Diagnostic yield of exome sequencing for prenatal diagnosis of fetal structural anomalies: A systematic review and meta-analysis

Rhiannon Mellis¹,² | Kathryn Oprych³ | Elizabeth Scotchman¹ | Melissa Hill¹,² | Lyn S Chitty¹,²

Objective: We conducted a systematic review and meta-analysis to determine the diagnostic yield of exome sequencing (ES) for prenatal diagnosis of fetal structural anomalies, where karyotype/chromosomal microarray (CMA) is normal.

Methods: Following electronic searches of four databases, we included studies with ≥10 structurally abnormal fetuses undergoing ES or whole genome sequencing. The incremental diagnostic yield of ES over CMA/karyotype was calculated and pooled in a meta-analysis. Sub-group analyses investigated effects of case selection and fetal phenotype on diagnostic yield.

Results: We identified 72 reports from 66 studies, representing 4350 fetuses. The pooled incremental yield of ES was 31% (95% confidence interval (CI) 26%–36%, \( p < 0.0001 \)). Diagnostic yield was significantly higher for cases pre-selected for likelihood of monogenic aetiology compared to unselected cases (42% vs. 15%, \( p < 0.0001 \)). Diagnostic yield differed significantly between phenotypic sub-groups, ranging from 53% (95% CI 42%–63%, \( p < 0.0001 \)) for isolated skeletal abnormalities, to 2% (95% CI 0%–5%, \( p = 0.04 \)) for isolated increased nuchal translucency.

Conclusion: Prenatal ES provides a diagnosis in an additional 31% of structurally abnormal fetuses when CMA/karyotype is non-diagnostic. The expected diagnostic yield depends on the body system(s) affected and can be optimised by pre-selection of cases following multi-disciplinary review to determine that a monogenic cause is likely.

Key points

What’s already known about this topic?

- Prenatal exome sequencing (ES) increases genetic diagnoses in fetuses with structural abnormalities and a normal karyotype and chromosomal microarray.
- Published diagnostic yields from ES are varied and may be influenced by study size, case selection and fetal phenotype.
1 | INTRODUCTION

Ultrasound scanning in pregnancy can reveal fetal structural abnormalities from the first trimester onwards, ranging from isolated minor anomalies to severe, multisystem disorders. These abnormalities may indicate an underlying genetic condition but making a definitive diagnosis can be difficult because many conditions have overlapping fetal presentations. A genetic diagnosis can clarify the prognosis for the fetus, informing parental decision making and clinical management during pregnancy and the perinatal period.

Conventional genetic testing for fetal anomalies comprises quantitative fluorescence PCR (QF-PCR) for rapid aneuploidy exclusion, and karyotype or chromosomal microarray (CMA) for detection of chromosomal and sub-chromosomal abnormalities. Together, these can provide a diagnosis in up to 40% of cases, but leave the majority undiagnosed. Targeted sequencing of individual genes can be offered in cases where clinical features or family history suggest a specific monogenic disorder but genome-wide next generation sequencing (NGS) is now being used more frequently to allow interrogation of multiple genes in a single test. Exome sequencing (ES) examines the 1%-2% of the genome that encodes proteins and harbours around 85% of known disease-causing genetic variants, and is generally less costly and easier to interpret than whole genome sequencing (WGS). ES can be applied in practice as whole ES (sequencing or analysing all exons), medical/clinical ES (sequencing or analysing only the exons of known Mendelian disease genes), or targeted ES (applying ‘virtual’ gene panels to ES data to limit analysis to genes of interest for a specific phenotype).

In the postnatal setting ES has proven a powerful tool for the diagnosis of children with neurodevelopmental delay but its application in prenatal diagnosis initially lagged behind. This is largely because long turnaround times precluded the return of results within the timeframe of a pregnancy. Additionally, incomplete or unrecognised fetal phenotypes complicate the interpretation of genetic variants from prenatal ES. Over recent years, rapid analytical pipelines have made it possible to return ES results during an ongoing pregnancy and it is now considered for prenatal diagnosis of fetal structural anomalies in the guidance of several professional bodies. Two large prospective studies demonstrated the diagnostic utility of ES for prenatal diagnosis of unselected fetal structural abnormalities, reporting diagnostic rates of 8.5%–10%, while numerous smaller studies report diagnostic rates above 80% in highly-selected case series. This body of literature has been previously reviewed but has rapidly expanded over the last 2 years.

As prenatal ES is increasingly adopted in clinical settings there remain important questions around best practice for implementation. It will be of interest to health professionals and policy makers to define expected diagnostic yields from prenatal ES for different fetal presentations. This information may help to manage parental expectations during pre-test counselling and will inform development of further guidelines on when and how to offer prenatal ES. In this systematic review and meta-analysis we aim to address the question ‘what is the diagnostic yield of ES for prenatal diagnosis in pregnancies complicated by fetal structural anomalies where fetal karyotype/microarray is normal?’ We also explore the diagnostic yield in different phenotypic sub-groups.

2 | METHODS

2.1 | Protocol and registration

We devised a systematic review protocol in line with PRISMA guidance and registered it prospectively on the PROSPERO international register of systematic reviews (reference CRD42020200600). Study authors agreed the protocol prior to conducting the searches. Any required small amendments were made with the consensus of all authors.

2.2 | Eligibility criteria

Studies were included in this review if they met the following criteria: (i) Retrospective or prospective cohorts of 10 or more pregnancies undergoing ES (whole, clinical or targeted) or WGS for diagnosis of fetal structural anomalies, (ii) CMA/karyotype was negative or non-diagnostic, (iii) Testing was initiated based on the prenatal phenotype, (iv) Full text report was available in English language.

The focus of this review is prenatal diagnosis of structural anomalies, therefore studies with mixed pre- and postnatal presentations were only included if it was possible to extract the prenatal cases. Similarly, for studies with mixed indications for testing (e.g. some cases initiated because of fetal phenotype and some because of family history) we only included studies where fetal anomaly cases could be extracted. Studies where ES was completed postnatally for some or all cases but had been initiated based on the prenatal phenotype alone were included. Only variants classified as pathogenic/likely pathogenic according to diagnostic-standard guidelines and determined to be causative of the fetal phenotype.
were considered in diagnostic yield, therefore studies using ES purely for novel candidate gene discovery were not included. We also excluded case reports, series of fewer than 10 cases, reviews, editorials and commentaries, and studies using gene panel testing rather than ES.

2.3 Information sources and search strategy

Electronic searches of four databases (MEDLINE, Embase, Cochrane library, and Web of Science) were conducted for records published between January 2010 and October 2021. This date range was chosen because prenatal ES was not used prior to 2010. The search was initially conducted on 27/08/2020 and repeated on 15/10/2021. Additional potentially relevant studies were identified by manually searching reference lists of relevant studies and published reviews, and conference proceedings of prenatal and genetics conferences of relevant major societies in last 3 years.

Search terms were variations on the keywords ‘prenatal diagnosis’ and ‘exome sequencing’. Alternative terms for ‘prenatal diagnosis’ included ‘fetal’, ‘fetus’, ‘prenatal’, and ‘antenatal’. Alternative terms for ‘exome sequencing’ included ‘whole exome sequencing’, ‘exome’, ‘whole genome sequencing’, ‘genome, human’ and ‘sequence analysis, DNA’. The full search strategy is detailed in supporting information S1. We chose relatively broad search terms in an effort to capture as many relevant studies as possible; for the same reason no search limits were applied apart from date range.

2.4 Study selection

After the removal of duplicates, two reviewers (RM and KO) independently screened titles and abstracts. For abstracts identified as potentially relevant, full text articles were retrieved and reviewed against the inclusion and exclusion criteria. Any disagreement between reviewers was resolved by discussion. Where the same data were presented more than once, the most recent study was selected.

2.5 Data extraction

The following data, where available, were extracted by two reviewers (RM and ES) into a dataset: study setting, sample size, study inclusion criteria, ES approach, phenotypes used for interpretation (prenatal only or prenatal plus postnatal/post-mortem), number of fetuses with diagnostic variants, variants of uncertain significance (VUS), incidental findings, gestation at testing, test turnaround time, pregnancy outcomes, and impact on management. For studies that performed CMA in parallel with sequencing, the cases with negative CMA were extracted, in order to be comparable with other studies where chromosomal abnormalities were ruled out prior to ES.

2.6 Quality assessment of included studies

The quality of the studies was assessed using a modified Standards for Reporting of Diagnostic Accuracy (STARD) checklist, comprising 22 items. The items for reporting considered most important for these studies (as they may influence the reported diagnostic yield) were: (i) clearly described selection criteria, (ii) whether diagnostic variants were confirmed by Sanger sequencing, (iii) whether American College of Medical Genetics and Genomics (ACMG) or equivalent standardised classification was used for variant interpretation, (iv) approach to analysing and reporting VUS and incidental findings addressed. For eight studies the assessment was done independently by two reviewers (RM and MH) with discrepancies resolved by discussion. Once consensus was reached on how to apply the criteria and reviewers’ scores were concordant on new study assessments, the remaining studies were assessed by RM only.

2.7 Data synthesis

The primary outcome of interest was the incremental diagnostic yield of ES (i.e. the proportion of cases in which a diagnostic genetic variant is detected by ES) over CMA/karyotype, expressed as a risk difference. This was calculated for each study with 95% confidence intervals (CI) and pooled for all studies in a meta-analysis, using a random effects model with inverse variance weighting. A subgroup analysis was performed to investigate the effect on diagnostic yield of pre-test case selection for higher likelihood of monogenic disease, where this information was available. Where studies reported data on isolated system anomalies classified by different phenotypic groups, a further subgroup analysis was performed to estimate pooled incremental yields by phenotypic group. Only phenotype groups reported in at least three studies were included in this analysis. Findings were displayed using forest plots. Between-study heterogeneity was assessed visually and using an I² statistic with a Q-test. Publication bias was assessed graphically using a funnel plot with Egger’s test. All statistical analyses were performed in R studio v1.3.1093 using the ‘metafor’ package.

3 RESULTS

Database searches identified 8123 unique records in total. After screening by title and abstract, and retrieving full text reports, 148 articles were assessed, of which 72 reports from 66 studies were deemed eligible and included in the final review (Figure 1, Table 1). Six reports examined specific phenotypic sub-groups from previously published larger cohorts, hence were not separate studies. These were excluded from the overall meta-analysis of incremental yield and the subgroup analysis by case selection, to avoid double-counting of cases. However, these reports provide updated information in their respective phenotypic groups so were included in the subgroup
analysis by phenotypic group, in place of the previously reported subgroups from their parent studies. Across 66 studies, this review represents 4350 probands undergoing prenatal ES.

3.1 Study characteristics

The characteristics of the included studies are summarised in Table 1. Of the 72 included reports, 48 (67%) were published in 2020-21. Three studies used WGS rather than ES, with or without applying phenotype-specific panels for analysis.\(^{29-31}\) Eighteen studies used a clinical exome sequencing (CES) approach in some or all cases, which here we define as targeting either the sequencing or the analysis to a specific gene panel.\(^{8,13,62-71,34,41-48,33,40}\) Eight studies targeted ES analysis more tightly using phenotype-specific gene panels,\(^{7,12,15,49-53}\) while the remainder took a whole ES approach.\(^{8,13,62-71,54,72-81,55,82-88,56-61}\) Where stated (15 studies),\(^{7,8,74,75,78,82,84,15,29,36,52,53,55,62,73}\) the median turnaround time for ES was 20 days (range 4–141). Studies included fetuses with a range of structural abnormality phenotypes, with some studies recruiting cases with fetal anomalies in a specific body system (e.g. renal anomalies or cardiac defects) and others including any structural anomaly. In total, 42/66 studies recruited cases with anomalies in a specific body system or broke down fetal abnormalities within the cohort into distinct phenotypic groups (Table 1). The quality of included reports, assessed using modified STARD criteria, was generally high (Figure 2).

3.2 Incremental yield of diagnostic variants from ES

The diagnostic yield of ES for fetal structural anomalies among the included studies ranged from 5% (95% CI 0.8–9.8%) to 89% (95% CI 72%–99%). The pooled incremental yield of ES over CMA/karyotype from all studies was 31% (95% CI 26%–36%, p < 0.0001), illustrated in the forest plot (Figure 3). Between-study heterogeneity was high, indicated by an I\(^{2}\) statistic of 94% (Q = 682, p < 0.0001). Testing for publication bias showed significant funnel plot asymmetry (z = 5.6, p < 0.0001 by Egger’s test; supporting information S1), indicating possible publication bias.

3.3 Effect of case selection on diagnostic yield

Some of the heterogeneity between included studies may be due to differing approaches to case selection. Some studies included fetuses with structural anomalies without any prior review of the clinical features to ascertain risk of a monogenic aetiology, whereas others selected only pregnancies where the pattern of fetal anomalies was
| Study                  | No. of cases with normal CMA | Inclusion criteria                                                                 | Sequencing and analysis approach | Diagnostic variants (%) | VUS or candidate genes (%) | Incidental findings (%) | Clinical impact (%) | Average TAT in days (range, if given) |
|-----------------------|------------------------------|-----------------------------------------------------------------------------------|----------------------------------|------------------------|--------------------------|------------------------|---------------------|---------------------------------------|
| Carss et al, 2014a    | 28                           | Fetuses and neonates with prenatal structural anomalies                            | Trio WES                         | 3/28 (11%)             | 5/28 (18%)              | ND                     | ND                  | ND                     |
| Drury et al, 2015a    | 24                           | Fetuses with structural abnormalities, and/or increased NT (>3.5 mm)               | Solo or Trio CES                 | 5/24 (21%)             | 1/24 (4%)               | 2/24 (8%)              | ND                  | ND                     |
| Pangalos et al, 2016  | 14                           | Fetuses with (mild or severe) structural anomalies                                 | Solo TES (758 prenatal disorders gene panel) | 6/14 (43%)             | 1/14 (7%)               | 0/14 (0%)              | ND                  | <10                    |
| Yates et al, 2017     | 84                           | Fetuses with US anomalies resulting in TOP or IUFD                                | Solo or Trio WES                 | 17/84 (20%)            | 45/84 (54%)             | 2/84 (2%)              | ND                  | ND                     |
| Aarabi et al, 2018a   | 20                           | ≥1 major structural defect detected prenatally                                     | Trio WES or CES                  | 1/20 (5%)              | 2/20 (10%)              | ND                     | ND                  | ND                     |
| Armes et al, 2018a    | 16                           | Fetal or early neonatal death, having undergone full autopsy, genetic aetiology suspected (PM) | Trio WGS with panel analysis     | 2/16 (13%)             | 6/16 (38%)              | ND                     | ND                  | ND                     |
| Boissel et al, 2018a,b| 101                          | Fetuses or stillborns with severe anomalies:                                      | Solo or Trio WES                 | 19/101 (19%)           | 7/101 (7%)              | ND                     | ND                  | ND                     |
| Chandler et al, 2018a,b| 16                          | Fetuses with suspected skeletal dysplasia on prenatal US                          | Trio TES (240 skeletal gene panel) | 13/16 (81%)            | 2/16 (13%)              | ND                     | NQ                  | 16.5 (11–41)            |
| Fu et al, 2018a,b     | 196                          | Fetuses with ultrasound detected structural malformations, excluding isolated increased NT ≥ 3.5 mm/cystic hygroma, and isolated soft markers | Solo or Trio CES                 | 47/196 (24%)           | 25/196 (13%)            | 12/196 (6%)            | ND                  | ~21                    |
| Leung et al, 2018a    | 33                           | Fetuses with structural anomalies detected on prenatal US                          | Trio WES                         | 3/33 (9%)              | 6/33 (18%)              | ND                     | ND                  | ND                     |
| Shamseldin et al, 2018a | 44                        | Severe fetal malformations, lethal NIHF or unexplained IUFD                       | Solo WES                         | 22/44 (50%)            | 15/44 (34%)             | ND                     | ND                  | ND                     |
| Zhou, X et al, 2018a,b| 12                          | Fetuses with suspected skeletal dysplasia on US                                  | Solo TES (363 skeletal dysplasia gene panel) | 10/12 (83%)            | ND                      | ND                     | ND                  | 42                     |
| Choy et al, 2019a,b   | 42                           | Fetuses with increased NT ≥ 3.5 mm /− structural anomalies                         | Solo WGS                         | 7/42 (17%)             | 18/42 (43%)             | ND                     | None - results not returned to families | 10                    |
| Daum et al, 2019a,b   | 77                           | Fetuses with US-detected                                                          | Solo or Trio WES                 | 16/77 (21%)            | ND                      | ND                     | NQ                  | 26 (19–33)             |
| Study                  | No. of cases with normal CMA | Inclusion criteria                                                                 | Sequencing and analysis approach | Diagnostic variants (%) | VUS or candidate genes (%) | Incidental findings (%) | Clinical impact (%) | Average TAT in days (range, if given) |
|-----------------------|------------------------------|------------------------------------------------------------------------------------|----------------------------------|--------------------------|---------------------------|------------------------|---------------------|---------------------------------------|
| De Koning et al, 2019 | 20                           | Fetuses with one or more structural anomalies, genetic etiology suspected          | Trio WES                         | 8/20 (40%)              | 1/20 (5%)                 | 3/20 (15%)              | 9/12 (75%)           | 10.5 (7–19)          |
| Greenbaum et al, 2019 | 45                           | Fetuses with abnormal US findings and/or relevant family history                   | Solo or Trio WES                 | 13/45 (29%)             | ND                        | ND                     | ND                  | Ongoing pregnancies: 15–20 Post-mortem: 15–60 |
| Liu et al, 2019ab     | 28                           | Fetuses with skeletal abnormalities on US                                           | Solo TES (363 skeletal dysplasia gene panel) | 16/28 (57%)             | 5/28 (18%)                | ND                     | ND                  | ND                     |
| Lord et al, 2019ab    | 610                          | Fetuses with any structural anomaly on US, including isolated increased NT ≥ 4 mm | Trio TES (1628 developmental disorder gene panel) | 52/610 (9%)             | 24/610 (4%)               | ND                     | ND                  | ND                     |
| Meier et al, 2019a    | 19                           | Severe fetal anomalies identified by US scan: (i) Two or more anomalies associated with a high risk for fetal or perinatal lethality and suspected genetic etiology, or (ii) familial recurrence of the fetal anomaly phenotype | Trio WES                         | 6/19 (32%)             | 6/19 (32%)                | ND                     | ND                  | ND                     |
| Normand et al, 2019a  | 146                          | Fetuses with at least one structural anomaly detected by prenatal imaging or autopsy | Solo or Trio WES                 | 46/146 (32%)            | ND                        | 3/62 (5%)              | 15/19 (79%)          | Rapid: 14 (7–38) Trio: 43 (13–78) Solo: 88 (18–141) |
| Petrovski et al, 2019b| 234                          | Any ultrasound identified fetal structural abnormalities, including isolated NT ≥ 3-5 mm | Trio WES                         | 24/234 (10%)           | 46/234 (20%)              | 4/234 (2%)             | ND                  | 28–56                  |
| Quinlan-Jones et al, 2019a | 27                      | Fetuses with a significant structural anomaly resulting in TOP, IUFD, or NND | Trio TES (1628 developmental disorder gene panel) | 10/27 (37%)             | 6/27 (22%)                | ND                     | ND                  | ND                     |
| Westphal et al, 2019b | 30                           | Fetuses and neonates with prenatally diagnosed CHDs (focus on severe and syndromic) | Trio WES                         | 6/30 (20%)              | 4/30 (13%)                | 3/30 (10%)             | ND                  | ND                     |
| Study                        | No. of cases with normal CMA | Inclusion criteria                                                                                                                                                                                                 | Sequencing and analysis approach                                                                 | Diagnostic variants (%) | VUS or candidate genes (%) | Incidental findings (%) | Clinical impact (%) | Average TAT in days (range, if given) |
|-----------------------------|------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------|--------------------------|---------------------------|--------------------------|----------------------|---------------------------------------|
| Aggarwal et al, 2020        | 32                           | Fetuses with structural abnormalities or unexplained IUFD, genetic aetiology suspected (PM)                                                                                                                   | Solo and Trio, WES or CES                                                                                                                              | 18/32 (56%)              | 4/32 (13%)                | ND                       | ND                   | ND                     |
| Becher et al, 2020          | 35                           | One or more fetal malformations or severe fetal hydrops, genetic aetiology suspected                                                                                                                        | Trio WES                                                                                                                                                 | 9/35 (26%)               | 7/35 (20%)                | 1/35 (3%)                | NQ                   | Urgent: 12 (9–16) Non-urgent: 120–180 |
| Chen et al, 2020²            | 105                          | Fetuses with structural anomalies, including isolated increased NT > 3.5mm                                                                                                                                     | Trio CES                                                                                                                                                 | 20/105 (19%)             | 12/105 (11%)              | ND                       | ND                   | 25                     |
| Corsten-Janssen et al, 2020 | 23                           | Fetuses with major structural anomalies                                                                                                                                                                       | Trio TES (~ 3850 OMIM disease gene panel with late onset diseases excluded)                                                                            | 8/23 (35%)               | ND                        | 11/23 (48%)              | 6/23 (26%)            | 14 (8–20)               |
| Deden et al, 2020²           | 54                           | Fetuses with multiple congenital anomalies. Isolated major anomaly or multiple soft markers only if high suspicion of genetic aetiology                                                                                                                     | Trio WES                                                                                                                                                 | 18/54 (33%)              | 2/54 (4%)                 | ND                       | 24/37 (68%)           | 10 (4–28)               |
| Deng et al, 2020²³           | 21                           | Fetuses with NIHF +/- other structural anomalies                                                                                                                                                               | Trio CES                                                                                                                                                 | 3/21 (14%)               | 2/21 (10%)                | ND                       | NQ                   | ND                     |
| Guo et al, 2020             | 40                           | Recurrent fetal congenital abnormalities or dysmorphic features                                                                                                                                               | Trio WES                                                                                                                                                 | 12/40 (30%)              | 20/40 (50%)               | ND                       | NQ                   | ND                     |
| Han et al, 2020²³           | 26                           | Fetal anomalies suggestive of a skeletal dysplasia                                                                                                                                                              | Trio CES                                                                                                                                                 | 23/26 (89%)              | 1/26 (4%)                 | 1/26 (4%)                | NQ                   | <14                    |
| Heide et al, 2020²³         | 62                           | Fetuses with abnormal corpus callosum (complete or partial agenesis, or short CC) detected on US +/- other anomalies                                                                                           | Trio TES (425 brain anomaly gene panel) Or 12/62 (19%)                                                                                                 | 6/65 (9%)                | ND                        | ND                       | NQ                   | 21.5 (9–53)            |
| Lefebvre et al, 2020        | 95                           | Fetuses with multiple congenital anomalies                                                                                                                                                                      | Solo WES                                                                                                                                                 | 24/95 (25%)              | 14/95 (15%)               | ND                       | ND                   | ND                     |
| Lei et al, 2020²³           | 163                          | Fetuses with CAKUT +/- other structural anomalies                                                                                                                                                              | Trio CES                                                                                                                                                 | 20/163 (12%)             | 2/163 (1%)                | 9/163 (6%)               | NQ                   | 14–84                  |
| Study                  | No. of cases with normal CMA | Inclusion criteria                                                                 | Sequencing and analysis approach | Diagnostic variants (%) | VUS or candidate genes (%) | Incidental findings (%) | Clinical impact (%) | Average TAT in days (range, if given) |
|-----------------------|-----------------------------|-----------------------------------------------------------------------------------|---------------------------------|--------------------------|----------------------------|------------------------|---------------------|---------------------------------------|
| Li, L et al, 2020ab   | 19                          | Fetuses with cerebellar vermis defects +/- other structural anomalies              | Solo or Trio CES                | 8/19 (42%)               | 2/19 (11%)               | 1/19 (5%)                | ND                  | 28–56                                 |
| Li, R et al, 2020ab   | 260                         | Fetuses with congenital heart defects +/- other structural anomalies               | Trio CES                        | 26/260 (10%)             | 16/260 (6%)              | 7/260 (3%)               | ND                  | 21–56                                 |
| Mone et al, 2020bc    | 197                         | Fetuses with prenatally detected congenital heart disease (excluding small muscular VSDs), extracted from the PAGE study extended cohort (Lord et al, 2019) |
|                       |                             | Trio TES (1628 developmental disorder gene panel)                                |                                 |                          |                           |                        | ND                  | ND                                    |
| Pooh et al, 2020a     | 16                          | Fetuses with anomalies detected on US at 11-13 weeks, genetic disease strongly suspected | Trio CES                        | 7/16 (44%)               | ND                        | ND                     | ND                  | 7–10                                 |
| Reischer et al, 2020ab| 11                          | Fetuses with decreased/absent fetal movement plus arthrogryposis of limbs       | Solo WES                        | 6/11 (55%)               | 1/11 (9%)                | ND                     | ND                  | ND                                    |
| Sparks et al, 2020ab  | 127                         | Fetuses with NIHF, fetal ascites, pleural or pericardial effusions, skin oedema, cystic hygroma, increased NT, or a combination of these conditions | Trio WES                        | 37/127 (29%)             | 12/127 (9%)              | 4/115                  | ND                  | Ongoing pregnancies: 14–28 Post-mortem: 56–84 |
| Sun et al, 2020ab     | 66                          | Fetuses with cardiac left-sided lesions                                           | Solo or Trio WES                | 13/66 (20%)              | 11/66 (17%)              | 1/66 (2%)               | ND                  | ND                                    |
| Vora et al, 2020a     | 102                         | Fetuses with isolated or multiple congenital anomalies                           | Trio WES (n = 99) or WGS (n = 3) | 21/102 (21%)             | 10/102 (10%)             | 9/204 (4%) in consenting parents | ND                  | 182–365                               |
| Xue et al, 2020ab     | 24                          | Fetuses with increased NT and no detectable structural malformations on US       | Trio WES                        | 3/24 (13%)               | 2/24 (8%)                | ND                     | ND                  | ND                                    |
| Yang et al, 2020ab    | 73                          | Fetuses with isolated first-trimester increased NT ≥35 mm                         | Trio CES                        | 4/73 (6%)                | 7/73 (10%)               | ND                     | ND                  | ND                                    |
| Study                        | No. of cases with normal CMA | Inclusion criteria                                                                 | Sequencing and analysis approach | Diagnostic variants (%) | VUS or candidate genes (%) | Incidental findings (%) | Clinical impact (%) | Average TAT in days (range, if given) |
|-----------------------------|-----------------------------|------------------------------------------------------------------------------------|----------------------------------|--------------------------|---------------------------|------------------------|---------------------|--------------------------------------|
| Zhou, X et al, 2020<sup>a,b</sup> | 41                          | Fetuses with isolated renal abnormalities on US                                    | Solo or Trio WES                 | 3/41 (7%)                | ND                        | 1/41 (2%)              | NQ                  | ND                                  |
| Qi et al, 2020<sup>a,b</sup> | 80                          | Fetuses with at least one ultrasonographic structural anomaly, including isolated increased NT | Trio CES (5000 OMIM genes) +/- WES (where sufficient DNA) | 27/80 (33.8%) | 5/80 (6.3%) | ND | ND | 14–28 |
| Rinaldi et al, 2020<sup>a</sup> | 30                          | Fetuses with structural malformation or ‘severe condition’ on US (e.g. growth restriction, absence of spontaneous movements) | Trio CES (5817 OMIM genes)       | 19/30 (63%)              | 0/30                     | ND                     | None in affected pregnancy - sequenced after end of pregnancy | ND |
| Al-Kouatly et al, 2021<sup>a</sup> | 22                          | Fetuses with confirmed NIHF after first trimester +/- other structural anomalies (strict phenotypic description for NIHF: at least two abnormal fluid collections) | Trio CES                          | 11/22 (50%)              | 5/22 (22.7%)            | ND                     | NQ                  | 14–21 |
| Aoi et al, 2021<sup>a</sup> | 17                          | Fetuses with ultrasonographic structural anomalies                               | Solo or Trio WES                 | 5/17 (29.4%)             | 1/17 (5.9%)             | ND                     | ND                  | ND                                  |
| Cao et al, 2021<sup>a,b</sup> | 11                          | Fetuses with abnormalities suggestive of fetal akinesia                           | Trio CES (4200 OMIM genes)       | 5/11 (45.5%)             | 3/11 (27.3%)            | ND                     | None in affected pregnancy - sequenced after end of pregnancy | ND |
| Correa et al, 2021<sup>a,b</sup> | 19                          | Fetuses with confirmed NIHF after first trimester +/- other structural anomalies (strict phenotypic description for NIHF: at least two abnormal fluid collections) | Solo WES                          | 7/19 (36.8%)             | 1/19 (5.3%)             | ND                     | ND                  | ND                                  |
| De Koning et al, 2021<sup>a,c</sup> | 18                          | Fetuses with CNS malformations detected by prenatal US; isolated or with other structural anomalies. (Some cases previously published in De Koning et al, 2019) | Trio WES                          | 10/18 (56%)              | 2/18 (11.1%)            | 2/18 (11.1%) | 12/18 (67%) | ND |
| Dempsey et al, 2021<sup>a</sup> | 52                          | Ongoing pregnancies with US-detected fetal structural                             | Trio WES with phenotype-specific panel analysis | 19/52 (37%) | 13/52 (25%) | ND | 17/43 (39.5%) in pregnancies | 17 |
| Study                        | No. of cases with normal CMA | Inclusion criteria                                                                 | Sequencing and analysis approach       | Diagnostic variants (%) | VUS or candidate genes (%) | Incidental findings (%) | Clinical impact (%) | Average TAT in days (range, if given) |
|-----------------------------|-----------------------------|-----------------------------------------------------------------------------------|----------------------------------------|--------------------------|---------------------------|------------------------|-------------------|--------------------------------------|
| He et al, 2021              | 94                          | Fetuses with structural anomalies, detected by prenatal US in the second trimester | Solo or Trio WES                      | 37/94 (39.4%)            | 14/94 (14.9%)             | 4/94 (4.3%)             | ND                | 42–56                                |
| Kucińska-Chahwan et al, 2021 | 26                          | Fetuses with skeletal abnormalities, excluding isolated talipes                     | Solo WES                              | 18/26 (69.2%)           | ND                        | ND                     | ND                | 84                                   |
| Lei et al, 2021             | 85                          | Fetuses with US-detected structural anomalies, other than skeletal abnormalities   | Solo WES                              | 16/85 (18.8%)           | 16/85 (18.8%)             | ND                     | ND                | 14–21                                |
| Liao et al, 2021            | 24                          | Fetuses with ultrasonographic diagnosis of abnormal sylvian fissure                | Solo, Duo or Trio WGS                | 12/24 (50%)             | ND                        | ND                     | ND                | ND                                   |
| Melli et al, 2021           | 213                         | Fetuses with increased NT ≥35 mm at 11–14 weeks of gestation +/- other anomalies, extracted from the PAGE and Columbia studies’ extended cohorts (Lord et al, 2019; Petrovski et al, 2019) | Trio TES (1628 developmental disorder gene panel) or WES | 28/213 (13.1%) | 8/213 (3.8%) | ND | ND | ND |
| Mone et al, 2021            | 28                          | Fetuses with NIHF +/- other anomalies, extracted from the PAGE study extended cohort (Lord et al, 2019) | Trio TES (1628 developmental disorder gene panel) | 7/28 (25%) | 2/28 (7.1%) | ND | ND | ND |
| Peng et al, 2021            | 38                          | Fetuses with suspected skeletal dysplasia on prenatal US                           | Solo or Trio WES                      | 25/38 (65.8%)           | 4/38 (10.5%)             | 12/38 (31.6%)          | ND                | ND                                   |
| Qiao et al, 2021            | 300                         | Fetuses with CHD detected by echocardiography                                       | Solo or Trio WES                      | 24/300 (8%)             | 32/300 (10.7%)           | incidental: 37/300 (12.3%) | ND                | ND                                   |
| Study                      | No. of cases with normal CMA | Inclusion criteria                                                                 | Sequencing and analysis approach | Diagnostic variants (%) | VUS or candidate genes (%) | Incidental findings (%) | Clinical impact (%) | Average TAT in days (range, if given) |
|---------------------------|-----------------------------|-----------------------------------------------------------------------------------|---------------------------------|-------------------------|---------------------------|-----------------------|---------------------|-------------------------------------|
| Sun et al, 2021<sup>a,b</sup> | 19                          | Fetuses with non-compaction cardiomyopathy on fetal echocardiogram                | Trio WES                        | 8/19 (42.1%)            | 8/19 (42.1%)             | ND                    | ND                  | ND                                  |
| Tang et al, 2021<sup>a,b</sup> | 15                          | Fetuses with suspected skeletal dysplasias on US scanning                         | Trio WES                        | 10/15 (66.7%)          | 1/15 (6.7%)             | ND                    | ND                  | ND                                  |
| Tolusso et al, 2021<sup>a</sup> | 20                          | Fetuses with congenital anomalies judged likely to be caused by an underlying monogenic condition | Trio WES                        | 8/20 (40%)            | 7/20 (35%)             | 1/20 (5%)             | 9/20 (45%)          | Ongoing pregnancies: 19.3 Terminated pregnancies: 56.8 |
| Wagner et al, 2021<sup>b,c</sup> | 15                          | Fetuses with NIHF detected on US. (Some cases previously published in Daum et al, 2019) | Trio WES                        | 7/15 (47%)             | ND                      | ND                    | ND                  | ND                                  |
| Zhang, F et al, 2021<sup>b</sup> | 12                          | Fetuses with isolated absent or hypoplastic nasal bone                            | Solo CES (2742 OMIM genes)      | 2/12 (16.7%)           | ND                      | ND                    | ND                  | ND                                  |
| Zhang, L et al, 2021<sup>a,b</sup> | 55                          | Fetuses with suspected skeletal dysplasias based on US features                  | Solo or Trio WES                | 35/55 (64%)            | 4/55 (7.3%)            | ND                    | ND                  | 21–70                               |
| Zhang, X et al, 2021<sup>a,b</sup> | 27                          | Fetuses with suspected skeletal dysplasias undergoing ToP before the third trimester | Trio WES (n = 17) or TES (n = 10 (505 skeletal dysplasia gene panel)) | 21/27 (77.8%)          | ND                      | ND                    | ND                  | ∼28                                 |
| Zhen et al, 2021<sup>b</sup> | 13                          | Fetuses with apparently isolated micrognathia identified on US at 11–14 weeks    | Trio CES (4200 OMIM disease genes) | 8/13 (61.5%)          | ND                      | ND                    | ND                  | ND                                  |
| Zhou, J et al, 2021<sup>a,b</sup> | 102                         | Fetuses with structural or growth anomalies identified by US scanning             | Trios WES and WGS in parallel   | 13/102 (12.7%)         | ND                      | ND                    | ND                  | 21 (15–27)                          |
| Zhou, X et al, 2021<sup>a,b</sup> | 28                          | Fetuses with recurrent NIHF +/- additional structural anomalies, where immune, infectious causes or thalassaemia ruled out | Trio WES                        | 10/28 (36%)           | 5/28 (18%)             | ND                    | ND                  | 28–42                               |
highly suggestive for a monogenic disorder (as determined by multidisciplinary teams [MDTs] usually including clinical genetics specialists). Seven studies were excluded from this analysis because it was not possible to determine how the cases were selected, or because mixed selection criteria were used. The other 59 studies were classified as selected or unselected cohorts (Table 2) for subgroup analysis (Figure 4). This demonstrated that the pooled incremental diagnostic yield of ES for selected cohorts was 42% (95% CI 35%–48%), which is significantly higher than that in unselected cohorts (15%, 95% CI 11%–18%; \( p < 0.0001 \)).

### 3.4 Effect of fetal phenotypic group on diagnostic yield

Further heterogeneity may arise from the varied types of fetal structural anomaly included in different studies. To investigate this, a second subgroup analysis was performed for 42 studies that reported cases disaggregated by phenotypic group, and where it was possible to distinguish isolated system anomalies from multisystem anomalies (Figure 5). Data from at least 3 studies were available for fetuses in each of 13 phenotypic subgroups based on affected body system (Table 3). The number of fetuses analysed in each subgroup varied widely, from seven cases with isolated abdominal wall abnormalities, to 773 cases with isolated cardiac abnormalities. The subgroups for multisystem anomalies, isolated skeletal anomalies, and isolated brain or central nervous system (CNS) anomalies were the next largest, with over 400 cases each. The subgroup analysis found that the incremental diagnostic yield of ES differed significantly between different phenotypic groups (\( I^2 = 96.4\%, Q = 177, p < 0.0001 \)). Table 3 details the pooled estimated incremental diagnostic yield for each subgroup, from highest to lowest. The highest diagnostic yields occurred in fetuses with skeletal abnormalities (53% [95% CI 42%–63%], \( p < 0.0001 \)), neuromuscular abnormalities, usually presenting as fetal akinesia deformation sequence (FADS), (37% [20%–54%], \( p < 0.0001 \)), and abnormalities in multiple systems (29% [22%–35%], \( p < 0.0001 \)). By contrast, the lowest rates of ES diagnoses occurred in fetuses with isolated increased nuchal translucency (NT; 2% [0%–5%], \( p = 0.04 \)), and those with gastrointestinal abnormalities (2% [−4 to 8%], \( p = 0.5 \)). There were no diagnoses from ES in 7 fetuses with abdominal wall defects and 38 fetuses with chest abnormalities.

### 3.5 Impact of prenatal ES on decision-making and clinical management

Only seven reports from six studies explicitly assessed the impact of prenatal ES on clinical management. These studies reported that ES results influenced at least one of (i) decisions about termination of pregnancy, (ii) pre- or perinatal management, (iii) future reproductive planning, or (iv) counselling on recurrence risk in 26%–79% of cases.\(^{3,26,52,62,73,74,82}\)
The findings of our review underscore the value of applying ES in the prenatal context. Meta-analysis results show an incremental diagnostic yield of 31% from ES over CMA/karyotype alone for prenatal diagnosis of fetal structural anomalies. This diagnostic yield could be considered a conservative estimate because we included only definite diagnoses (variants classified as pathogenic/likely pathogenic and deemed to be causing the fetal phenotype) in our definition of diagnostic yield. Variants reported as ‘possibly diagnostic’, ‘probably diagnostic’, or ‘potentially relevant’ were not included in our analysis but may have been re-classified as diagnostic after further study. For example, one study of 19 fetuses with severe multiple anomalies of suspected genetic aetiology found a diagnostic yield of 31.6% when strictly using ACMG guidelines, but this rose to 63% when including variants not reportable by ACMG criteria but which were plausible based on animal models, in silico prediction tools or functional studies. Other studies have also shown potentially increased diagnostic yields when probable diagnoses are included.

In this way, diagnostic yields can increase over time, as has been previously documented in large postnatal ES studies such as the UK Deciphering Developmental Disorders (DDD) study, where iterative reanalysis was performed after several years. This results from improved bioinformatics methodologies, newly discovered gene-disease associations and, in the case of prenatal ES, evolving knowledge of fetal presentations of known disorders. For example, in one study from our review, Petrovski et al. reported four variants which were re-classified (from bioinformatic signatures) to diagnostic variants over the course of the study period in light of new published evidence or additional phenotypes emerging postnatally.

Our second key finding is that the diagnostic yield of ES for fetal anomalies is higher when cases are pre-selected for likelihood of a monogenic aetiology by MDTs with expertise in genetics. This is intuitive but, as prenatal ES is increasingly implemented in clinical diagnostic settings, it highlights the value of pre-test review and MDT collaboration to direct testing where it is most likely to yield a diagnosis. Furthermore, our findings demonstrate that diagnostic yield varies significantly between phenotypic groups: the highest yield observed was for skeletal dysplasias (53%) and the lowest yields were for fetuses with isolated increased NT (2%) and isolated gastrointestinal anomalies (2%).

This understanding of likely diagnostic yield for different fetal presentations may inform clinical guidelines on offering prenatal ES, however our findings should be interpreted with care. Classification of cases by phenotype and comparison across studies is challenging because of different approaches to the classification (e.g. whether ‘soft markers’ are classed as separate abnormalities, and whether cases were categorised strictly based on isolated system anomalies or based on the system that best describes the predominant phenotype in the fetus). Moreover, fetal phenotypes evolve as organ development progresses, so classification of the same case may differ depending on the gestation at assessment. Further research is needed to clarify the diagnostic utility of prenatal ES for isolated anomalies in systems that have been less thoroughly investigated to date. For example, in this review, for fetuses with chest (respiratory) abnormalities, fetal growth restriction, gastrointestinal anomalies...
and abdominal wall defects, the sample sizes were too small to reach statistical significance in estimates of pooled diagnostic yield.

Studies examining other aspects of clinical utility such as impact on clinical management and parental decision-making will be an important complement to this, however we found only six to include in our review. Three of these studies found that negative as well as diagnostic results impacted upon parental and clinical decision-making, leading Deden et al. to conclude...
FIGURE 4  Forest plot showing individual and pooled incremental yield of prenatal exome sequencing (ES) over karyotype/chromosomal microarray analysed by pre-test case selection criteria: “unselected” cohorts underwent no prior review to determine clinical features indicating higher likelihood of monogenic aetiology; “selected” cohorts underwent pre-test expert review to select fetuses with clinical features indicating higher likelihood of monogenic disorder or to rule out non-genetic conditions.
### TABLE 2  Effect of case selection on incremental diagnostic yield of ES over CMA

| Study                          | Number of probands with normal CMA | Diagnostic yield (%) |
|-------------------------------|------------------------------------|----------------------|
| **Unselected fetuses: NO prior review/selection for features indicating higher likelihood of monogenic disease** |                                     |                      |
| Quinlan-Jones et al, 2019     | 27                                 | 10/27 (37%)          |
| Qi et al, 2020                | 80                                 | 27/80 (33.8%)        |
| Lefebvre et al, 2020          | 95                                 | 24/95 (25.3%)        |
| Fu et al, 2018                | 196                                | 47/196 (24%)         |
| Drury et al, 2015             | 24                                 | 5/24 (20.8%)         |
| Sun et al, 2020               | 66                                 | 13/66 (19.7%)        |
| Heide et al, 2020             | 62                                 | 12/62 (19.4%)        |
| Chen et al, 2020              | 105                                | 20/105 (19%)         |
| Boissel et al, 2018           | 101                                | 19/101 (18.8%)       |
| Lei et al, 2021               | 85                                 | 16/85 (18.8%)        |
| Choy et al, 2019              | 42                                 | 7/42 (16.7%)         |
| Zhou, J et al, 2021           | 102                                | 13/102 (12.7%)       |
| Xue et al, 2020               | 24                                 | 3/24 (12.5%)         |
| Lei et al, 2020               | 163                                | 20/163 (12.3%)       |
| Carss et al, 2014             | 28                                 | 3/28 (10.7%)         |
| Petrovski et al, 2019         | 234                                | 24/234 (10.3%)       |
| Li, R et al, 2020             | 260                                | 26/260 (10%)         |
| Leung et al, 2018             | 33                                 | 3/33 (9.1%)          |
| Lord et al, 2019              | 610                                | 52/610 (8.5%)        |
| Qiao et al, 2021              | 300                                | 24/300 (8%)          |
| Zhou, X et al, 2020           | 41                                 | 3/41 (7.3%)          |
| Yang et al, 2020              | 73                                 | 4/73 (5.5%)          |
| Aarabi et al, 2018            | 20                                 | 1/20 (5%)            |
| **Pooled estimated diagnostic yield for unselected subgroup** | 2771                               | 15% [95% CI = 11%–18%] p < 0.0001 |

| **Selected fetuses: Prior expert review/selection for clinical features indicating higher likelihood of monogenic disorder or ruling out non-genetic conditions** |                                     |                      |
| Han et al, 2020               | 26                                 | 23/26 (88.5%)        |
| Chandler et al, 2018          | 16                                 | 13/16 (81.3%)        |
| Zhang, X et al, 2021          | 27                                 | 21/27 (77.8%)        |
| Kucinska-Chahwan et al, 2021  | 26                                 | 18/26 (69.2%)        |
| Tang et al, 2021              | 15                                 | 10/15 (66.7%)        |
| Zhou, X et al, 2018           | 12                                 | 8/12 (66.7%)         |
| Peng et al, 2021              | 38                                 | 25/38 (65.8%)        |
| Zhang, L et al, 2021          | 55                                 | 35/55 (63.6%)        |
| Rinaldi et al, 2020           | 30                                 | 19/30 (63.3%)        |
| Liu et al, 2019               | 28                                 | 16/28 (57.1%)        |
| Aggarwal et al, 2020          | 32                                 | 18/32 (56.3%)        |
| Reischer et al, 2020          | 11                                 | 6/11 (54.5%)         |
| Al-Kouatly et al, 2021        | 22                                 | 11/22 (50%)          |
| Shamseldin et al, 2018        | 44                                 | 22/44 (50%)          |

(Continues)
that ‘the efficacy of prenatal [ES] should be evaluated by more variables than diagnostic yield alone.’ This is an area where the need for further research is pressing, including more studies exploring the perspectives of parents undergoing prenatal ES to investigate what personal utility they derive from it.

4.1 | Strengths and limitations

This systematic review is the most detailed and comprehensive in this area to date. It included 72 reports of 66 studies, many of which were published in the last 12 months. The reporting of research in the included studies was generally of high quality with a standardised approach to reporting which made it easier to compare study characteristics, and we were able to perform meta-analyses for a range of phenotypic groups as well as overall. There are however signs of possible publication bias, as indicated by funnel plot asymmetry showing a paucity of published studies with small sample size and low diagnostic yield (supporting information S1). This could reflect publication bias or indeed may represent a true relationship between small sample size and higher diagnostic yield: among the studies included in this review, smaller cohorts were often highly selected (usually after review by clinical genetics specialists) for cases where the clinical presentation suggested a monogenic aetiology. The higher a priori likelihood of a monogenic disorder among such cases translates to higher diagnostic yields, whereas the largest studies to date have been unselected cohorts, including fetuses with isolated and minor abnormalities and anomalies associated with multifactorial aetiology, as well as those more likely to have a monogenic disorder.

The main limitation of our review was the high degree of heterogeneity between included studies, limiting the precision of comparisons. On performing subgroup analyses to investigate the effect on ES diagnostic yield of case selection criteria or fetal phenotypic group, there remained substantial heterogeneity within many of the subgroups as well as between them, which suggests that neither of these factors alone fully explain the observed heterogeneity amongst

| Study                         | Number of probands with normal CMA | Diagnostic yield (%) |
|-------------------------------|------------------------------------|----------------------|
| Cao et al, 2021              | 11                                 | 5/11 (45.5%)         |
| Pooh et al, 2020             | 16                                 | 7/16 (43.8%)         |
| Li, L et al, 2020            | 19                                 | 8/19 (42.1%)         |
| Sun et al, 2021              | 19                                 | 8/19 (42.1%)         |
| De Koning et al, 2019        | 20                                 | 8/20 (40%)           |
| Tolusso et al, 2021          | 20                                 | 8/20 (40%)           |
| Correa et al, 2021           | 19                                 | 7/19 (36.8%)         |
| Dempsey et al, 2021          | 52                                 | 19/52 (36.5%)        |
| Zhou, X et al, 2021          | 28                                 | 10/28 (35.7%)        |
| Corsten-Janssen et al, 2020  | 23                                 | 8/23 (34.8%)         |
| Deden et al, 2020            | 54                                 | 18/54 (33.3%)        |
| Meier et al, 2019            | 19                                 | 6/19 (31.6%)         |
| Normand et al, 2019          | 146                                | 46/146 (31.5%)       |
| Guo et al, 2020              | 40                                 | 12/40 (30%)          |
| Aoi et al, 2021              | 17                                 | 5/17 (29.4%)         |
| Sparks et al, 2020           | 127                                | 37/127 (29.1%)       |
| Becher et al, 2020           | 35                                 | 9/35 (25.7%)         |
| Daum et al, 2019             | 77                                 | 16/77 (20.8%)        |
| Vora et al, 2020             | 102                                | 21/102 (20.6%)       |
| Westphal et al, 2019         | 30                                 | 6/30 (20%)           |
| Deng et al, 2020             | 21                                 | 3/21 (14.3%)         |
| Armes et al, 2018            | 16                                 | 2/16 (12.5%)         |
| **Pooled estimated diagnostic yield for selected subgroup** | **1293** | **42% [95% CI = 35%–48%] p < 0.0001** |

Note: The bold and underlined formatting was just for emphasis in defining the two subgroups.

Abbreviations: CMA, chromosomal microarray; ES, exome sequencing.
Studied systems included skeletal, respiratory, neuromuscular/FADS, and multisystem anomalies. The numbers of cases with normal chromosomes varied, ranging from 120 to 307 cases per study. The diagnostic yield of prenatal exome sequencing (ES) varied across studies, with yields ranging from 1.8% to 48.2%, and the risk difference varied from -0.67 to 0.37.

**Figure 5:** Forest plot showing individual and pooled incremental yield of prenatal exome sequencing (ES) over karyotype/chromosomal microarray analysed by phenotypic sub-group in studies where fetuses with anomalies confined to a single body system were distinguishable from those with multi-system anomalies.

The studies also varied in sample size, sequencing and analysis approach (gene agnostic or targeted), approach to variant interpretation, and whether or not interpretation considered postnatal/post-mortem phenotype information in addition to prenatal phenotypes. All of these factors are likely to influence the diagnostic yield. For example, even when adhering to standardised...
ACMG variant interpretation guidelines, there may be some subjectivity in how the guidelines are applied and how it is judged whether a variant explains the fetal phenotype. Studies done in a purely research setting without reporting results back to families may have interpreted variants with candidate causality more generously, leading to higher reported diagnostic yields than in studies that used

**FIGURE 5** (Continued)
more conservative reporting standards. In addition, for the phenotype subgroup analysis we only included studies that disaggregated cases by phenotypic group and distinguished cases with a single affected body system from those with multiple system anomalies. This resulted in the exclusion of 24/66 studies from this subgroup analysis. Manual inspection of the raw phenotype data from these studies revealed that 24/66 studies contained data that were not suitable for inclusion in the analysis.
studies could enable extraction and classification of more cases in their phenotypic groups and allow a more granular analysis of the incremental yield of ES in different phenotypic groups, as has been reported recently for congenital heart disease, fetal hydrops, and isolated increased NT respectively. 

5 | CONCLUSION

Prenatal ES is a powerful tool to improve the diagnosis of monogenic conditions in fetuses with structural abnormalities. However, it also presents challenges for interpreting variants in the face of incomplete and non-specific fetal phenotypes while also aiming to minimise identification of VUS and incidental findings. In ongoing pregnancies, results must be returned quickly if they are to inform decisions on termination of pregnancy or prenatal management. Prenatal ES is rapidly being adopted in many clinical settings and its successful implementation will require robust laboratory infrastructure and bioinformatics pipelines, expert parental counselling, multidisciplinary collaboration between fetal medicine and clinical and laboratory genetics specialists to offer and interpret tests appropriately, and clear guidelines to ensure equity of access. The need to sequence fetus-parent trios in parallel to minimise turn-around times means that ES remains relatively costly, so in many health systems it may be necessary to triage cases to allocate finite resources. Further research is needed on the clinical impact of prenatal ES, to better understand which pregnancies benefit most and how to prioritise potential cases in a way that maximises benefit to families, and is efficient for healthcare systems.

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CONFLICT OF INTEREST
None of the authors have any conflict of interests to declare.

ETHICS STATEMENT
This work is a systematic review of literature and no ethical committee approval was required.

DATA AVAILABILITY STATEMENT
Data sharing is not applicable to this article as no new data were created or analysed in this study. The data that support the findings of this study are derived from the cited publicly available published studies included in the review.

| Phenotypic group | Cases analysed | Pooled estimated diagnostic yield [95% CI], p-value |
|------------------|----------------|---------------------------------------------------|
| Skeletal         | 424            | 53% [42%–63%], p < 0.0001                           |
| Neuromuscular/Fetal akinesia deformation sequence (FADS) | 33            | 37% [20%–54%], p < 0.0001                           |
| Multisystem      | 698            | 29% [22%–35%], p < 0.0001                           |
| Hydrops/Oedema   | 137            | 22% [14%–31%], p < 0.0001                           |
| Central nervous system (CNS) | 417            | 17% [12%–22%], p < 0.0001                           |
| Cardiac          | 773            | 11% [7%–16%], p < 0.0001                           |
| Craniofacial     | 99             | 9% [1%–17%], p = 0.02                              |
| Congenital anomalies of kidneys and urinary tract (CAKUT) | 278            | 9% [5%–12%], p < 0.0001                           |
| Fetal growth restriction | 28            | 4% [–9 to 17%], p = 0.58                           |
| Isolated increased nuchal translucency (NT) | 290            | 2% [0%–5%], p = 0.04                              |
| Gastrointestinal | 60             | 2% [–4 to 8%], p = 0.5                             |
| Respiratory/Chest | 38            | 0 [−7 to 7%], p = 1                                |
| Abdominal wall   | 7              | 0 [−31% to 31%], p = 1                             |

Note: Phenotypic groups refer to fetuses with one or more anomalies in a single body system. Fetuses with anomalies in more than one system are classified as ‘Multisystem’. The ‘isolated increased NT’ group contains a mixture of (i) cases with isolated increased NT at presentation where it was unspecified whether additional anomalies developed later in pregnancy, and (ii) cases where isolated increased NT remained isolated throughout pregnancy. The bold and underlined formatting was just for emphasis in defining the two sub-groups.

Abbreviations: CMA, chromosomal microarray; ES, exome sequencing.
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