Trial name or title	Genetic non invasive prenatal testing: A clinical and technical experience of 3000 cases with fol- low-up.
Target condition and refer-	Target conditions: T21, T18, T13, 45,X, 47,XXY, 47,XYY and 47,XXX.
ence standard(s)	Reference standard: fetal karyotype of amniotic fluid.

Torres 2015 (Continued)

Index and comparator tests

gNIPT by MPSS.

Commercial test: TrisoNIM® prenatal test by NIMGenetics Genomics.

Starting date	Not reported.
Contact information	Juan C Cigudosa
	NIMGenetics Genomics
	Madrid, Spain.
Aim to study	To show a NIPT protocol, called TrisoNIM [®] , which has been partially performed in our laboratory, based in massive parallel sequencing.
Funding source or sponsor of the study	Not reported.
Information about the authors contacted	Author were contacted on: 29 February, 22 March, 15 and 27 June 2016. Reply received on: 20 June 2016.
Notes	Full poster received from the authors.

Van Wymersch 2015

Trial name or title	Introduction of noninvasive prenatal testing for fetal trisomies: preliminary results and conse- quences on invasive samplings.
Target condition and refer- ence standard(s)	Target conditions: T21, T18, T13, 45,X, 47,XXY, 47,XYY and 47,XXX. Microdeletion syndromes can also be detected. Reference standards: fetal karyotype of chorionic villi or amniotic fluid or medical record from birth.
Index and comparator tests	gNIPT by MPSS. Blood samples were collected before invasive procedure. Cutpoint: not reported. Commercial test: NIFTY™ test (Bejing Genomics Institute, Hong-Kong, China).
Starting date	December 2013.
Contact information	Dr Van Wymersch Didier, Service de Gynécologie Obstétrique, Centre Hospitalier de Luxembourg, 4 Rue Barblé, L-1210 Luxembourg E-mail: vanwymersch.didier@chl.lu
Aim to study	To analyse a year of gNIPT implantation in our institute and to analyse gNIPT implication in chro- mosomal abnormalities screening politic.
Funding source or sponsor of the study	No reported. Samples analysed at BGI.
Information about the authors contacted	Author was contacted on: 12 September 2016. Reply received on: 29 September 2016.



Van Wymersch 2015 (Continued)

Notes

This publication showed the first 683 samples. At the time of writing, authors have a much larger population of 2132 pregnant women. No false negative results have been observed to date for all the pregnancies that have already come to term.

Willems 2014	
Trial name or title	The first 3000 Non-Invasive Prenatal Tests (NIPT) with the Harmony test in Belgium and the Netherlands.
Target condition and reference standard(s)	Target conditions: T21, T18 and T13. Reference standard: fetal karyotype of chorionic villi or amniotic fluid.
Index and comparator tests	gNIPT by TMPS.
	Commercial test: Harmony™ prenatal test by Ariosa Diagnostics, Inc.
Starting date	Recruitment period: March 2013 to December 2013.
Contact information	Patrick Willems
	patrick.willems@genetic-diagnostic.net
Aim to study	To report the results of the first 3000 consecutive gNIPT tests performed in pregnant women from Belgium and the Netherlands.
Funding source or sponsor of the study	Not reported. Ariosa Diagnostics, Inc made sequencing and analysis.
Information about the authors contacted	No need for further contact.
Notes	

Yu 2014a

Trial name or title	Maternal non-invasive fetal DNA test used in prenatal diagnosis.
Target condition and reference standard(s)	Target conditions: T21, T18, T13 and 45,X. Reference standards: cytogenetic tests and postnatal follow-up.
Index and comparator tests	gNIPT by MPSS by BGI-Shenzhen.
Starting date	Patients recruited in 2012.
Contact information	Yu M or Fei S.
	Beijing United Family Hospital.
Aim to study	To determine gNIPT accuracy in Chinese population.
Funding source or sponsor of the study	Not reported.

Yu 2014a (Continued)	
Information about the authors	Author was contacted on: 15 May 2016.
contacted	BGI was contacted on: 19 May 2016.
	No reply received from the author or BGI.
Notes	Conference abstract. Some women were still pregnant at the time of writing their conference abstract.

Zwiefelhofer 2014

Prenatal detection of fetal aneuploidy on the Ion Torrent Proton™ platform.
Target condition: T21. Reference standard: fetal karyotype.
gNIPT by MPSS on the Proton™ platform.
Not reported.
Sequenom, Inc.
To examine the performance of a gNIPT for fetal aneuploidy on the Ion Torrent Proton™ platform.
Study funded by Sequenom, Inc.
Author was contacted on: 19 April and 15 June 2016. Reply received on: 22 June 2016.
Full poster received from authors. This study includes 156 samples including 16 women carrying a T21 fetus. All patient samples were correctly identified according to their karyotype results.

CVS: chorionic villi sampling

gNIPT: genomics-based non-invasive prenatal testing

MPSS: massively parallel shotgun sequencing

TMPS: targeted massively parallel sequencing

DATA

Presented below are all the data for all of the tests entered into the review.

Table Tests. Data tables by test

Test	No. of studies	No. of participants
1 MPSS T21	41	50133
2 MPSS T18	38	49003
3 MPSS T13	29	46090



SMPSS 47, XXX 5 5449 SMPSS 47, XXY 8 6588 SMPSS 47, XXY 8 6629 SMPSS 47, XYY 8 6629 SMPSS 41, Zaneuploidies 44 50864 SMPSS, autosomes 43 50453 SMPSS, SCA 14 7911 SIT MPSS, SCA 14 7911 SIT MPSS, T13 10 22868 SIT MPS 47, XXX 6 2214 SIT MPS 47, XXX 2 586 SIT MPS 47, XXY 4 1021 SIT MPS 47, XXY 2 358 SIT MPS 41, Zaneuploidies 13 34473 SIT MPS, SCA 6 2214 SIT MPS, SCA 6 2214 SIT MPS, SCA 6 2214 SIT Traditional screening tests T21 2 1	Test	No. of studies	No. of participants
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rMPSS 47,XYY 8 6629 SMPSS all 7 aneuploidies 44 50864 DMPSS, autosomes 43 50453 L0 MPSS, SCA 14 7911 L1 TMPS T21 16 32487 L2 TMPS T18 12 30319 L3 TMPS T13 10 22868 L4 TMPS 45,X 6 2214 L5 TMPS 47,XXY 4 1021 L6 TMPS 47,XXY 2 358 L8 TMPS all 7 aneuploidies 21 35275 L9 TMPS, autosomes 18 34473 L9 TMPS, SCA 6 2214 L7 TMPS 47,XXY 2 358 L9 TMPS, autosomes 18 34473 L9 TMPS, autosomes 18 34473 L9 TMPS, SCA 6 2214 L1 Traditional screening tests, autosomes 5 24279 L2 Traditional screening tests T18 2 17753	5 MPSS 47, XXX	5	5449
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MPSS, autosomes 43 50453 10 MPSS, SCA 14 7911 11 TMPS T21 16 32487 12 TMPS T18 12 30319 13 TMPS T13 10 22868 14 TMPS 45,X 6 2214 15 TMPS 47,XXX 2 586 16 TMPS 47,XYY 4 1021 17 TMPS 47,XYY 2 358 18 TMPS all 7 aneuploidies 21 35275 19 TMPS, autosomes 18 34473 20 TMPS, SCA 6 2214 21 Traditional screening tests, autosomes 5 24279 21 Traditional screening tests T18 2 17753	7 MPSS 47,XYY	8	6629
14 7911 11 TMPS T21 16 32487 12 TMPS T18 12 30319 13 TMPS T13 10 22868 14 TMPS 45,X 6 2214 15 TMPS 47,XXX 2 586 16 TMPS 47,XXY 4 1021 17 TMPS 47,XYY 2 358 18 TMPS all 7 aneuploidies 21 35275 19 TMPS, autosomes 18 34473 10 TMPS, SCA 6 2214 17 Taditional screening tests, autosomes 5 24279 21 Traditional screening tests T18 2 17753	8 MPSS all 7 aneuploidies	44	50864
11 TMPS T21 16 32487 12 TMPS T18 12 30319 13 TMPS T13 10 22868 14 TMPS 45,X 6 2214 15 TMPS 47,XXX 2 586 16 TMPS 47,XXY 4 1021 17 TMPS 47,XYY 2 358 18 TMPS all 7 aneuploidies 21 35275 19 TMPS, succomes 18 34473 10 TMPS, SCA 6 2214 11 Traditional screening tests, autosomes 5 24279 12 Traditional screening tests T18 2 17753	9 MPSS, autosomes	43	50453
12 30319 13 TMPS T13 10 22868 14 TMPS 45,X 6 2214 15 TMPS 47,XXX 2 586 16 TMPS 47,XXY 4 1021 17 TMPS 47,XYY 2 358 18 TMPS all 7 aneuploidies 21 35275 19 TMPS, sutosomes 18 34473 20 TMPS, SCA 6 2214 21 Traditional screening tests, autosomes 5 24279 22 Traditional screening tests T18 2 17753	10 MPSS, SCA	14	7911
10 22868 14 TMPS 45,X 6 2214 15 TMPS 47,XXX 2 586 16 TMPS 47,XXY 4 1021 17 TMPS 47,XYY 2 358 18 TMPS all 7 aneuploidies 21 35275 19 TMPS, autosomes 18 34473 20 TMPS, SCA 6 2214 21 Traditional screening tests, autosomes 5 24279 22 Traditional screening tests T18 2 17753	11 TMPS T21	16	32487
14 TMPS 45,X 6 2214 15 TMPS 47,XXX 2 586 16 TMPS 47,XXY 4 1021 17 TMPS 47,XYY 2 358 18 TMPS all 7 aneuploidies 21 35275 19 TMPS, sutosomes 18 34473 20 TMPS, SCA 6 2214 21 Traditional screening tests, autosomes 5 24279 22 Traditional screening tests T18 2 17753	12 TMPS T18	12	30319
15 TMPS 47,XXX 2 586 16 TMPS 47,XXY 4 1021 17 TMPS 47,XYY 2 358 18 TMPS all 7 aneuploidies 21 35275 19 TMPS, autosomes 18 34473 20 TMPS, SCA 6 2214 21 Traditional screening tests, autosomes 5 24279 22 Traditional screening tests T21 2 17753 23 Traditional screening tests T18 2 17747	13 TMPS T13	10	22868
16 TMPS 47,XXY4102117 TMPS 47,XYY235818 TMPS all 7 aneuploidies213527519 TMPS, autosomes183447320 TMPS, SCA6221421 Traditional screening tests, autosomes52427922 Traditional screening tests T2121775323 Traditional screening tests T18217747	14 TMPS 45,X	6	2214
LT TMPS 47,XYY2358L8 TMPS all 7 aneuploidies2135275L9 TMPS, autosomes1834473L0 TMPS, SCA62214L1 Traditional screening tests, autosomes524279L2 Traditional screening tests T21217753L3 Traditional screening tests T18217747	15 TMPS 47,XXX	2	586
L8 TMPS all 7 aneuploidies213527519 TMPS, autosomes183447320 TMPS, SCA6221421 Traditional screening tests, autosomes52427922 Traditional screening tests T2121775323 Traditional screening tests T18217747	16 TMPS 47,XXY	4	1021
19 TMPS, autosomes183447320 TMPS, SCA6221421 Traditional screening tests, autosomes52427922 Traditional screening tests T2121775323 Traditional screening tests T18217747	17 TMPS 47,XYY	2	358
20 TMPS, SCA6221421 Traditional screening tests, autosomes52427922 Traditional screening tests T2121775323 Traditional screening tests T18217747	18 TMPS all 7 aneuploidies	21	35275
21 Traditional screening tests, autosomes52427922 Traditional screening tests T2121775323 Traditional screening tests T18217747	19 TMPS, autosomes	18	34473
22 Traditional screening tests T2121775323 Traditional screening tests T18217747	20 TMPS, SCA	6	2214
23 Traditional screening tests T18 2 17747	21 Traditional screening tests, autosomes	5	24279
	22 Traditional screening tests T21	2	17753
24 Traditional screening tests T13 1 11185	23 Traditional screening tests T18	2	17747
	24 Traditional screening tests T13	1	11185

Test 1. MPSS T21.

Test 2. MPSS T18.

Test 3. MPSS T13.

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Test 4. MPSS 45,X.

Test 5. MPSS 47, XXX.

Test 6. MPSS 47,XXY.

Test 7. MPSS 47,XYY.

Test 8. MPSS all 7 aneuploidies.

Test 9. MPSS, autosomes.

Test 10. MPSS, SCA.

Test 11. TMPS T21.

Test 12. TMPS T18.

Test 13. TMPS T13.

Test 14. TMPS 45,X.

Test 15. TMPS 47,XXX.

Test 16. TMPS 47,XXY.



Test 17. TMPS 47,XYY.

Test 18. TMPS all 7 aneuploidies.

Test 19. TMPS, autosomes.

Test 20. TMPS, SCA.

Test 21. Traditional screening tests, autosomes.

Test 22. Traditional screening tests T21.

Test 23. Traditional screening tests T18.

Test 24. Traditional screening tests T13.

ADDITIONAL TABLES

Target	Affected births ^a	Clinical features	Prognosis
condition	/100,000		
T21	140 to 230 ^{b,c}	Intellectual disability (mild to moderate), neurodevel- opmental problems, characteristic dysmorphic fea- tures, congenital defects (cardiac (44% to 58%) and gastrointestinal system (4% to 10%)), vision or hear- ing impairment (38% to 80%) and obstructive sleep apnoea syndrome (57%) ^{d,e}	Mean and median life ex- pectancies are estimated to be 51 and 58 years old ^f
T18	59¢	Severe intellectual disability and a wide range of sig- nificant malformations (cardiac defects, gastrointesti- nal system defects, renal anomalies, central nervous system defects (apnoea and seizures)) ^{d,g}	Most affected fetuses die in utero. Median survival has been estimated at 14 days (95% confidence interval (CI) 10 to 20) and 8% (95% CI 4 to 14) reach 1 year of age ^h

Table 1. Characteristics of target conditions (Continued)

Т13	23 ^c	Severe intellectual disability, seizures and several dysmorphic features, malformations of the extremi- ties, cardiac defects, renal anomalies, and abdominal wall defects ^{d,i}	Most affected fetuses die in utero. Median survival time has been estimated at 10 days (95% CI 7 to 19) and 8% (95% CI 4 to 14) reach 1 year of age ^h
45,X	30 to 50 ^{c,j}	Learning disabilities (70%), short stature, congenital heart diseases (30%) and gonadal dysgenesis (90% with amenorrhoea and infertility due to early ovarian failure) ^{k,l}	Mortality in 45,X women is 3- fold higher than in the gen- eral population with an aver- age life span of 69 years ^m
47,XXY	12 ^c	Learning disabilities (> 75%), small testes (> 95%), azoospermia (> 95%), male infertility (91% to 99%), decreased testosterone level (63% to 85%) and gy- naecomastia (38% to 75%) ^{l,n}	Life expectancy is slightly shorter (approximately 2 years) than euploid men ⁿ
47,XXX	6 <i>c</i>	Developmental delays (motor and speech), learning or intellectual disability, attention deficits (25% to 35%), mood disorders (anxiety and depression), tall stature (80% to 89%), clinodactyly (42% to 65%), hy- potonia in infancy (55% to 71%), genitourinary mal- formations and congenital heart defects ^o	Mortality significantly in- creased with a median sur- vival age of 70.9 years com- pare to 81.7 years for euploid females ^p
47,XYY	3с	Developmental delays (speech, language and motor), attention deficit disorder (52%), tall stature (78%), central adiposity, macrocephaly (33%), hypotonia (63%), clinodactyly (52%), hypertelorism (59%) and testicular enlargement for age (50%) but no increase in genital anomalies ⁹	Mortality increased with a re- duction of life span of 10.3 years compared to euploid men ^r

45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, T21: trisomy 21, T18: trisomy 18, T13: trisomy 13. ^{*a*}Including live births, fetal deaths and terminations of pregnancy.

b(Christianson 2006; Parker 2010) c(Wellesley 2012) d(Driscoll 2009) e(Irving 2012; Weijerman 2010) f(Wu 2013b) g(Cereda 2012) *h*(Wu 2013a) ⁱ(Chen 2009) j(Stochholm 2006) k(Karnis 2012; Mazzanti 1998; Sybert 2004) ¹(Tyler 2004) *m*(Saenger 1996; Schoemaker 2008) n(Groth 2013) o(Tartaglia 2010) P(Stochholm 2010b) 9(Bardsley 2013; Leggett 2010) r(Stochholm 2010a).

Test name	Method	Aneuploidy	Reported	Reported	Reported
(Company,			sensitivity	specificity	false positive
Genomics-based nor	n-invasive prenatal te	sting for detection of	fetal chromosomal aneu	ploidy in pregnant women (Review)	234

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country)			% (95% CI)	% (95% CI)	rate %
Bambni™	MPSS	T21	100.0 (ND)	> 99.9 (ND)	< 0.1
Test		T18	100.0 (ND)	> 99.9 (ND)	< 0.1
(Berry Genomics		T13	100.0 (ND)	> 99.9 (ND)	< 0.1
Co. Ltd, China)		45,X	100.0 (ND)	99.8 (ND)	0.0
		47,XXX	100.0 (ND)	100.0 (ND)	0.1
		47,XXY	100.0 (ND)	100.0 (ND)	0.0
		47,XYY	100.0 (ND)	100.0 (ND)	0.0
GENOMOM	MPSS	T21, T18	99.0 (ND)	ND	ND
(Genome Care,		and T13			
Korea)		SCA	95.0 (ND)	ND	ND
Harmony™	Oligo TMPS	T21	> 99.0 (ND)	> 99.9 (ND)	< 0.1
prenatal test		T18	97.4 (ND)	> 99.9 (ND)	< 0.1
(Ariosa Diagnostics,		T13	93.8 (ND)	> 99.9 (ND)	< 0.1
Inc., USA)		45,X ^b	96.3 (81.7 to 99.8)	99.5 (98.1 to 99.9)	0.5
		47,XXX ^b	100.0 (ND)	99.5 (98.1 to 99.9)	0.5
		47,XXY ^b	100.0 (61.0 to 100.0)	100.0 (99.0 to 100.0)	0.0
IONA [®] test	MPSS	T21	> 99.0 (ND)	> 99.0 (ND)	< 1.0
(Premaitha Health		T18	> 99.0 (ND)	> 99.0 (ND)	< 1.0
plc, UK)		T13	> 99.0 (ND)	> 99.0 (ND)	< 1.0
(Laboratoire	MPSS	T21, T18	> 99.8 (ND)	> 99.8 (ND)	< 0.2
CERBA, France)		and T13			
MaterniT21™	MPSS	T21	99.1 (96.6 to 99.9)	99.9 (99.7 to 99.9)	0.1
Plus test		T18	> 99.9 (93.9 to 100.0)	99.6 (99.3 to 99.7)	0.4
(Sequenom Inc.,		T13	91.7 (61.0 to 99.0)	99.7 (98.5 to 99.5)	0.3
USA)		combined sex aneuploidies	96.2 (ND)	99.7 (ND)	0.3
MomGuard™	MPSS	T21, T18, T13,	> 99.0 (ND)	ND	ND
(LabGenomics,		45,X, 47,XXX,			



Table 2. Reported accuracy of commercially available genomics-based non-invasive prenatal tests^a (Continued)

Korea)		47,XXY, 47,XY	ſY		
NIFTY™ test	MPSS	T21	99.2 (ND)	100 (ND)	0
(Bejing Genomics		T18	98.2 (ND)	100 (ND)	0
Institute (BGI),		T13	100 (ND)	100 (ND)	0
China)		45,X	> 99.9 (ND)	> 99.9 (ND)	< 0.1
Panorama™	SNP TMPS	T21	> 99.9 (ND)	100 (ND)	0
prenatal test ^c		T18	> 96.4 (ND)	> 99.9 (ND)	< 0.1
(Natera, Inc., USA)		T13	> 99.9 (ND)	100 (ND)	0
		45,X	> 92.9 (ND)	> 99.9 (ND)	< 0.1
PrenaTest [®]	MPSS	T21	98.7 (ND)	99.9 (ND)	0.1
(LifeCodexx AG,		T18	100 (ND)		
Germany)		T13	100 (ND)		
		45,X	90.9 (ND)	98.8 (ND)	1.2
		47,XYY	100 (ND)		
Prendia	MPSS	T21	100.0 (88.8 to 100.0)	100.0 (98.0 to 100.0)	0.0
(Genesupport,		T18	95.8 (76.8 to 99.7)	100.0 (97.0 to 100.0)	0.0
Switzerland)		T13	100.0 (74.6 to 100.0)	100.0 (98.1 to 100.0)	0.0
		45,X	100.0 (74.6 to 100.0)	100.0 (98.1 to 100.0)	0.0
		47,XXX	100.0 (46.2 to 100.0)	100.0 (98.2 to 100.0)	0.0
Tranquility	MPSS	T21	99.9 (ND)	99.8 (ND)	0.2
(Genoma,		T18	99.9 (ND)	99.9 (ND)	0.1
Switzerland)		T13	99.9 (ND)	99.7 (ND)	0.3
verifi [®] prenatal	MPSS	T21	99.5 (98.7 to 99.5)	99.8 (98.9 to 99.9)	0.2
test		T18	97.3 (94.2 to 98.2)	99.7 (99.5 to 99.9)	0.3
(Illumina, Inc., USA)		T13	98.0 (95.6 to 98.9)	99.8 (99.8 to 99.9)	0.2
		45,X	95.0 (75.1 to 99.9)	99.0 (97.6 to 99.7)	1.0
VisibiliT™	MPSS	T21	> 99.0 (80.8 to 100)	> 99.9 (99.5 to 100)	< 0.1
(Sequenom Inc.,		T18	> 99.0 (65.5 to 100)	> 99.9 (99.5 to 100)	< 0.1
USA)					

45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, T21: trisomy 21, T18: trisomy 18, T13: trisomy 13 CI: confidence interval, MPSS: massively parallel shotgun sequencing, ND: no data available, TMPS: targeted massively parallel sequencing and SNP: single nucleotide polymorphism.

a(Ariosa Diagnostics 2016; BGI 2014; BGI 2016; Berry Genomics 2016; Genoma 2016; Genome Care 2016; Illumina 2014; Illumina 2016; LabGenomics 2016; LifeCodexx 2016; Natera 2016; Genesupport 2016; Premaitha Health plc 2016; Sequenom 2016).

^b(Hooks 2014).

^cDNA of maternal and paternal origin are needed.

Screening tests	First trimester	Second trimester				
	(before 14 weeks' gesta- tion)	(14 to 20 weeks' gestation)				
Ultrasonography	NT measurement	Various morphologic measurements that modify the prior risk established				
Combined test	 hCG (free β or total) PAPP-A NT measurement 	NA				
Triple test	NA	 hCG (free β or total) uE3 AFP 				
Quadruple test	NA	 hCG (free β or total) uE3 AFP inhibin A 				
Sequential test ^b	 free β hCG PAPP-A NT measurement 	 Invasive test is offered if 1st trimester result is positive Quadruple test is offered if 1st trimester result is negative 				
Contingent test ^b	 free β hCG PAPP-A NT measurement 	 Invasive test is offered if 1st trimester result is positive Quadruple test is offered after an intermediate 1st trimester result No test is offered after a low-risk result 				
Serum integrated test ^c	• PAPP-A	Triple or Quadruple test				
Integrated test ^c	PAPP-ANT measurement	Quadruple test				

Table 3. Traditional screening tests (mostly for T21)^a

Maternal age is often included in the algorithm for prenatal screening tests. AFP: alpha-fetoprotein, hCG: human chorionic gonadotropin, NA: not applicable, NT: nuchal translucency, PAPP-A: pregnancy associated plasma protein A and uE3: unconjugated estriol. *a*(Gekas 2009; Okun 2008; Wald 2005).

^bA test result was available after first-trimester screening test.

^cSingle test result available after second-trimester screening test.

Study ID	Target condi-	Study design and	Prior risk	Index test details	Cutpoint	Reference	Comparator
	tion(s)	participants				standard	
MPSS							
Alberti 2015	T21	 Case-control study (1:2) from a prospective cohort 976 singleton pregnancies enrolled, 183 were analysed 	High risk	 Illumina HiSeq 2000 se- quencer without multiplex- ing In-house test FF measured 	Z score of 3	Fetal karyoty- pe ^a	
Benachi 2015	T21, T18, T13	 Blinded retrospective study 900 singleton or twin pregnancies enrolled, 886 were analysed 	High risk	 Illumina v3 flow-cell on a HiSeq 1500 sequencer in 12-plex Commercial - Laboratoire CERBA FF measured 	Z score of 3 for T21; 3.95 for T18 and T13	Fetal kary- otype or neonatal clin- ical examina- tion	
Bianchi 2012	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Nested case-control study (1:4) from a prospective co- hort (MELISSA) 2882 singleton pregnancies enrolled, 503 for T21, 502 for T18, 501 for T13 and 489 for 45,X were analysed 	High risk	 Illumina HiSeq 2000 sequencer in 6-plex Commercial test - Verinata FF measured 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype	
Bianchi 2013	T21, T18, T13, 45,X	 Retrospective study from stored plasma 2882 singleton pregnancies enrolled, 113 were analysed 	High risk	 Illumina TrueSeq 3.0 se- quencing chemistry Commercial test - Verinata 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype	
Bianchi 2014a	T21, T18, T13	 Blinded prospective cohort study 2052 singleton pregnancies enrolled, 1952 for T21 and T18, and 1914 for T13 were analysed 	High, low and without prior risk	 Illumina HiSeq 2000 in 8-plex Commercial - verifi[®] prenatal test FF measured 	NCV of 4; rese- quenced if NCV is between 3 and 4	Fetal or post- natal kary- otype, neona- tal clinical ex- amination or medical record from birth	Standard screening (T21 only wit mixed cut- points) whicl include first- trimester combined test or a sec- ond-trimester result

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							(quadruple, serum inte- grated, fully integrated, or sequential).
Bijok 2014	T21, T18, T13	 Prospective cohort study 10 singleton pregnancies enrolled, 9 were analysed 	High risk	 Illumina Genome Analyzer Ilx or HiSeq 2000 sequencer in multiplex Commercial - NIFTY[™] test, BGI-Shenzhen FF measured 	NR	Fetal kary- otype	
Canick 2012	T21, T18, T13	 Case-control study 4664 pregnant women enrolled, 27 multifetal pregnancies were analysed 	High risk	 Illumina HiSeq 2000 sequencer in 4-plex Commercial test - Sequenom, Inc. FF measured 	Z score of 3	Fetal kary- otype	
Chen 2011	T18, T13	 Nested case-control study from prospective and retro- spective cohorts 392 singleton pregnancies enrolled, 289 were analysed 	High risk	 Illumina Genome Analyzer Ilx in 2-plex Commercial test - Se- quenom, Inc. 	Z score of 3	Fetal kary- otype	
Chiu 2011	T21	 Blinded case-control study (1:5) from prospective and retrospective cohorts 824 singleton pregnancies enrolled, 753 were analysed by 8-plex method and 314 by 2-plex method 	Mostly high (> 1/300) and some in- termediate risk (between 1/300 and 1/1000)	 Illumina Genome Analyzer II in 8-plex and 2-plex Commercial test - Se- quenom, Inc. FF measured 	Z score of 3	Fetal kary- otype	
Ehrich 2011	T21	 Blinded case-control study (1:11) from prospective co- hort 480 pregnant women en- rolled, 449 were analysed 	High risk	 Illumina Genome Analyzer Ilx sequencer in 4-plex Commercial test - Se- quenom, Inc. FF measured 	Z score of 2.5	Fetal kary- otype	

Table 4. Characteristics of included studies by type of gNIPT (Continued)

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Fiorentino 2016	T21, T18, T13	 Blinded prospective cohort study 7103 singleton pregnancies enrolled, 7082 were analysed 	Mostly high risk and without prior risk	 Illumina HiSeq 2500 sequencer in 15-plex, SAFeR™ algorithm. Commercial - Genoma's prenatal test FF measured 	NCV of 4; aneu- ploidy suspect- ed if NCV is be- tween 3 and 4	Fetal kary- otype or neonatal clin- ical examina- tion
Hou 2012	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Prospective cohort study 308 singleton pregnancies enrolled, 205 were analysed 	High risk	 Illumina HiSeq 2000 se- quencer Commercial test - BGI- Shenzhen 	NR	Fetal kary- otype
Huang 2014	T21, T18	 Blinded prospective cohort study 189 twin pregnancies en- rolled, 189 were analysed 	High risk	 Illumina Genome Analyzer Ilx or HiSeq 2000 sequencer Commercial test - BGI- Shenzhen 	L score of 1 and t score of 2.5 in- cluding warning zone	Fetal kary- otype
Jeon 2014	T21, T18	 Prospective cohort study 155 singleton pregnancies enrolled, 155 were analysed 	High risk	 Ion Torrent PGM or HiSeq 2000 sequencers, 10 sam- ples per Chip Commercial test - Genome Care 	Z score of 2.566 for T21; 2.459 for T18.	Fetal kary- otype
Jiang 2012	T21, T18, T13, 45,X, 47,XXY, 47, XYY	 Prospective cohort study 903 pregnant women enrolled, 903 were analysed 	High risk	 Illumina Genome Analyzer Ilx or HiSeq 2000 sequencer in multiplex Commercial - NIFTY™ test, BGI-Shenzhen FF measured 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype
Johansen 2016	T21, T18, T13	 Prospective cohort study 375 singleton pregnancies enrolled, 173 were analysed 	High risk	 Ion Proton[™] sequencer in 5-plex In-house test FF measured 	Z score of 4 (unclassified if Z score is be- tween 3 and 4) and WISECON- DOR of 1%	Fetal kary- otype
Ke 2015	T21, T18, T13	 Prospective cohort study 2340 singleton pregnancies enrolled, 2340 were analysed 	High risk	 High throughput sequenc- ing platform Commercial test - BGI- Shenzhen 	T score of 3	Fetal kary- otype or new- born outcome

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Kim 2016	T21	 Blinded prospective cohort study 101 pregnant women en- rolled, 101 were analysed 	High risk	 Ion Proton[™] sequencer in multiplex Commercial test - Genome Care 	Z score of 2.10 for Ion Proton™	Fetal kary- otype
Lau 2012	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Blinded prospective cohort study 108 singleton pregnancies enrolled, 108 were analysed 	Mostly high risk	 Illumina HiSeq 2000 sequencers in 12-plex Commercial - NIFTY™ test, BGI-Shenzhen 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype
Lee 2015	T21, T18, T13 and SCA (no case found)	 Blinded prospective cohort study 93 singleton and multife- tal pregnancies enrolled, 92 were analysed 	High risk	 Illumina MiSeq sequencer in 12-plex or NextSeq se- quencer in 96-plex Commercial test - Mom- Guard[™], LabGenomics FF measured 	Z score of 4 (in- termediate risk if Z score is be- tween 2.5 and 4) for T21 and T18; 2.8 for T13 (intermediate risk if Z score is between 1.9 and 2.8)	Fetal or neonatal karyotype
Lefkowitz 2016	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Retrospective cohort, blind- ed case-control study 5321 pregnant women en- rolled but 1222 were selected and 1166 were analysed 	High risk	 Illumina HiSeq 2000 sequencer in 6-plex or uniplex Commercial test - Sequenom, Inc. FF measured 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype
Liang 2013	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Blinded prospective cohort study 435 singleton and twin preg- nancies enrolled, 412 were analysed 	High risk	 Illumina HiSeq 2000 sequencer in 8-plex or 12-plex Commercial test - Berry Genomics Co. Ltd. FF measured 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype
Liu 2012	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Prospective cohort study 153 pregnant women enrolled, 153 were analysed 	High risk	Illumina HiSeq sequencer in multiplex.	Z score of 3	Fetal kary- otype
Ma 2016	T21, T18, T13	Blinded retrospective (archived samples) and prospective cohorts study	High and low risk	 Sequencing on BGISEQ-1000 in 16 or 24- plex Commercial test - BGI- Shenzhen 	Z score of 3	Fetal kary- otype or post- natal fol- low-up

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		 10,598 singleton pregnan- cies enrolled, 10,579 were analysed 				
Mazloom 2013	45,X, 47,XXX, 47,XXY, 47,XYY	 Blinded prospective cohort study 1975 singleton pregnancies enrolled, 411 samples from the validation set were analysed 	High risk	 Illumina v3 flow-cell on a HiSeq 2000 sequencer in 12-plex Laboratory test develop- ment by Sequenom, Inc. FF measured 	Different cut- points used for the four SCA ^b	Fetal kary- otype
Palomaki 2012	T21, T18, T13	 Nested case-control study (1:3) 4664 pregnant women en- rolled but 1988 singleton pregnancies were selected and 1971 were analysed 	High risk	 Illumina HiSeq 2000 sequencer in 4-plex Commercial test - Sequenom, Inc. FF measured 	Z score of 3 for T21; 3.88 for T18; 7.17 for T13	Fetal kary- otype
Papa- georghiou 2016a	T21, T18, T13	 Retrospective cohort, case- control study (1:9) 442 singleton and twin pregnancies enrolled, 426 singleton pregnancies were analysed 	High risk	 Ion Proton[™] sequencer in 8-plex Commercial - IONA[®] test, Premaitha Health (public limited company in UK) FF measured 	Likelihood ratio of 1 and mater- nal age-adjust- ed probability risk score	Fetal kary- otype or med- ical record from birth
Papa- georghiou 2016b	T21, T18, T13	 Retrospective cohort, case- control study (1:9) 442 singleton and twin preg- nancies enrolled, 11 twin pregnancies were analysed 	High risk	 Ion Proton[™] sequencer in 8-plex Commercial - IONA[®] test, Premaitha Health (public limited company in UK) FF measured 	Likelihood ratio of 1 and mater- nal age-adjust- ed probability risk score	Fetal kary- otype or med- ical record from birth
Poon 2016	T21, T18, T13	 Retrospective cohort, blind- ed nested case-control study 242 singleton pregnancies enrolled, 241 were analysed 	High risk	 Ion Proton[™] sequencer, IONA[®] software algorithm Commercial - IONA[®] test, Premaitha Health (public limited company in UK) FF measured 	NR (authors used the same gNIPT than Pa- pageorghiou 2016a)	Fetal kary- otype
Porreco 2014	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Blinded prospective cohort study 4170 singleton pregnancies enrolled, 3322 for auto- 	High risk	 Illumina HiSeq 2000 sequencer in 12-plex Commercial test - Sequenom, Inc. 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype or med- ical record from birth

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		Studed studies by type of gNIP somes, 3278 for 45,X and 47,XXX and 3201 for 47,XXY and 47,XYY were analysed		FF measured			
Sehnert 2011	T21, T18, T13, 45,X	 Retrospective (archived samples) cohort study 1014 singleton and multifetal pregnancies enrolled but only 47 singleton pregnancies in the test set were analysed in this review. 	High risk	 Illumina Genome Analyzer Ilx sequencer in uniplex Commercial test - Verinata 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype	
Shaw 2014	T21, T18, T13, 45,X, 47, XXX, 47,XXY, 47,XYY	 Prospective cohort study 201 singleton and multife- tal pregnancies enrolled, 200 were analysed 	High and low risk	 Illumina v2 HiSeq 2000 se- quencer in 12-plex Commercial test - Berry Ge- nomics Co. Ltd. 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype or med- ical record from birth	
Song 2013	T21, T18, T13, 45,X, 47,XXX, 47, XXY, 47,XYY (SCA data not shown in this review)	 Blinded prospective cohort study 1916 singleton pregnancies enrolled, 1741 were analysed 	Without prior risk	 Illumina v2 HiSeq2000 in 12-plex Commercial test- Berry Ge- nomics Co. Ltd. 	Z score of 3	Fetal or post- natal kary- otype or med- ical record from birth	Triple test fo T21 and T18 (cutpoint of in 270).
Song 2015	T21, T18, T13, 45,X, 47,XXX, 47,XYY	 Blinded prospective cohort study 213 singleton pregnancies enrolled, 204 were analysed 	High risk	 Illumina v2 HiSeq 2000 sequencer in 12-plex Commercial test - Berry Genomics Co. Ltd. FF measured 	Z score of 3	Fetal kary- otype or neonatal clin- ical examina- tion or both	
Stumm 2014	T21, T18, T13	 Prospective cohort, blinded study for T21 and unblinded for T18 and T13 522 singleton pregnancies enrolled, 472 were analysed 	High risk	 Illumina HiSeq 2000 sequencer in 12-plex (DAP.21 algorithm without CG correction) Commercial test - Life-Codexx AG FF measured 	MAD-based Z score of 3 for T21; 3.2 for T18; 3.9 for T13	Fetal kary- otype	
Sukhikh 2015	T21, T18, T13, 45,X	 Prospective cohort study 200 pregnant women enrolled, 200 were analysed 	High risk	 Ion Proton[™] sequencer In-house test 	T score of 5 for T21 and T18; 4 for T13; 0.04 Chrom. X and	Fetal kary- otype	

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		cluded studies by type of gNIP			0.04 Chrom. Y for 45,X	
Sung-Hee 2015	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Retrospective study 918 singleton pregnancies enrolled, 901 were analysed 	High risk	 Illumina Genome Analyzer Ilx or HiSeq 2000 sequencer in 12-plex Commercial - NIFTY[™] test, BGI-Shenzhen FF measured 	L score of 1 and t score of 2.5	Fetal kary- otype or med- ical record from birth
Tynan 2016	T21, T18, T13	 Blinded retrospective cohort study 1100 singleton pregnancies enrolled, 1048 were analysed 	High and without prior risk	 Illumina HiSeq 2000 or HiSeq 2500 sequencers in multiplex Commercial - VisibiliT[™] test, Sequenom, Inc. FF measured 	risk score of 1%	Fetal kary- otype or med- ical record from birth
Wang 2014	T21, T18, T13, 45,X	 Prospective cohort study 136 singleton pregnancies enrolled, 136 were analysed 	High risk	 Illumina HiSeq 2000 sequencer Commercial - NIFTY™ test, BGI-Shenzhen 	NR	Fetal or neonatal karyotype or clinical ex- amination at 42 days after birth or both
Wang 2015a	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Prospective cohort study 917 pregnant women enrolled, 917 were analysed 	High risk	 Illumina v2 HiSeq 2000 flow cell on a HiSeq sequencer Commercial test - Berry Ge- nomics Co. Ltd 	Z score of 3 for T21, T18 and T13; -3 for Chrom. X and 3 for Chrom. Y for sex Chrom. classification.	Fetal kary- otype or clini- cal follow-up to 6 months from birth
/ao 2014	T21, T18, T13 and SCA (SCA data not shown in this review)	 Retrospective study 5950 singleton pregnancies enrolled, 5530 were analysed 	High, low and without prior risk	 Illumina Genome Analyzer Ilx or HiSeq 2000 sequencer in 12-plex Commercial - NIFTY[™] test, BGI-Shenzhen FF measured 	Different cut- points used for autosomes and SCA ^b	Fetal kary- otype or clini- cal follow-up
Zhang 2016	T21, T18, 45,X, 47,XXX (SCA data not	 Blinded prospective cohort study 87 singleton pregnancies en- rolled, 87 were analysed 	High risk	 Illumina HiSeq 2000 sequencer in 12-plex Commercial test - Berry Genomics Co. Ltd. 	Z score of 3 for T21 (no other cutpoint report- ed)	Fetal or neonatal karyotype or neonatal clin-

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	shown in this review)					ical examina- tion
Zhou 2014a	T21, T18, T13	 Blinded prospective cohort study 306 singleton pregnancies enrolled, 301 were analysed 	High, low and without prior risk	 Illumina Genome Analyzer IIx or HiSeq 2000 sequencer in 12-plex Commercial - NIFTY[™] test, BGI-Shenzhen FF measured 	L score of 1 and t score of 2.5	Fetal or neonatal karyotype or birth outcome
Zhou 2014b	T21, T18, T13	 Blinded prospective cohort study 7705 singleton pregnancies enrolled, 3950 were analysed 	High, low and without prior risk	 Illumina Genome Analyzer Ilx or HiSeq 2000 sequencer in 12-plex Commercial - NIFTY[™] test, BGI-Shenzhen FF measured 	L score of 1 and t score of 2.5	Fetal or neonatal karyotype or birth outcome
TMPS						
Ashoor 2012	T21, T18	 Nested case-control study (1:3) from a prospective co- hort 400 singleton pregnancies enrolled, 397 were analysed 	High risk	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial - Harmony[™] prenatal test, Ariosa Diagnostics, Inc. 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal kary- otype
Ashoor 2013	T13	 Blinded prospective cohort study 2167 singleton pregnancies enrolled, 1949 were analysed 	High and low risk	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial - Harmony™ prenatal test, Ariosa Diagnostics, Inc. FF measured 	FORTE risk score of 1%	Fetal kary- otype or neonatal clin- ical examina- tion
Bevilacqua 2015	T21, T18, T13	 Prospective cohort study 515 multifetal pregnancies enrolled, 340 were analysed Women with singleton preg- nancies were excluded (in- complete 2 x 2 table). 	High and without prior risk	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial - Harmony[™] prenatal test, Ariosa Diagnostics, Inc. FF measured 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal or neonatal karyotype

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Comas 2015	T21, T18, T13, 45,X, 47,XXX, 47, XXY, 47,XYY (SCA data not shown in this review)	 Blinded prospective cohort study 333 singleton pregnancies enrolled, 312 were analysed 	High and without prior risk	 DANSR assay (FORTE algorithm) or SNP-based method Commercial - Panorama[™] test, Natera, Inc. or Harmony[™] prenatal test, Ariosa Diagnostics, Inc. FF measured 	Harmony™ pre- natal test: NR (usually Harmo- ny™ prenatal test uses FORTE risk score of 1%) Panorama™ test: NR	Fetal kary- otype or neonatal clin- ical examina- tion
del Mar Gil 2014	T21, T18, T13	 Retrospective cohort study 207 multifetal pregnancies enrolled, 192 twin pregnan- cies were analysed 	Without prior risk	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial - Harmony™ prenatal test, Ariosa Diagnostics, Inc. FF measured 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal kary- otype
Gil 2016	T21, T18, T13	 Prospective cohort study 11,692 singleton pregnancies enrolled, 3633 were analysed 	High and interme- diate risk ^c	 DANSR assay (usually with FORTE algorithm) Commercial - Harmony™ prenatal test, Ariosa Diag- nostics, Inc. 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal or post- natal kary- otype or neonatal clin- ical examina- tion
Hall 2014	T13	 Case-control study (1:3)/1000 singleton preg- nancies enrolled, 64 were analysed. 	High risk	 SNP-based method (NATUS algorithm), Illumina Genome Analyzer IIx or HiSeq sequencer, 11,000 or 19,488-plex targeted PCR Commercial - Natera's prenatal test FF measured 	NR	Fetal kary- otype or ge- netic testing of cord blood, buccal, saliva or products of conception
Hooks 2014	45,X, 47,XXX, 47, XXY, 47,XYY	 Case-control study from archived samples 432 singleton pregnancies enrolled, 414 were analysed 	High risk	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial - Harmony™ prenatal test, Ariosa Diagnostics, Inc. FF measured 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal kary- otype

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Jackson 2014	T21, T18, T13	 Prospective cohort study 1228 pregnant women enrolled, 1161 were analysed 	High and low risk	 DANSR assay (FORTE algorithm) Commercial - Harmony[™] prenatal test, Ariosa Diagnostics, Inc. 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal kary- otype or med- ical record from birth	
Korostelev 2014	T21, T18, T13, 45,X, 47,XXX, 47, XXY, 47,XYY	 Prospective cohort study 1968 singleton pregnancies enrolled, 685 were analysed 	High and without prior risk	 SNP-based method (NATUS algorithm), Illumina Genome Analyzer IIx or HiSeq sequencer, > 19,000-plex targeted PCR Commercial - Natera's prenatal test FF measured 	NR	Fetal kary- otype or med- ical record from birth	
Nicolaides 2012	T21, T18	 Retrospective study from archived plasma 2230 singleton pregnancies enrolled, 1949 were analysed 	Without prior risk	 DANSR assay (usually with FORTE algorithm) Commercial - Harmony™ prenatal test, Ariosa Diag- nostics, Inc. FF measured 	Risk score of 1%	Fetal kary- otype or neonatal clin- ical examina- tion	First-trimeste combined tes (cutpoint of 1 in 150).
Nicolaides 2013	T21, T18, T13, 45,X, 47,XXX, 47,XXY, 47,XYY	 Blinded prospective cohort study 242 singleton pregnancies enrolled, 229 were analysed 	High risk	 SNP-based method (NATUS algorithm), Illumina Genome Analyzer IIx or HiSeq sequencer, 19,488-plex targeted PCR Commercial - Natera's prenatal test FF measured 	NR	Fetal kary- otype	
Nicolaides 2014a	45,X, 47,XXX, 47,XXY, 47,XYY	 Case-control study (archived samples) 177 singleton pregnancies enrolled, 172 were analysed 	High risk	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial - Harmony™ prenatal test FF measured 	FORTE risk score of 1%	Fetal kary- otype	
Norton 2012	T21, T18	 Blinded prospective cohort study 4002 singleton pregnancies enrolled, 3080 were analysed 	High risk	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial test- Ariosa Diagnostics, Inc. 	FORTE risk score of 1%	Fetal kary- otype	

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able 4. Chara	acteristics of inc	cluded studies by type of gNIP1	(Continued)	• FF measured			
Norton 2015	T21, T18, T13	 Blinded prospective cohort study 18,955 singleton pregnan- cies enrolled, 15,841 were analysed 	Without prior risk	 DANSR assay (FORTE algorithm) Commercial - Harmony™ prenatal test, Ariosa Diagnostics, Inc. FF measured 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal or post- natal kary- otype, neona- tal clinical ex- amination or medical record from birth	First-trimeste combined tes (cutpoint of 1 in 270 for T21 and 1 in 150 for T18 and T13).
Pergament 2014	T21, T18, T13, 45,X	 Blinded prospective cohort study 1064 singleton pregnancies enrolled, 963 were analysed 	High and low risk	 SNP-based method (NATUS algorithm), Illumina Genome Analyzer IIx or HiSeq sequencer, 19,488-plex targeted PCR Commercial - Natera's prenatal test FF measured 	NR	Fetal kary- otype or ge- netic testing of cord blood, buccal, saliva or products of conception or birth outcome	
Persico 2016	T21, T18, 45,X, 47,XXX, 47,XXY, 47,XYY	 Blinded prospective cohort study 259 singleton pregnancies enrolled, 249 were analysed 	High risk	 SNP-based method (NATUS algorithm), Illumina Genome Analyzer IIx or HiSeq sequencer, 19,488-plex targeted PCR Commercial - Natera's prenatal test FF measured 	Risk score of 1%	Fetal kary- otype	
Quezada 2015	T21, T18, T13	 Prospective cohort study 2905 singleton pregnancies enrolled, 2785 were analysed 	Without prior risk	 DANSR assay (FORTE algorithm) Commercial - Harmony™ prenatal test FF measured 	NR (usually Har- mony™ pre- natal test us- es FORTE risk score of 1%)	Fetal or post- natal kary- otype, neona- tal clinical ex- amination or medical record from birth	First-trimeste combined tes (cutpoint of 1 in 100 for T21).
Saman- go-Sprouse 2013	45,X, 47,XXX, 47,XXY, 47,XYY	 Blinded prospective cohort study 201 singleton pregnancies (with known SCA and eu- ploid pregnancies) enrolled, 186 were analysed 	High and low risk	 SNP-based method (NATUS algorithm), Illumina HiSeq sequencer, 19,488-plex targeted PCR Commercial - Natera's prenatal test 	NR	Fetal kary- otype or ge- netic testing of cord blood, buccal, saliva	

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		• FF measured		or products of conception
 Sparks 2012a T21, T18 Case-control study from prospective cohort 338 singleton pregnancient enrolled, 167 were analysed 	25 	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial test- Ariosa Diagnostics, Inc. FF measured 	NR	Fetal kary- otype
 Verweij 2013 T21 Blinded prospective coho study 595 singleton pregnancie enrolled, 504 were analysed 	25	 DANSR assay (FORTE algorithm), Illumina HiSeq 2000 in 96-plex Commercial test- Ariosa Diagnostics, Inc. FF measured 	FORTE risk score of 1%	Fetal kary- otype
T13: trisomy 13. <i>a</i> Fetal karyotype include traditional banding techniques, spectra fluorescence polymerase chain reaction. <i>b</i> Different cutpoints used for autosomes or SCA as follows: Bianchi 2012: NCV of 4 (aneuploidy suspected if NCV is between 2.5 4 and NCV for Chrom. Y of 2.5 for 47,XXX; NCV for Chrom. X between with NCV for Chrom. Y is two times greater than expected NCV Chro Bianchi 2013: NCV of 4 (aneuploidy suspected if NCV is between 3 a Jiang 2012: t score of 3 and logarithmic LR of 1 for T21, T18 and T1 concentration by Chrom. X and Y independently for 47,XXY and 47,Y Lau 2012: Z score of 3 for T21, T18 and T13; if female fetus, Z score of Y of 3 for 47,XXY. Lefkowitz 2016: Z score of 3 for T21; S score of 3.95 for T18 and T13; Liang 2013: Z score of 3 for T21; S score of 3,95 for T18 and T13; Porreco 2014: Z score of 3 for T21; Z score of 3,95 for T18 and T13; regions between -2.5 and -3.5); Z score of -3.5 for 47,XYY with Chrom Sehnert 2011: NCV of 4 (unclassified if NCV is between 2.5 and 4) for	and 4) for T21, T18, ar -2.5 and 2.5 and NCV f m. X. nd 4) for T21, T18, and 3; if female fetus, t sco YY. or Chrom. X of -3 for 45 Z scores for SCA see M Chrom. X and ± 3 for Cl tween 2.5 and 3.5); Z s Y with Chrom. Y repres score of 3.5 for 47,XXX b. Y representation; Z s T21, T18, and T13; NC	nd T13; NCV for Chrom. X of -4 and or Chrom. Y > 33 for 47,XXY; NCV T13; NCV for Chrom. X of -3 and ore of -2.5 for 45,X and 47,XXX; t 5,X; if female fetus, Z score for Ch azloom 2013. hrom. Y for sex chromosome cla score of -3.5 for 45,X (non-repor sentation. ((non-reportable regions betwee core between -3.5 and 3.5 for 47	nd NCV for Chrom for Chrom. X of	. Y of 2.5 for 45,X; NCV for Chrom. X of 4 and NCV for Chrom. Y of 4 for 47,XYY of 3 for 45,X. bined with estimation of fetal ccfDNA 7,XXX; if male fetus, Z score for Chrom. ween -2.5 and -3.5); Z score of -3.5 for score of -3.5 for 45,X (non-reportable Y representation.
SDs from the mean of female samples for sex chromosome classific Shaw 2014: Z score of 3 for T21, T18, and T13; Z score of -3 for Chron Yao 2014: T score of 2.5 for T21, T18 and T13; if female fetus, T score estimation of fetal ccfDNA concentration by Chrom. X (expected values estimated by chromosome Y to that estimated by chromosome X) by	n. X and 3 for Chrom. Y pre for Chrom. X of -2. ue of zero) for 47,XXY;	5 for 45,X and 2.5 for 47,XXX; if if male fetus, T score for Chrom.	male fetus, T scor	

Table 4. Characteristics of included studies by type of gNIPT (Continued)

^cPregnant women with a first-trimester combined test selected for their risk of fetal aneuploidy (cutpoint of 1 in 100 for high risk and 1 in 101 to 1 in 2500 for intermediate risk).





Company	Number of studies	Number of affected/unaffected	Number of studies	Number of studies	Number of studies with
		pregnancies ^a	with pregnant	with high- risk	mixed risk ^b cohort
			women with- out	pregnant	
			prior risk of	women	
			fetal aneu- ploidy		
Ariosa	15	594/32,302	4	6	5
Diagnostics, Inc.					
Bejing Genomics	12	427/24,724	0	7	5
Institute (BGI)					
Sequenom, Inc.	9	904/8486	0	7	2
Berry Genomics	6	147/3414	1	4	1
Co. Ltd					
Natera, Inc.	6	276/2103	0	3	3
Illumina, Inc.	4	273/2342	0	3	1
In-house	3	114/442	0	3	0
Premaitha	3	99/579	0	3	0
Health plc					
Genome Care	2	21/235	0	2	0
CERBA	1	113/745	0	1	0
Genoma	1	105/6977	0	0	1
LabGenomics	1	8/84	0	1	0
LifeCodexx AG	1	55/417	0	1	0
Not reported	1	5/148	0	1	0
Total	65	3141/82,998	5	42	18

Table 5. Manufacturers of gNIPT used in the included studies by prior risk of fetal aneuploidy

^{*a*}We included pregnancies with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies. ^{*b*}Mixed-risk cohort included a mix of pregnant women without prior risk, low risk or high risk of fetal aneuploidy.

Table 6. Reasons for patient exclusion

Study ID	dy ID Number of preg- Reasons for exclusion nant women en- rolled		Number of women with results for 2 x 2 table analysis
Alberti 2015	976	 701 not selected for the case-control study 23 selected for reference set 8 selected for pretesting phase 47 low amount of DNA 11 low fetal fraction DNA or assay failure 3 haemolysed samples 	183
		Total: 793	
Ashoor 2012	400	• 3 samples failed amplification and sequencing	397
Ashoor 2013	2167	 165 selected for first phase (case-control study not included in this review) 53 failed amplification or sequencing Total: 218 	1949
Benachi 2015	900	 8 without reference standard result 6 low fetal fraction DNA or result appeared atypical 	886
		Total: 14	
Bevilacqua 2015	2362	 1847 not selected 159 without follow-up 11 failed samples 5 failed samples and were without follow-up 	340
		Total: 2022	
Bianchi 2012	2882	• 127 ineligible	503 (T21)
		 45 without karyotype 85 multifetal pregnancies	502 (T18)
		2091 not selected for this case-control study	501 (T13)
		 2 for tracking issue 16 without fetal DNA detected	489 (45,X)
		Total: 2366	
		In addition, other samples excluded from 2 x 2 tables for cen- sored complex karyotype:	
		 13 for T21 14 for T18 15 for T13 27 for 45,X 	
Bianchi 2013	2882	• 2769 not selected for the study	113
Bianchi 2014a	2052	10 failed blood quality control	1952 (T21 and T18)
		72 without clinical outcome17 without gNIPT result28 without standard screening result	1914 (T13)

Table 6. Reasons for patient exclusion (Continued) • 1 without gNIPT result and without standard screening result Total for T21 and T18: 100 Total for T13: 128 Bijok 2014 10 • 1 low fetal fraction DNA 9 • 4637 not selected for the case-control study Canick 2012 4664 27 • 103 selected for reference control Chen 2011 392 289 Chiu 2011 • 46 failed quality control for blood sampling 824 753 (8-plex) • 12 without karyotype • 2 twin pregnancies • 11 failed quality control for sequencing Total: 71 (8-plex) Comas 2015 333 • 17 without follow-up 312 • 3 unrepeated tests 1 failed test second time^a and without follow-up Total: 21 del Mar Gil 2014 207 • 11 low fetal fraction DNA 192 • 4 laboratory processing failures Total: 15 Ehrich 2011 • 13 preanalytic failure (including 9 for low plasma volume and 480 449 4 processing errors) • 18 failed quality control at second time (including 7 for low fetal fraction DNA) Total: 31 Fiorentino 2016 7103 21 failed quality control (unrepeated tests) 7082 Gil 2016 • 7994 patients did not undergo a gNIPT 3633 11,692 • 45 failed tests first time^b 20 failed tests second time Total: 8059 Hall 2014 > 1000 About 932 samples not selected for the case-control study 64 • 4 failed quality control Total: 936 Hooks 2014 432 • 18 low fetal fraction DNA, unusually high variation in ccfDNA 414 counts or failed QC Hou 2012 • 103 patients did not undergo a gNIPT 205 308 NR Huang 2014 189 189 Jackson 2014 1228 • 7 with other abnormal ultrasound 1161 • 14 opted for CVS without gNIPT

		 32 declined all testing 14 failed tests twice	
		Total: 67	
Jeon 2014	155	NR	155
Jiang 2012	903	NR	903
Johansen 2016	375	191 not selected for validation set11 low fetal fraction DNA	173
		Total: 202	
Ke 2015	2340	NR	2340
Kim 2016	101	NR	101
Korostelev 2014	1968	1043 without follow-up240 samples did not undergo a gNIPT	685
		Total: 1283	
Lau 2012	108	NR	108
Lee 2015	93	• 1 low fetal fraction DNA	92
Lefkowitz 2016	5321	 4099 not selected for the study 11 for incomplete follow-up 3 with confirmed mosaicism 11 low fetal fraction DNA 29 for technical reasons 2 for maternal event 	1166 (autosomes) 1144 (SCA)
		Total: 4155 (autosomes)	
		In addition:	
		22 sequencing failures for SCA	
		Total: 4177 (SCA)	
Liang 2013	435	11 without karyotype12 failed quality control	412
		Total: 23	
Liu 2012	153	NR	153
Ma 2016	10,598	 14 with incomplete follow-up 5 failed quality control	10,579
		Total: 19	
Mazloom 2013	1975	• 1564 selected for the training set	411
Nicolaides 2012	2230	 181 ineligible 46 low fetal fraction DNA 54 assay failures 	1949

Table 6. Reasons for patient exclusion (Continued) Total: 281

		Total: 281					
Nicolaides 2013	242	• 13 failed quality control	229				
Nicolaides 2014a	177	 1 failed quality control 4 low fetal fraction DNA	172				
		Total: 5					
Norton 2012	4002	774 ineligible57 low fetal fraction DNA91 assay failures	3080				
		Total: 922					
Norton 2015	18,955	 381 ineligible 64 withdrawn 384 handling errors 308 without standard screening test result 1489 without follow-up 192 low fetal fraction DNA 83 no fetal fraction DNA 213 high assay variance or assay failures 	15,841				
		Total: 3114					
Palomaki 2012	4876	 2888 not selected for this study 17 failed tests second time (mostly for low fetal fraction DNA) 	1971				
		Total: 2905					
Papageorghiou 2016a	442	 11 twin not selected 3 low fetal fraction DNA 2 failed quality control 	426				
		Total: 16					
Papageorghiou 2016b	442	 426 singleton not selected 3 low fetal fraction 2 failed quality control 	11				
		Total: 431					
Pergament 2014	1064	 13 not selected (other aneuploidies) 85 samples failed quality control for all five chromosomes (including 65 for low fetal fraction DNA) 	963 (T21) 964 (T18 and 45,X)				
		Total: 98	965 (T13)				
		In addition,					
		 3 samples failed only for T21 (total for T21: 101) 2 samples failed only for T18 and 45,X (total for T18 and 45,X: 100) 1 sample failed only for T13 (total for T13: 99) 					
D 1 0000		1 sample failed only for T13 (total for T13: 99)					
Persico 2016	259	 8 low fetal fraction DNA 2 failed internal quality control	249				

Table 6. Reasons for patient exclusion (Continued)

		Total: 10	
Poon 2016	242	1 low fetal fraction DNA	241
Porreco 2014	4170	 320 for insufficient sample volume 390 failed quality control 24 with incomplete follow-up 6 without invasive procedure 	3322 (T21, T18, T13 3278 (45,X, 47,XXX) 3201 (47,XXY, 47,XYY)
		In addition,	
		 54 failed quality control and 54 for complex autosome kary- otypes^c (total: 108 for autosomes) 102 failed quality control or other^d and 50 for complex SCA 	
		 102 Tailed quality control of other and 50 for complex SCA karyotype (total: 152 for 45,X and 47,XXX) 182 low fetal fraction DNA or other^d and 47 for complex SCA karyotype (total: 229 for 47,XXY and 47,XYY) 	
Quezada 2015	2905	 66 without follow-up 1 lost in mail 38 low fetal fraction DNA 15 assay failures 	2785
		Total: 120	
Samango-Sprouse 2013	201	 12 low fetal fraction DNA or poor DNA quality 2 without gNIPT result 1 with conflicting algorithm metrics 	186
		Total: 15	
Sehnert 2011	1014	 895 not selected for sequencing 71 selected for training set 1 twin pregnancy 	47
		Total: 967	
Shaw 2014	201	• 1 for early GA	200
Song 2013	1916	 102 without follow-up 64 failed quality control 9 failed quality control and without follow-up 	1741
		Total: 175	
Song 2015	213	 8 without follow-up 1 failed quality control	204
		Total: 9	
Sparks 2012a	338	• 171 selected for training set	167
Stumm 2014	522	 8 without reference standard 9 without consent 1 previously analysed 14 failed sequencing quality control 18 failed libraries 	472

Table 6. Reasons for patient exclusion (Continued)

Total: 50

200	NR	200
918	 8 ineligible 9 without follow-up	901
	Total: 17	
1100	 28 library preparation failures or failed quality control 24 for discretionary non reporting	1048
	Total: 52	
595	 75 ineligible 7 low fetal fraction DNA 9 laboratory processing failures or specimen issues 	504
	Total: 91	
136	NR	136
917	NR	917
5950	• 420 without follow-up	5530
87	NR	87
306	• 5 without follow-up	301
7705	 4 low fetal fraction DNA 3751 without follow-up	3950
	Total: 3755	
	918 1100 595 136 917 5950 87 306	918• 8 ineligible • 9 without follow-up Total: 171100• 28 library preparation failures or failed quality control • 24 for discretionary non reporting Total: 52595• 75 ineligible • 7 low fetal fraction DNA • 9 laboratory processing failures or specimen issues Total: 91136NR917NR5950• 420 without follow-up87NR306• 5 without follow-up7705• 4 low fetal fraction DNA • 3751 without follow-up

ccfDNA: circulating cell-free DNA, CVS: chorionic villi sampling, GA: gestational age, gNIPT: genomics-based non-invasive prenatal testing, NR: not reported by authors.

^{*a*}Second time: sample failed the second gNIPT assay.

^bFirst time: sample failed the initial gNIPT assay.

^cComplex autosome karyotypes are mosaic, triploidies, unbalanced rearrangements with missing or duplicated genetic material. ^dOther are copy number variation of the X chromosome is confounded by maternal component and cannot be determined.

Table 7. Proportion of pregnant women with a reference standard and assay failure during gNIPT process

	<u> </u>				00 1	
Study ID	Failure rate at		Failure rate of	Final failure rate	Aneuploid ^b	Euploid ^b
	first attempt	tests ^a	Tate of	total (%)	samples	samples
	(%)	(%)	repeated tests		(%)	(%)
			(%)			
MPSS						
Alberti 2015	61/244 (25%)	0	NA	61/244 (25%)	NR	NR

Benachi 2015	42/892 (4.7%)	42 (100%) with second	6/42 (14.3%)	6/892 (0.7%)	2.7%	0.4%
		aliquot				
Bianchi 2012	16/519 (3.1%)	0	NA	16/356 (3.1%)	NR	NR
Bianchi 2014a	18/1970 (0.9%)	0c	NA	T21 and T18: 18/1970 (0.9%)	NR	NR
				T13: 18/1932 (0.9%)		
Bijok 2014	1/10 (10.0%)	0	NA	1/10 (10.0%)	50%	0%
Chiu 2011	11/764 (1.4%)	0	NA	11/764 (1.4%)	NR	NR
Ehrich 2011	20/467 (4.3%)	20 (100%) re- sequenced	18/20 (90%)	18/467 (3.9%)	NR	NR
Fiorentino 2016	100/7103 (1.4%)	79 (79%) with new	0 (0%)	21/7103 (0.3%)	0%	0.3%
		sampling				
Johansen 2016	NR	2 with second aliquot or	NR	11/184 (6%) ^d	5.8%	6.1%
		resequenced were in the				
		grey zone (be- tween				
		affected and unaffected)				
Lee 2015	1/93 (1.1%)	0	NA	1/93 (1.1%)	NR	NR
Lefkowitz 2016	Autosomes: 42/1208 (3.5%)	0	NA	Autosomes: 42/1208 (3.5%)	Autosomes: 3.8%	Autosomes 3.4%
	SCA: 64/1208 (5.3%)			SCA: 64/1208 (5.3%)	SCA: 29.7%	SCA: 4.5%
Liang 2013	12/424 (2.8%)	0	NA	12/424 (2.8%)	NR	NR
Ma 2016	5/10,584 (0.05%)	0	NA	5/10,584 (0.05%)	NR	NR
Mazloom 2013	21/432 (4.9%)	0	NA	21/432 (4.9%)	11.8%	4.3%
Palomaki 2012	110/1988 (5.5%)	105 (95.5%) with second	17/110 (15.5%)	17/1988 (0.9%)	1.0%	0.8%
		aliquot and 5 (4.5%)				
		resequenced				
Papageorghiou 2016a	5/431 (1.2%)	0	NA	5/431 (1.2%)	NR	NR

Table 7. Proportion of pregnant women with a reference standard and assay failure during gNIPT process (Continued)

Table 7. Proportion of pregnant women with a reference standard and assay failure during gNIPT process (Continued) Papageorghiou

2016b

20100						
Poon 2016	1/242 (0.4%)	0	NA	1/242 (0.4%)	0%	0.5%
Porreco 2014	Autosomes:	0	NA	Autosomes: 108/3430	NR	NR
	108/3430 (3.1%)			(3.1%)		
	45,X and 47,XXX:			45,X and 47,XXX: 152/3430 (4.4%)		
	152/3430 (4.4%)			47,XXY and 47,XYY:		
	47,XXY and 47,XYY:			229/3430 (6.7%)		
	229/3430 (6.7%)					
Song 2013	73/1814 (4.0%)	0	NA	73/1814 (4.0%)	0%	4.0%
Song 2015	1/205 (0.5%)	0	NA	1/205 (0.5%)	NR	NR
Stumm 2014	32/504 (6.3%)	0	NA	32/504 (6.3%)	3.5%	6.7%
Sung-Hee 2015	21/908 (2.3%)	16 (76.2%) with new	2/16 (12.5%)	7/908 (0.8%)	NR	NR
		sampling				
Tynan 2016	52/1100 (4.7%)	0	NA	52/1100 (4.7%)	0%	4.9%
Yao 2014	0	0	NA	0	NA	NA
Zhou 2014a	0	0	NA	0	NA	NA
Zhou 2014b	141/3954 (3.6%)	141 (100%) with new	4/141 (2.8%)	4/3954 (0.1%)	NR	NR
		sampling				
Overall range of f	inal assay failure for MI	PSS		0% to 25%	0% to 50%	0% to 6.7%
TMPS						
Ashoor 2012	3/400 (0.8%)	0	NA	3/400 (0.8%)	0%	1%
Ashoor 2013	53/2002 (2.6%)	0	NA	53/2002 (2.6%)	0%	2.7%
Bevilacqua 2015	29/356 (8.1%)	26 (90%) with 2 nd	13/26 (50%)	16/356 (4.5%)	NR	NR
		aliquot				
Comas 2015	9/316 (2.8%)	6 (67%) with new	1/6 (16.7%)	4/316 (1.3%)	NR	NR
		sampling				
del Mar Gil 2014	15/207 (7.2%)	0	NA	15/207 (7.2%)	23%	6%

Table 7. Proportion of pregnant women with a reference standard and assay failure during gNIPT process (Continued)

Gil 2016	99/3698 (2.8%)	54 (54,5%) with new	20/54 (37%)	65/3698 (1.8%)	NR	NR
		sampling				
Hall 2014	4/68 (5.9%)	0	NA	4/68 (5.9%)	11.8%	3.9%
Hooks 2014	18/432 (4.2%)	0	NA	18/432 (4.2%)	NR	NR
Jackson 2014	NR	NR	14 (NR)	14/1175 (1.2%)	NR	NR
Nicolaides 2012	100/2049 (4.9%)	0	NA	100/2049 (4.9%)	9.1%	4.9%
Nicolaides 2013	13/242 (5.4%)	0	NA	13/242 (5.4%)	6.3%	5.2%
Nicolaides 2014a	5/177 (2.8%)	0	NA	5/177 (2.8%)	5.1%	1.7%
Norton 2012	148/3228 (4.6%)	0	NA	148/3228 (4.6%)	NR	NR
Norton 2015	488/16,329 (3.0%)	0	NA	488/16,329 (3.0%)	20.6%	2.9%
Pergament 2014	T21: 88/1051 (8.4%)	0	NA	T21: 88/1051 (8.4%)	All five	All five
	T18, 45,X: 87/1052 (8.3%)			T18, 45,X: 87/1052 (8.3%)	chromo- somes	chromo somes
	T13: 86/1053 (8.2%)			T13: 86/1053 (8.2%)	(n = 85): 15.2%	(n = 85): 7.1%
Persico 2016	10/259 (3.9%)	0	NA	10/259 (3.9%)	8.4%	2.1%
Quezada 2015	122 ^e /2838 (4.2%)	110 (90.1%) with new	41/110 (37.3%)	53/2838 (1.9%)	4.1%	1.8%
		sampling				
Saman- go-Sprouse 2013	15/201 (7.5%)	0	NA	15/201 (7.5%)	6.3%	7.6%
Verweij 2013	51/520 (9.8%)	51 (100%)	16/51	16/520 (3.1%)	NR	NR
		with 2 nd	(31.4%)	NR		
		aliquot				
Overall range of f	inal assay failure for TM	IPS		0.8% to 7.5%	0% to 23%	1% to 7.63%

CVS: chorionic villi sampling, FF: fetal fraction DNA, GA: gestational age, NA: not applicable, NR: not reported by authors, QC: quality control. ^aRepeated tests included second aliquot (aliquot from first sampling), resequenced (same library) or new sampling.

^baneuploid: proportion of failed samples of aneuploid cases out of all aneuploid tested with reference standard and gNIPT result. euploid: proportion of failed samples of euploid cases out of all euploid tested with reference standard and gNIPT result.

^cAuthors decided to resequence 12 samples with gNIPT results. They were in the grey zone (between affected and unaffected) and were resequenced in uniplex. All repeated tests were in affected or unaffected zone.

^dOnly the final failure rate was reported. The failure rate at first attempt was not reported nor the failure rate of repeated tests. ^eAuthor reported 123 failed tests but this number included one sample lost in the mail and so did not undergo the sequencing process.



Table 8. Data for 47,XXX, 47,XXY and 47,XYY according to the prior risk of fetal aneuploidy and gNIPT approach

Test		Number of	Number of	Number of
		studies	affected preg- nancies	unaffected pregnan- cies ^a
47,XXX				
Selected high risk	MPSS	5	8	5441
pregnant women	TMPS	2	6	580
47,XXY				
Selected high risk	MPSS	7	14	6466
pregnant women	TMPS	3	8	827
47,XYY				
Selected high risk	MPSS	7	11	6418
pregnant women	TMPS	1	3	169

^{*a*}Unaffected pregnancies: we included pregnancies with any other aneuploidy than the one under analysis with all euploid cases as "unnon affected".

Test subgroups		Number of	Number of	Number of	Sensitivity ^b	Specificity ^b
		studies	affected	unaffected	% (95% CI)	% (95% CI)
			pregnan- cies	pregnan- cies ^a		
Pregnancy	y type					
Autosome	s (T21, T18 and T1	13 combined), u	nselected popu	lation		
MPSS	singleton	1	11	1730	100 (74.1 to 100)	99.9 (99.7 to 100)
TMPS	singleton	3	107	20,468	95.5 (87.4 to 98.4)	99.9 (99.8 to 100)
	multifetal	1	11	181	90.9 (62.3 to 98.4)	100 (97.9 to 100)
Autosome	s (T21, T18 and T1	13 combined), se	elected high-ris	k population		
MPSS	singleton	19	1087	11,180	98.3 (97.3 to 98.9)	99.6 (99.5 to 99.7)
	multifetal	3	21	206	95.2 (72.9 to 99.3)	100 (98.2 to 100) ^c
TMPS	singleton	7	378	4282	98.9 (97.2 to 99.6)	99.9 (99.8 to 100)
SCA (45,X,	47,XXX, 47,XXY a	nd 47,XYY comb	ined), selected	high-risk popu	lation	
MPSS	singleton	7	101	4690	88.3 (52.9 to 98.1)	99.3 (97.5 to 99.8)

Table 9. Subgroup analyses of MPSS and TMPS (type of pregnancy and gestational age)

TMPS		4	96	968	93.8 (86.8 to 97.2)	99.6 (98.1 to 99.9)
Gestation	al age					
Autosome	s (T21, T18 and T1	3 combined	l), unselected p	opulation		
MPSS	≤29 weeks	1	11	1730	100 (74.1 to 100)	99.9 (99.7 to 100)
TMPS	≤15 weeks	4	118	20,649	94.9 (89.1 to 97.7)	99.9 (99.8 to 99.9)
Autosome	s (T21, T18 and T1	3 combined	l), selected high	risk population	1	
MPSS	≤15 weeks	3	49	532	100 (92.7 to 100) ^c	100 (99.3 to 100) ^c
	≤29 weeks	12	594	4605	98.3 (96.9 to 99.1)	99.3 (99.0 to 99.5)
	≤42 weeks	13	729	7831	98.9 (95.0 to 99.8)	99.9 (99.8 to 99.9)
TMPS	≤15 weeks	2	128	498	99.2 (95.7 to 99.9) ^c	100 (99.2 to 100) ^c
	≤29 weeks	2	33	535	97.0 (84.7 to 99.5) ^c	100 (99.3 to 100) ^c
	≤42 weeks	2	163	3084	99.4 (95.8 to 99.9)	99.9 (99.7 to 100)
SCA (45,X,	47,XXX, 47,XXY a	nd 47,XYY c	ombined), selec	ted high-risk po	pulation	
MPSS	≤15 weeks	1	2	202	0.00 (0.00 to 65.8)	99.5 (97.2 to 99.9)
	≤29 weeks	5	58	996	86.5 (63.1 to 96.0)	95.1 (93.5 to 96.3)
	≤42 weeks	5	89	6103	95.8 (80.3 to 99.2)	99.6 (99.4 to 99.7)
TMPS	≤15 weeks	2	58	343	93.1 (83.0 to 97.4)	99.7 (98.0 to 100)
	≤42 weeks	1	34	380	97.1 (85.1 to 99.5)	98.9 (97.3 to 99.6)

45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, T21: trisomy 21, T18: trisomy 18, T13: trisomy 13 CI: confidence interval, MPSS: massively parallel shotgun sequencing, SCA: sex chromosome aneuploidies, TMPS: targeted massively parallel sequencing.

*a*We included pregnancies with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies. *b*For two or more studies, the sensitivities and specificities are the summary estimates obtained from meta-analysis. *c*Simple pooling used to obtain summary estimates of sensitivity, specificity or both.

Table 10. Direct comparisons of gNIPT and traditional screening tests for autosomes (T21, T18 and T13 combined) in unselected population of pregnant women undergoing aneuploidy screening

Study	Sensitivity (es) %	true positives/cas-	Difference % (95% CI)	Specificity (tru %	e negatives/unaffected ^a)	Difference % (95% Cl)
	MPSS	Traditional screening tests		MPSS	Traditional screening tests	

Table 10. Direct comparisons of gNIPT and traditional screening tests for autosomes (T21, T18 and T13 combined) in unselected population of pregnant women undergoing aneuploidy screening (Continued)

		-			-	
Song 2013	100 (11/11)	54.6 (6/11)	45.5 (10.0 to 72.0)	99.9 (1729/1730)	86.0 (1487/1730)	14.0 (12.4 to 15.7)
	TMPS	Traditional screening tests		TMPS	Traditional screening tests	
Nicolaides 2012	100 (10/10)	100 (10/10)	0.00 (-27.8 to 27.8)	99.9 (1937/1939)	95.5 (1852/1939)	4.38 (3.51 to 5.40)
Norton 2015	98.0 (49/50)	78.0 (39/50)	20.0 (7.44 to 33.3)	99.9 (15,779/15,791)	94.1 (14,860/15,791)	5.82 (5.46 to 6.20)
Quezada 2015	91.5 (43/47)	100 (49/49)	-8.51 (-19.9 to 0.40)	99.7 (2730/2738)	95.6 (2663/2787)	4.16 (3.40 to 5.00)

CI: confidence interval, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing. *a*We included pregnancies with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies.

Test	Number of	Number of	Number of	Summary sensitivity	Summary specificity	P value ^b
	studies	affected	unaffected	% (95% CI)	% (95% CI)	
		pregnan- cies	pregnan- cies ^a			
Case-cont	trol studies exclu	ded				
Autosome	es (T21, T18 and T	13 combined), s	selected high-ris	sk population		
MPSS	22	696	11,293	98.3 (95.1 to 99.4)	99.9 (99.8 to 100)	0.72
TMPS	4	219	3,813	98.6 (95.8 to 99.6)	99.9 (99.8 to 100)	
SCA (45,X,	, 47,XXX, 47,XXY a	ind 47,XYY com	bined), selected	l high-risk population		
MPSS	10	98	5,872	91.9 (73.8 to 97.9)	99.5 (98.8 to 99.8)	0.41
TMPS	2	6	472	93.8 (86.8 to 97.2)	99.6 (98.1 to 99.9)	
Exclusion	of studies with l	ess than 10 pre	gnancies with	aneuploidy		
Autosome	es (T21, T18 and T	13 combined), s	selected high-ris	sk population		
MPSS	21	1458	13,921	98.7 (96.8 to 99.4)	99.8 (99.5 to 100)	0.07
TMPS	7	378	4,282	98.9 (97.2 to 99.6)	99.9 (99.8 to 100)	
SCA (45,X,	, 47,XXX, 47,XXY a	ind 47,XYY com	bined), selected	l high-risk population		
MPSS	6	130	5,761	94.5 (80.6 to 98.6)	99.4 (97.6 to 99.8)	0.28
TMPS	2	90	496	94.4 (87.3 to 97.7)	99.0 (97.6 to 99.6)	_



45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, T21: trisomy 21, T18: trisomy 18, T13: trisomy 13 CI: confidence interval, MPSS: massively parallel shotgun sequencing, SCA: sex chromosome aneuploidies, TMPS: targeted massively parallel sequencing.

^{*a*}We included pregnancies with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies. ^{*b*}The P value indicates the statistical significance of the difference in model fit and was obtained from likelihood ratio tests comparing models with and without a covariate for test type.

APPENDICES

Appendix 1. Screening tests and medical terms glossary^a

Terms	Definitions
Amniocentesis	Invasive procedure under continuous ultrasound guidance (performed between 15 to 19 weeks of gestational age). A sterile needle is passed through the mother's abdomen, uterus and amniotic sac. A sample of fetal cells present in the amniotic fluid surrounding the fetus is aspirated with a syringe and sent for analysis to test for a range of chromosomal and inherited disorders.
Aneuploidy	The state of having a different (additional or missing) number of chromosomes than the 23 pairs normally present in humans.
Attention deficit disorder (ADD)	ADD is a neurodevelopmental disorder defined by impairing levels of inattention and disorganisa- tion. Inattention manifests behaviourally in ADD as wandering off task, lacking persistence, having difficulty sustaining focus and being disorganised.
Case-control study	In the context of diagnostic accuracy, existing records are used to identify a group of people known to have the target condition (cases) and another group (controls) without the target condition. The control group may consist of healthy individuals or those with other conditions similar to the target condition Cases and controls are then compared with respect to certain variables hypothesised to increase the risk of having the disease.
Chorionic villus sampling (CVS)	An abdominal or cervical procedure performed under continuous ultrasound guidance to obtain a sample of placental tissue for chromosomal or genetic analysis (between 12 to 19 weeks of ges- tational age). The range of chromosomal and genetic conditions that can be detected is similar to those for amniocentesis.
Clinodactyly	Permanent deflection of one or more fingers.
Cut-off	Synonyms: cutpoint or threshold.
Cutpoint	A value for a test result measured on an ordinal or continuous scale which divides the group of peo- ple tested into a group at lower risk of the condition being screened for and a group at higher risk (for whom further investigations may be offered). Synonyms: cut-off or threshold.
Detection rate	The proportion of affected individuals with a positive screening result. The detection rate is the same as the sensitivity of a test.
Developmental delay	An individual with this neurodevelopmental disorder fails to meet expected developmental mile- stones in several areas of intellectual functioning.
Diagnostic accuracy	The ratio of true positive and true negative results to the total number of test results (true posi- tives, true negatives, false positives and false negatives). Represents the level of agreement be- tween the information from the index test and the reference standard.



(Continued)

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(continueu)	
Diagnostic test	A test recognised as having best performances to provide sufficient information allowing a definite diagnosis (as opposed to screening test results that need to be confirmed before a final diagnosis can be reached).
Dysgenesis	Defective or abnormal formation of an organ or part, primarily during embryonic development. Gonodal dysgenesis is a defective development of the gonads, which may be accompanied by ab- normalities of the sex chromosomes.
False negative	A negative test result in someone with the target condition.
False positive	A positive test result in someone without the target condition.
Fluorescence in situ Hybridisation	Describes a type of DNA analysis by the hybridisation of fluorescently-labelled probes complemen- tary to certain genomic regions. In the context of fetal aneuploidy detection, describes a diagnos- tic test in which chromosome-specific fluorescently-labelled DNA probes are used on uncultured cells from chorionic villi or amniotic fluid to assess the number of homologous chromosome copies present.
High risk of fetal aneuploidy	A pregnancy is considered at high risk of fetal aneuploidy if the result of the prenatal screening test puts the fetus at increased risk for aneuploidy.
Hypertelorism	Abnormal distance between two paired organs.
Intellectual disability or intellectual developmental disorder	This neurodevelopmental disorder is characterised by deficits in general mental abilities, such as reasoning, problem solving, planning, abstract thinking, judgment, academic learning and learning from experience. It is a condition diagnosed before age 18. In the past, the term mental retardation was used to describe this condition but this term is no longer used.
Invasive procedure	A method used to obtain a biological sample that involves significant disruption of the physical in- tegrity of a patient. Examples include amniocentesis and chorionic villi sampling.
Karyotype	A photomicrograph of an individual's chromosomes arranged in a standard format and visualised by various staining methods, showing the number, size, and shape of each chromosome; used to correlate chromosomal anomalies with specific diseases (karyotyping). In humans, there are a total of 23 pairs of homologous chromosomes (total of 46 chromosomes).
Learning disability	Learning disability refers to inadequate development of specific academic, language and speech skills such as reading disability, mathematics disability and writing disability.
Low risk of fetal aneuploidy	A pregnancy is considered at low risk of fetal aneuploidy if the result of the prenatal screening test puts the fetus at decreased risk for aneuploidy.
Meta-analysis	The use of statistical techniques in a systematic review to integrate the results of included studies. Sometimes misused as a synonym for systematic reviews which may or may not include a meta- analysis.
Mosaic	An individual who has some cells with an unusual genetic or chromosomal make-up while the rest of the cells in the body have the typical genetic or chromosomal constitution.
Mixed risk population	Mixed risk population included a mixture of selected pregnant women with low, high or no prior risk of fetal aneuploidy.
Negative predictive value	A measure of test performance. Defined as the proportion of people with a negative test result who do not have the target condition.



(Continued)	
Nuchal translucency scan	The thickness of fluid in the tissue space within the nape of the fetal neck typically measured by ul- trasonography. An increased amount of fluid is associated with Down syndrome and other struc- tural or genetic anomalies.
Positive predictive value	A measure of test performance. Defined as the proportion of people with a positive test result who do have the target condition.
Probability	The chance or risk of an event happening.
Prospective study	A study in which a group of individuals is followed through time in order to detect the occurrence of a disease or another outcome of interest.
Reference standard	The best available test to detect the presence or absence of the target condition.
Retrospective study	A study in which all or part of the data collection occurred before initiation of the study.
Screening	Testing asymptomatic people for the likelihood of the presence of a disease, either with the aim of reducing risk of an adverse outcome, or with the aim of giving information about risk.
Seizure	A sudden attack, spasm, or convulsion caused by abnormal electrical conduction in the brain.
Sensitivity	A measure of test performance. Defined as the proportion of individuals with the target condition who have a positive test result. Higher sensitivity values means that a higher proportion of affected individuals will be detected by the test (few false negatives).Sensitivity is the same as the detection rate.
Single nucleotide polymorphism	Single nucleotide polymorphisms are the most common type of genetic variation among people. A difference in a single DNA nucleotide (A, T, C or G) in a DNA sequence.
Specificity	A measure of test performance. Defined as the proportion of individuals without the target condi- tion who have a negative test result. Higher specificity values means that a smaller proportion of unaffected individuals will be wrongly classified as having the target condition (few false positives).
Threshold	Synonyms: cutpoint or cut-off.
True negative	An individual with a negative test result who does not have the target condition.
True positive	An individual with a positive test result who has the target condition.
Trisomy	Three copies of a particular chromosome rather than the usual pair.
Unselected pregnant women	A pregnant women who did not undergo any prenatal screening test at the time of enrolment.

^aAdapted in part from the United Kingdom National Screening Committee Glossary, MedlinePlus Medical Encyclopedia, American Psychiatric Association and The Cochrane Collaboration's Glossary of terms (APA 2013; Cochrane Glossary 2014; MedlinePlus 2014; UK Screening Glossary 2012).

Appendix 2. List of acronyms and abbreviations

Acronyms or abbreviations	Terms
45,X	monosomy X or Turner syndrome



(Continued)	
47,XXX	trisomy X or triple X syndrome
47,XXY	Klinefelter syndrome
aCGH	array comparative genomic hybridisation
AFP	alpha-fetoprotein
Bioch/US	biochemical or ultrasound or both screening test
ccfDNA	circulating cell-free DNA
Chrom. 21	chromosome 21
CVS	chorionic villi sampling
FISH	fluorescence in situ hybridisation
gNIPT	genomics-based non-invasive prenatal testing
hCG	human chorionic gonadotropin
HSROC	hierarchical summary receiver operating characteristic
MPSS	massively parallel shotgun sequencing
NA	not applicable
ND	no data available
NGS	next generation sequencing
NR	not reported
NT	nuchal translucency
PAPP-A	pregnancy associated plasma protein A
QF-PCR	quantitative fluorescent polymerase chain reaction
QUADAS-2	QUality Assessment of Diagnostic Accuracy Studies
Ref. Chrom	reference chromosome
RS	reference standard
SCA	sex chromosome aneuploidy
SNP	single nucleotide polymorphism
T13	trisomy 13 or Patau syndrome
T18	trisomy 18 or Edward syndrome
T21	trisomy 21 or Down syndrome

Cochrane
Library

(Continued)	
TMPS	targeted massively parallel sequencing
uE3	unconjugated estriol

Appendix 3. Index test technical details

Typically, blood samples from pregnant women are obtained by venous puncture in the first or second trimester. After two centrifugation steps, plasma is separated from maternal whole blood and ccfDNA is extracted from plasma with commercial kits. DNA is converted into a genomic library where each of the DNA fragments are ligated with platform specific adapters. For TMPS only, libraries are clonally amplified before being sequenced. Then the libraries of several pregnant women are loaded on a next generation sequencer. The produced sequencing reads are aligned on a reference human genome to their respective chromosomal location and the number of sequence reads from each chromosome is computed (Rothberg 2011). MPSS randomly sequences DNA fragments from across the whole genome while TMPS sequences DNA fragments from selected regions (Figure 1). Ultimately, all gNIPT for aneuploidies rely on assigning sequence reads of DNA fragments to their chromosome of origin and comparing total number or proportions of reads or single nucleotide polymorphisms (SNP) genotype between each chromosome of interest (e.g. 13, 18, 21, X and Y) and a reference set of chromosomes. A Z score (or other statistics) are computed and a patient-specific risk can be assessed based on a risk threshold determined from read counts from a series of known euploid and aneuploid pregnancies. For MPSS, the counts from chromosomes of interest are normalised using the counts from all other chromosome sequences, while, for TMPS, the counts are normalised against a subset of selected sequences. Bioinformatic approaches vary according to the testing approach (MPSS or TMPS) and research team. Besides the use of normalised chromosome read counts, TMPS also allows for the use of additional allelic information when polymorphic loci such as SNP are targeted, such as an estimate of fetal DNA concentration (fetal DNA proportion) (Liao 2012). Thus, while MPSS produces a larger number of total sequence reads, TMPS will generates a larger number of reads from each targeted chromosomes.

Appendix 4. Search strategy

MEDLINE (Ovid)		
Steps	Text words and subject headings	Sets of search
1	'cell-free dna'.mp	Index test
2	'cell free dna'.mp	
3	cfdna.mp	
4	ffdna.mp	
5	cffdna.mp	
6	'free foetal dna'.mp	
7	'free fetal dna'.mp	
8	nipd.mp	
9	nipt.mp	
10	(non invasive or noninvasive or non-invasive).mp	
11	(genetic adj2 (diagnos* or detect* or test* or screen*)).mp	
12	exp Genetic Testing/	
13	exp Sequence Analysis, DNA/	



(Continued)		
14	((antenatal or ante natal) adj2 (diagnos* or detect* or test* or screen*)).mp	
15	((prenatal or pre natal) adj2 (diagnos* or detect* or test* or screen*)).mp	_
16	exp Prenatal Diagnosis/	
17	or/1-16	-
18	maternal.mp	Patient description
19	exp Pregnancy/	
20	exp Pregnancy Complications/	
21	pregnant.mp	
22	pregnanc*.mp	
23	exp Fetus/	_
24	fetus.mp	-
25	foetus.mp	
26	fetal.mp	-
27	foetal.mp	-
28	or/18-27	-
29	trisom*.mp	Target condition
30	aneuploid*.mp	-
31	(down* adj syndrome*).mp	
32	exp Aneuploidy/	-
33	exp Trisomy/	-
34	exp Down Syndrome/	-
35	chromosome disorders.mp	-
36	or/29-35	-
37	or/1-5,11-13	Combined sets
38	6 or 7 or 8 or 9 or 14 or 15 or 16	-
39	36 and 37 and 28 and 10	-
40	38 and 37 and 36	-
41	39 or 40	Final combined set



Embase (Embase.com)

Steps	Text words and subject headings	Sets of search
1	'cell-free dna'	Index test
2	'cell free dna'	
3	cfdna	
4	ffdna	
5	cffdna	
6	'free foetal dna'	
7	'free fetal dna'	
8	nipd	
9	nipt	
10	'non invasive'	
11	noninvasive	
12	'non-invasive'	
13	genetic NEXT/1 (diagnos* or screen* or test* or detect*)	
14	'genetic screening'/exp	
15	'genetic testing'/exp	
16	'sequence analysis dna'/exp	
17	antenatal NEXT/1 (diagnos* or screen* or test* or detect*)	
18	prenatal NEXT/1 (diagnos* or screen* or test* or detect*)	
19	pre?natal NEXT/1 (diagnos* or screen* or test* or detect*)	
20	'prenatal diagnosis'/exp	
21	1 OR 2 OR 3 OR 4 OR 5 OR 6 OR 7 OR 8 OR 9 OR 10 OR 11 OR 12	
	OR 13 OR 14 OR 15 OR 16 OR 17 OR 18 OR 18 OR 19 OR 20	
22	antenatal	Patient description
23	prenatal	
24	pre?natal	



25 maternal 26 foetus 27 fetus 28 foetal 29 fetal 30 pregnant* 31 pregnant 32 'pregnant woman'/exp 34 'pregnant woman'/exp 35 'pregnant woman'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 30 OR 31 OR 35 37 trisom* 38 aneuploid* 39 down' NEXT/1 syndrome 40 'aneuploidy'/exp 41 'aneuploidy/exp 42 'trisomy'/exp 43 'down syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 30 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45 47 21 and 36 and 46	(Continued)		
27 fetus 28 foetal 29 fetal 30 pregnanc* 31 pregnant 32 'pregnancy/exp 33 'pregnancy complications'/exp 34 'pregnant woman'/exp 35 'pregnant woman'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 37 trisom* 38 aneuploid* 39 down* NEXT/1 syndrome 40 'aneuploidy'/exp 41 'aneuploidy'/exp 42 'trisomy'/exp 43 'down syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	25	maternal	
28 foetal 29 fetal 30 pregnanc* 31 pregnant 32 'pregnancy'/exp 33 'pregnancy complications'/exp 34 'pregnant woman'/exp 35 'pregnant woman'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 37 trisom* 38 aneuploid* 39 down* NEXT/1 syndrome 40 'aneuploid/'exp 41 'aneuploid//exp 42 'trisomy'/exp 43 'downs syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	26	foetus	
29fetal30pregnanc*31pregnant32'pregnancy/exp33'pregnancy complications'/exp34'pregnant woman'/exp35'pregnant women'/exp3622 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 3537trisom*38aneuploid*39down* NEXT/1 syndrome40'aneuploid/'exp41'aneuploid/'exp42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	27	fetus	
30 pregnanc* 31 pregnant 32 'pregnancy'(exp 33 'pregnancy complications'/exp 34 'pregnant woman'/exp 35 'pregnant woman'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 37 trisom* 38 aneuploid* 39 down* NEXT/1 syndrome 41 'aneuploidy'/exp 42 'trisomy'/exp 43 'downs syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	28	foetal	
31 pregnant 32 'pregnancy'/exp 33 'pregnancy complications'/exp 34 'pregnant woman'/exp 35 'pregnant woman'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 37 trisom* 38 aneuploid* 39 down* NEXT/1 syndrome 40 'aneuploid'/exp 41 'aneuploidy'/exp 43 'downs syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	29	fetal	
32 'pregnancy'/exp 33 'pregnancy complications'/exp 34 'pregnant woman'/exp 35 'pregnant women'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 37 trisom* 38 aneuploid* 39 down* NEXT/1 syndrome 40 'aneuploidy'/exp 41 'aneuploidy'/exp 42 'trisomy'/exp 43 'downs syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	30	pregnanc*	
33 'pregnancy complications'/exp 34 'pregnant woman'/exp 35 'pregnant women'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 37 trisom* 38 aneuploid* 39 down* NEXT/1 syndrome 40 'aneuploid'/exp 41 'aneuploidy'/exp 42 'trisomy'exp 43 'down syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	31	pregnant	
34 'pregnant woman'/exp 35 'pregnant women'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 37 trisom* 38 aneuploid* 39 down* NEXT/1 syndrome 40 'aneuploid//exp 41 'aneuploid//exp 42 'trisomy'/exp 43 'down syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	32	'pregnancy'/exp	
35 'pregnant women'/exp 36 22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35 Target condition 37 trisom* Target condition 38 aneuploid* 1000000000000000000000000000000000000	33	'pregnancy complications'/exp	
3622 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29 OR 30 OR 31 OR 32 OR 33 OR 34 OR 35Target condition37trisom*Target condition38aneuploid*39down* NEXT/1 syndrome40'aneuploid'/exp41'aneuploidy'/exp42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	34	'pregnant woman'/exp	
OR 30 OR 31 OR 32 OR 33 OR 34 OR 3537trisom*Target condition38aneuploid*39down* NEXT/1 syndrome40'aneuploid'/exp41'aneuploidy'/exp42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	35	'pregnant women'/exp	
37trisom*Target condition38aneuploid*39down* NEXT/1 syndrome40'aneuploid'/exp41'aneuploidy'/exp42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	36	22 OR 23 OR 24 OR 25 OR 26 OR 27 OR 28 OR 29	
38 aneuploid* 39 down* NEXT/1 syndrome 40 'aneuploid'/exp 41 'aneuploidy'/exp 42 'trisomy'/exp 43 'downs syndrome'/exp 44 'down syndrome'/exp 45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45		OR 30 OR 31 OR 32 OR 33 OR 34 OR 35	
39down* NEXT/1 syndrome40'aneuploid'/exp41'aneuploidy'/exp42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	37	trisom*	Target condition
40'aneuploid'/exp41'aneuploidy'/exp42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	38	aneuploid*	
41'aneuploidy'/exp42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	39	down* NEXT/1 syndrome	
42'trisomy'/exp43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	40	'aneuploid'/exp	
43'downs syndrome'/exp44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	41	'aneuploidy'/exp	
44'down syndrome'/exp45'chromosome disorders'4637 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	42	'trisomy'/exp	
45 'chromosome disorders' 46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	43	'downs syndrome'/exp	
46 37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	44	'down syndrome'/exp	
	45	'chromosome disorders'	
47 21 and 36 and 46 Final combined set	46	37 OR 38 OR 39 OR 40 OR 41 OR 42 OR 43 OR 44 OR 45	
	47	21 and 36 and 46	Final combined set

Web of Science (ISI)

Steps Text words and	subject headings	Sets of search
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(Continued)		
1	TOPIC: (down* syndrome) OR TOPIC: (trisom*) OR TOPIC: (aneuploid*)	Target condition
2	TOPIC: (pregnan*)	Patient description
3	TOPIC: (dna) OR TOPIC: (blood)	Index test
4	TOPIC: (pre?natal screen*) <i>OR</i> TOPIC: (prenatal screen*) <i>OR</i> TOPIC: (pre?natal test*)	Index test
	OR TOPIC: (prenatal test*) OR TOPIC: (genetic test*) OR TOPIC: (genetic screen*)	
	<i>OR</i> TOPIC: (prenatal diagnos*) <i>OR</i> TOPIC: (pre?natal diagnos*) <i>OR</i> TOPIC: (detection)	
	<i>OR</i> TOPIC: (genetic diagnos*) <i>OR</i> TOPIC: (non invasive) <i>OR</i> TOPIC: (non-invasive)	
	OR TOPIC: (noninvasive)	
5	#1 AND #2 AND #3 AND #4	Final combined set

Cochrane Register of Diagnostic Test Accuracy Studies

Hand search: in diagnostic test accuracy database, there are 18 publications from the Cochrane Pregnancy and Childbirth group.

Clinicaltrials.gov

Steps	Text words and subject headings	Sets of search
1	(down syndrome OR trisomy OR aneuploidy)	Target condition
2	(testing OR screening OR diagnosis OR detection)	Index test
3	#1 AND #2	Final combined set

European Clinical Trials Register		
Text words and subject headings	Sets of search	
pregnan*	Population	
trisom* OR aneuploid*	Target condition	
#1 OR #2	Final combined set	
	Text words and subject headings pregnan* trisom* OR aneuploid*	Text words and subject headings Sets of search pregnan* Population trisom* OR aneuploid* Target condition



Who ICTRP

Steps	Text words and subject headings	Sets of search
1	screen* OR detect* OR diagnos* OR test* OR pregnan*	Index test and population
2	«down syndrome » OR trisom* OR aneuploid*	Target condition
3	#1 AND #2	Final combined set

NTIS.gov	NTIS.gov		
Steps	Text words and subject headings	Sets of search	
1	(down syndrome OR trisomy OR aneuploidy)	Target condition	
2	(testing OR screening OR diagnosis OR detection)	Index test	
3	#1 AND #2	Final combined set	

OpenGrey		
Steps	Text words and subject headings	Sets of search
1	"down syndrome" OR trisom* OR aneuploid*	Target condition
2	screen* OR detect* OR diagnos* OR test* OR pregnan*	Index test and population
3	#1 AND #2	Final combined set

National Guideline Clearing House (NGCH)		
Text words and subject headings	Sets of search	
diagnosis (guideline category)	Index test	
screening (guideline category)	Index test	
aneuploid OR trisomy OR « down syndrome »	Target condition	
(#1 OR #2) AND #3	Final combined set	
	Text words and subject headings diagnosis (guideline category) screening (guideline category) aneuploid OR trisomy OR « down syndrome »	



TheseNet

Steps	Text words and subject headings	Sets of search
1	trisomy	Target condition
2	screening	Index test
3	#1 AND #2	Final combined set

These Canada P	ortal	
Steps	Text words and subject headings	Sets of search
1	trisomy OR (down AND syndrome)	Target condition
2	screening	Index test
3	#1 AND #2	Final combined set

Appendix 5. Data collection form for study classification during full-text assessment

Heading	Detailed instructions	Data
Study ID	Last name of the first author and year of publication	Name:
		Year:
Reference details	Details allowing identification of the publication	Journal:
		Volume:
		Issue:
		Pages:
		Accession number (e.g. PMID ^a):
Multiple reports	For example, duplicate publications or follow-up studies.	Study ID:
of this study	Provide the study ID linked to this classified study	
Type of report	Check the appropriate box	Journal article #
		Conference/abstract #
		Ongoing trial #
		Others #
		Specify:
Language	In which language was the study written	Language:



(Continued)		Translation needed? Yes # No #
Eligibility	Provide reason for exclusion or awaiting classification (e.g. why authors should be contacted and what issues should be clarified)	Study excluded? Yes # No # Reason: Awaiting classification? Yes # No # Reason:
Report author contact details for further information	Date when the authors were contacted (<i>dd/mm/yyyy</i>)	No need for further contact # Authors have been contacted on: Reply received on:
Review author ID	Who completed the form	Name:
Date of classification	(dd/mm/yyyy)	Date:
Notes, questions or reminders		

^aPMID: PubMed identifier.

Appendix 6. QUADAS-2 tool for assessing methodological quality of included studies

	Signalling question	Signalling question	Signalling question	Risk of bias	Concerns about applicability
Domain 1: Pa	atient selection				
Patient	Was a consecutive or random	Was a case-con-	Did the study	Could the se-	Are there concerns
selection	sample of patients enrolled?	trol design avoid- ed?	avoid inap- propriate ex- clusions?	lection of pa- tients have in- troduced bias?	that the included patients and setting do not match the re view question?
	Yes: if all consecutive or ran- dom samples or convenient samples or all eligible pregnant women were enrolled.	Yes: if a case-con- trol design was avoided.	Yes: if the study avoided inappropriate exclusions.	Low risk: if 'yes' for all sig- nalling ques- tion.	Low concern: if the selected pregnant women represent the women indicated
	No: if selected pregnant women were enrolled.		High or un- clear risk: if	by the review ques- tion ^a .	
Unclear: if this was not clearUnclear: if thisfrom the report.was not clear from the report.	was excluded based on fam- ily's situation, maternal age, ethnicity, ma- ternal cancer history, type	'no' or 'unclear' was reported for at least one signalling ques- tion.	High concern: if selected pregnant women differ from those targeted by the review question ^a . Unclear concern: if		
			of pregnan- cy, gestational age, assisted reproductive technology or		insufficient informa- tion was available.



(Continued)

any other aneuploidies.

Unclear: if this was not clear from the report.

Index	Were the index test results in- terpreted without knowledge	If a threshold was	Could the con- duct or inter-	Are there concerns
test ^b	of the results of the reference standard?	used, was it pre- specified?	pretation of the index test have intro- duced bias?	that the index test, its conduct, or in- terpretation dif- fer from the review question?
	 Yes: if the gNIPT results were interpreted without knowledge of the results of the reference standard^c. No: if the gNIPT results were interpreted with knowledge of the results of the reference standard^c. Unclear: if this was not clear from the report. 	Yes: if criteria for a positive test were prespecified. No: if the criteria for a positive test were not prespec- ified. Unclear: if this was not clear from the report.	Low risk: if 'yes' for all sig- nalling ques- tion. High or un- clear risk: if 'no' or 'unclear' was reported for at least one signalling ques- tion.	Low concern: if the gNIPT was per- formed such as de- scribed in the review question ^a . High concern: if gNIPT vary from those specified in the review question. Unclear concern: if insufficient informa- tion was available.
Domain 3: Re	ference standard			
Reference Standard ^c	Is the reference standard ^c likely to correctly classify the target condition ^d ?	Were the refer- ence standard re- sults interpreted without knowl- edge of the re- sults of the index test ^b ?	Could the ref- erence stan- dard, its con- duct, or its in- terpretation have intro- duced bias?	Are there concerns that the target con- dition as defined by the reference standard does not match the review question?
	Yes: if one appropriate refer- ence standard ^c was used. No: if pregnant women did not undergo appropriate reference	Yes: if karyotype results were in- terpreted without knowledge of re- sults of the index	Low risk: if 'yes' for all sig- nalling ques- tions. High or un-	Low concern: if the reference standard- s ^c were used as de- scribed in the review question ^a .

(Continued)

Domain 4: Flow and timing

Flow and timing	Was there an appropriate in- terval between gNIPT and ref- erence standard?	Did all analysed patients receive the reference standard?	Were all pa- tients in- cluded in the analysis?	Could the pa- tient flow have introduced bias?
	Yes: if the interval between blood collection for gNIPT and fluid collection for reference standard ^c was more than one day (only if blood collection oc- curred after fluid collection). If blood collection occurred be- fore fluid collection, there is no time limit ^e . No: if the interval between blood collection for gNIPT and the fluid collection for reference standard ^c was less than one day if the blood collection occurred after the fluid collection. Unclear: if this was not clear from the report.	Yes: if all pregnant women analysed have appropriate reference stan- dard ^c . No: if some preg- nant women analysed do not have a karyotype result. Unclear: if this was not clear from the report.	Yes: if all preg- nant women recruited in- to the study were included in the analy- sis or if failed samples oc- curred before NGS process. No: if all preg- nant women recruited in- to the study were not in- cluded in the analysis or if failed sam- ples occurred during NGS process. Unclear: if this was not clear from the report.	Low risk: if 'yes' for all sig- nalling ques- tions. High or un- clear risk: if 'no' or 'unclear' was reported for at least one signalling ques- tion.

^aReview question: what is the diagnostic accuracy of massively parallel shotgun sequencing (MPSS) and targeted massively parallel sequencing (TMPS) using circulating cell-free DNA (ccfDNA) in maternal blood for the detection of common fetal aneuploidies (T21, T18, T13, 45,X, 47,XXY, 47,XXX and 47,XYY) in pregnant women according to their prior risk of fetal aneuploidy?

^bIndex test refers to genomics-based non-invasive prenatal testing (gNIPT) methods such as MPSS or TMPS.

^cThe appropriate reference standard is karyotyping (traditional banding techniques or spectral karyotyping from invasive methods like chorionic villi sampling or amniocentesis), chromosome analysis (e.g. FISH, aCGH and QF-PCR), clinical examination or medical record from birth (for T21, T18 or T13). For sex chromosome aneuploidies, only fetal karyotype was appropriate reference standard because they usually have a normal phenotype.

^dTarget conditions (aneuploidies) are T21, T18, T13, 45,X, 47,XXY, 47,XXX and 47,XYY. ^eTarget conditions (aneuploidies) do not vary over time.

Appendix 7. gNIPT accuracy in mixed prior risk of fetal aneuploidy

Test	Number of	Number of	Number of unaffected	Sensitivity ^b	Specificity ^b	P value ^c
	studies	affected	pregnan- cies ^a	% (95% CI)	% (95% CI)	



(Continued)		pregnan- cies				
T21, mixed risi	k					
MPSS	10	445	30,962	96.0 (93.7 to 97.4)	99.9 (99.9 to 100)	
TMPS	6	169	6925	98.2 (94.6 to 99.4)	100 (99.9 to 100)	
Difference betw	ween MPSS	and TMPS		-2.27 (-4.97 to 0.43)	-0.04 (-0.08 to -0.002)	0.10
Tradition- al screening tests ^d	1	3	1909	100 (43.9 to 100)	96.4 (95.5 to 97.1)	
T18, mixed ris	k					
MPSS	9	113	30,637	100 (98.3 to 100) ^e	99.9 (99.8 to 100)	
TMPS	4	53	5569	98.1 (87.8 to 99.7)	99.9 (99.8 to 100)	
Tradition- al screening tests	1	1	1905	100 (20.7 to 100)	99.4 (99.0 to 99.7)	
T13, mixed ris	kf					
MPSS	8	27	30,384	100 (87.5 to 100) ^e	100 (> 99.9 to 100) ^e	
TMPS	5	31	8362	78.6 (48.3 to 93.5)	99.9 (99.8 to 100)	
45X, mixed ris	k					
MPSS	2	12	296	91.7 (58.7 to 98.8)	100 (98.7 to 100) ^e	
TMPS	2	22	1128	90.9 (70.0 to 97.7)	99.9 (99.4 to 100)	
47,XXY, mixed	risk					
MPSS	1	1	107	ND		
TMPS	1	2	184	ND		
47,XYY, mixed	risk					
MPSS	1	1	199	ND		
TMPS	1	1	185	ND		
Autosomes (T2	21, T18 and	T13 combined),	mixed risk			
MPSS	10	585	30,822	96.9 (95.2 to 98.1)	99.9 (99.9 to 99.9)	
TMPS	7	253	8793	96.0 (92.8 to 97.9)	99.8 (99.7 to 99.9)	
Difference bet	ween MPSS	and TMPS		0.88 (-1.90 to 3.65)	0.07 (-0.02 to 0.16)	0.25

Coch Libr	nrane ary	Trusted evidence. Informed decisions. Better health.			Cochrane Database of Systematic Reviews
(Continued) Tradition- al screening	1	4	1908	100 (51.0 to 100)	95.8 (94.8 to 96.6)
tests SCA (45,X, 47,	 XXX, 47,X	XY and 47,XYY combi	ned), mixed	risk	
MPSS	2	14	294	92.9 (63.0 to 99.0)	100 (98.7 to 100) ^e
TMPS	2	25	1125	92.0 (73.1 to 98.0)	99.9 (99.4 to 100)

45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, ND: no data available, T21: trisomy 21, T18: trisomy 18, T13: trisomy 13, CI: confidence interval, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, SCA: sex chromosome aneuploidies.

^aWe included pregnancies with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies.

^bFor two or more studies, the sensitivities and specificities are the summary estimates obtained from meta-analysis. Sensitivity and specificity, and their 95% CIs are reported as percentages.

^cThe P value indicates the statistical significance of the difference in model fit and was obtained from likelihood ratio tests comparing models with and without a covariate for test type.

^dTraditional screening test are first trimester combined test, second trimester quadruple test, second trimester fully integrated test, second trimester sequential test or second trimester triple test.

eSimple pooling used to obtain summary estimates of sensitivity and/or specificity.

^fTest comparison analysis did not converge.

Appendix 8. Investigation of heterogeneity

Test subgrou	ips	Number of studies	Number of affected pregnancies	Number of unaffected pregnancies ^a
Reference st	andard			
Autosomes, ı	unselected population			
MPSS	mixed reference standard ^b	1	11	1730
TMPS	karyotyping ^c	1	11	181
	mixed reference standard	3	107	20,468
Autosomes, s	elected high-risk population			
MPSS	karyotyping	22	1075	7028
	mixed reference standard	10	433	8769
TMPS	karyotyping	7	378	4282

(Continued)

SCA, selected high-risk population

JCA, selected				
MPSS	karyotyping	10	134	3943
	mixed reference standard	2	17	3509
TMPS	karyotyping	4	96	968
Ethnicity				
Autosomes, i	unselected population			
MPSS	more than 50% Asian ^d	1	11	1730
TMPS	more than 50% Caucasian ^e	3	107	20,468
	not reported	1	11	181
Autosomes, s	selected high-risk population			
MPSS	more than 50% Asian	14	206	6589
	more than 50% Caucasian	7	843	6262
	not reported	11	459	2946
TMPS	more than 50% Caucasian	3	237	3744
	not reported	4	141	538
SCA, selected	l high-risk population			
MPSS	more than 50% Asian	5	25	1852
	more than 50% Caucasian	5	96	4286
	not reported	2	30	1314
TMPS	more than 50% Caucasian	1	56	116
	not reported	3	40	852
	notreportea	3	40	852

45,X: Turner syndrome, 47,XXX: triple X syndrome, 47,XXY: Klinefelter syndrome, T21: trisomy 21, T18: trisomy 18, T13: trisomy 13, MPSS: massively parallel shotgun sequencing, TMPS: targeted massively parallel sequencing, SCA: sex chromosome aneuploidies

^aWe included pregnancies with any other aneuploidy than the one under analysis with all euploid cases as "unaffected" pregnancies.

^bMixed RS: include karyotyping and neonatal clinical examination or medical records from birth.

^cKaryotyping: include fetal karyotyping performed on cells obtained from chorionic villi sampling (CVS), amniotic fluid, placental tissue, a fetus lost by miscarriage or other equivalent and recognised methods on the same materials.

^dMore than 50% Asian: in the cohort, more than 50% of all pregnant women were Asian ethnicity.

eMore than 50% Caucasian: in the cohort, more than 50% of all pregnant women were Caucasian ethnicity.

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HISTORY

Protocol first published: Issue 7, 2015 Review first published: Issue 11, 2017

Date	Event	Description
2 July 2015	Amended	Minor changes to author affiliations.

CONTRIBUTIONS OF AUTHORS

FR initiated this review project and recruited all authors. MB co-ordinated the production of the review.

MB, CL, JB, YT and FR drafted the protocol and all authors critically commented on the protocol. The final protocol was read and approved by all authors.

MB, CL and WW developed and applied the literature search. MB, CL, JB and LN applied eligibility criteria, collected, filled the data form and assessed methodological quality of the included studies. MB contacted authors of the included studies. YT checked data collection, undertook statistical analysis and data synthesis. MB, CL and FR wrote the draft of the review. MB, CL, JB, LN, YT, SL, YG and FR contributed with redrafting. SL, FL, YG and FR provide senior clinical input. FR oversaw the review process. The final review was read and approved by all authors.

DECLARATIONS OF INTEREST

Mylene Badeau: none known.

Jonatan Blais: none known.

Yves Giguère (and some other members of the review team - FR, SL, FL) are investigators in a Research Project funded under the auspices of Genome Canada and the Canadian Institutes for Health Research (both non-for-profit organisations funded by the Canadian government) but that call for some mandatory in-kind contributions from other partners. This Research Project thus receives in-kind funding from private corporations which either offer commercial NIPT tests (Ariosa Diagnostics Inc, San Jose, CA) or offer reagents and/or equipment that can be used to perform NIPT assays (Life Technologies Inc, NY, USA; Illumina, San Diego, CA, USA; QIAGEN, Hilden, GER; Perkin Elmer, Waltham, MASS, USA). This funding is at arms length from the scientific components of the research project. The present review is not funded by this Project. Some of the authors may eventually publish results from clinical trials that could be considered in this review. None of the four authors involved in gNIPT studies will take part in the selection of studies, nor in any decisions/analyses related to their own studies. All authors declared no other conflict of interest

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DIFFERENCES BETWEEN PROTOCOL AND REVIEW

In the "reference standards" section in the methods, the protocol did not specify that neonatal clinical examination or medical records from birth are not appropriate reference standards for SCA diagnosis. These aneuploidies (SCA) usually have a normal phenotype and can not be detected with neonatal clinical examination or medical records from birth.

In the "sensitivity analyses" section, of the protocol, the authors stated they intended to investigate "studies where not all pregnant women received neither reference standard (no karyotyping confirmation nor birth follow-up); authors who have taken for granted that the baby is normal". We decided to remove this analysis because this goes against one of our criteria for considering studies for this review (pregnant women with MPSS or TMPS and a reference standard). These type of studies were excluded.

We decide to remove Google scholar from our electronic searches databases list. Google scholar found more than 100,000 publications about our topic but only the first 1000 are retrievable. This database is not reproducible and search fields did not allow us to specify the search strategy. All first most relevant publications were already found with other databases.

We changed OpenSIGLE to Opengrey because the first has been replaced by the second.

Although studies used different cutpoints, there was little or no variation in threshold and no requirement to estimate the correlation between sensitivity and specificity across studies in a meta-analysis. Therefore, we did not estimate summary ROC curves using the HSROC model. As the cutpoints were regarded as qualitative, we estimated summary sensitivities and specificities using random-effects and



fixed-effect logistic regression models, and simple pooling as appropriate. Further details are available in the statistical analysis and data synthesis section. We used the Stata software package for the analyses instead of SAS.

We made changes in the QUADAS-2 tool. In domain 1, at third signalling question, we added "maternal cancer history, type of pregnancy, gestational age, assisted reproductive technology" in "No" answer. In concerns of applicability in domain 3, concerns about applicability, we modified conditions for low and high concern according to the review question. In domain 4, at second signalling question, we added "analysed" for clarify and we changed "yes" answer by removing "karyotype result" for "appropriate reference standard".

INDEX TERMS

Medical Subject Headings (MeSH)

*Aneuploidy; Cell-Free Nucleic Acids [*blood]; Chromosome Disorders [*diagnosis] [genetics]; Disorders of Sex Development [diagnosis] [genetics]; Fetal Diseases [*diagnosis] [genetics]; High-Throughput Nucleotide Sequencing [*methods]; Pregnancy, High-Risk; Prenatal Diagnosis [*methods]

MeSH check words

Female; Humans; Pregnancy