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[Overview of Reviews]

Interventions during pregnancy to prevent preterm birth: an overview of Cochrane systematic reviews

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ABSTRACT

Background

Preterm birth (PTB) is a major factor contributing to global rates of neonatal death and to longer-term health problems for surviving infants. Both the World Health Organization and the United Nations consider prevention of PTB as central to improving health care for pregnant women and newborn babies. Current preventative clinical strategies show varied efficacy in different populations of pregnant women, frustrating women and health providers alike, while researchers call for better understanding of the underlying mechanisms that lead to PTB.

Objectives

We aimed to summarise all evidence for interventions relevant to the prevention of PTB as reported in Cochrane systematic reviews (SRs). We intended to highlight promising interventions and to identify SRs in need of an update.

Methods

We searched the Cochrane Database of Systematic Reviews (2 November 2017) with key words to capture any Cochrane SR that prespecified or reported a PTB outcome. Inclusion criteria focused on pregnant women without signs of preterm labour or ruptured amniotic membranes. We included reviews of interventions for pregnant women irrespective of their risk status. We followed standard Cochrane methods.

We applied GRADE criteria to evaluate the quality of SR evidence. We assigned graphic icons to classify the effectiveness of interventions as: clear evidence of benefit; clear evidence of harm; clear evidence of no effect or equivalence; possible benefit; possible harm; or unknown benefit or harm. We defined clear evidence of benefit and clear evidence of harm to be GRADE moderate- or high-quality evidence with a confidence interval (CI) that does not cross the line of no effect. Clear evidence of no effect or equivalence is GRADE moderate- or high-quality evidence with a narrow CI crossing the line of no effect. Possible benefit and possible harm refer to GRADE low-quality evidence with a clear effect (CI does not cross the line of no effect) or GRADE moderate- or high-quality evidence with a wide CI. Unknown harm or benefit refers to GRADE low- or very low-quality evidence with a wide CI.

Main results

We included 83 SRs; 70 had outcome data. Below we highlight key results from a subset of 36 SRs of interventions intended to prevent PTB.

Outcome: preterm birth

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Clear evidence of benefit

Four SRs reported clear evidence of benefit to prevent specific populations of pregnant women from giving birth early, including midwife-led continuity models of care versus other models of care for all women; screening for lower genital tract infections for pregnant women less than 37 weeks' gestation and without signs of labour, bleeding or infection; and zinc supplementation for pregnant women without systemic illness. Cervical cerclage showed clear benefit for women with singleton pregnancy and high risk of PTB only.

Clear evidence of harm

No included SR reported clear evidence of harm.

No effect or equivalence

For pregnant women at high risk of PTB, bedrest for women with singleton pregnancy and antibiotic prophylaxis during the second and third trimester were of no effect or equivalent to a comparator.

Possible benefit

Four SRs found possible benefit in: group antenatal care for all pregnant women; antibiotics for pregnant women with asymptomatic bacteriuria; pharmacological interventions for smoking cessation for pregnant women who smoke; and vitamin D supplements alone for women without pre-existing conditions such as diabetes.

Possible harm

One SR reported possible harm (increased risk of PTB) with intramuscular progesterone, but this finding is only relevant to women with multiple pregnancy and high risk of PTB. Another review found possible harm with vitamin D, calcium and other minerals for pregnant women without pre-existing conditions.

Outcome: perinatal death

Clear evidence of benefit

Two SRs reported clear evidence of benefit to reduce pregnant women's risk of perinatal death: midwife-led continuity models of care for all pregnant women; and fetal and umbilical Doppler for high-risk pregnant women.

Clear evidence of harm

No included SR reported clear evidence of harm.

No effect or equivalence

For pregnant women at high risk of PTB, antibiotic prophylaxis during the second and third trimester was of no effect or equivalent to a comparator.

Possible benefit

One SR reported possible benefit with cervical cerclage for women with singleton pregnancy and high risk of PTB.

Possible harm

One SR reported possible harm associated with a reduced schedule of antenatal visits for pregnant women at low risk of pregnancy complications; importantly, these women already received antenatal care in settings with limited resources.

Outcomes: preterm birth and perinatal death

Unknown benefit or harm

For pregnant women at high risk of PTB for any reason including multiple pregnancy, home uterine monitoring was of unknown benefit or harm. For pregnant women at high risk due to multiple pregnancy: bedrest, prophylactic oral betamimetics, vaginal progesterone and cervical cerclage were all of unknown benefit or harm.

Authors' conclusions

Implications for practice

The overview serves as a map and guide to all current evidence relevant to PTB prevention published in the Cochrane Library. Of 70 SRs with outcome data, we identified 36 reviews of interventions with the aim of preventing PTB. Just four of these SRs had evidence of clear

benefit to women, with an additional four SRs reporting possible benefit. No SR reported clear harm, which is an important finding for women and health providers alike.

The overview summarises no evidence for the clinically important interventions of cervical pessary, cervical length assessment and vaginal progesterone because these Cochrane Reviews were not current. These are active areas for PTB research.

The graphic icons we assigned to SR effect estimates do not constitute clinical guidance or an endorsement of specific interventions for pregnant women. It remains critical for pregnant women and their healthcare providers to carefully consider whether specific strategies to prevent PTB will be of benefit for individual women, or for specific populations of women.

Implications for research

Formal consensus work is needed to establish standard language for overviews of reviews and to define the limits of their interpretation.

Clinicians, researchers and funders must address the lack of evidence for interventions relevant to women at high risk of PTB due to multiple pregnancy.

PLAIN LANGUAGE SUMMARY

Ways to help pregnant women avoid preterm birth

What is the issue?

Preterm birth, or being born before 37 weeks of pregnancy, is a major reason why newborns die and may also mean long-term disability for surviving infants. There are many ways healthcare providers try to prevent women from having their babies too early. Pregnant women may be encouraged to take vitamins, reduce smoking, take medicines for infections or attend regular healthcare visits. Our overview looks at different ways (or interventions) to prevent preterm birth. We searched for relevant papers in the Cochrane Library on 2 November, 2017.

Why is this important?

Preterm birth is devastating and costly for women, families and health systems. We aimed to summarise relevant information for pregnant women, healthcare workers and researchers.

What evidence did we find?

We included 83 systematic reviews with evidence about whether or not the intervention was able to reduce pregnant women's chance of having a preterm birth or a baby death. Seventy of these reviews had information about preterm birth. We categorised the evidence we found as: clear benefit or harm; no effect; possible benefit or harm; or unknown effect.

Outcome: preterm birth

Clear benefit

We were confident that the following interventions were able to help specific populations of pregnant women avoid giving birth early: midwife-led continuity models of care versus other models of care for all women; screening for lower genital tract infections; and zinc supplementation for pregnant women without systemic illness. Cervical stitch (cerclage) was of benefit only for women at high risk of preterm birth and with singleton pregnancy.

Clear harm

We found no treatment that increased women's chance of giving birth preterm.

Possible benefit

The following interventions may have helped some groups of pregnant women avoid preterm birth, but we have less confidence in these results: group antenatal care for all pregnant women; antibiotics for pregnant women with asymptomatic bacteriuria; pharmacological interventions for smoking cessation; and vitamin D supplements alone for women without health problems.

Possible harm

We found two interventions that may have made things worse for some pregnant women: intramuscular progesterone for women at high risk of preterm birth with multiple pregnancy; and taking vitamin D supplements, calcium and other minerals for pregnant women without health problems.

Outcome: perinatal death

Clear benefit

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We were confident in evidence for midwife-led continuity models of care for all pregnant women; and for fetal and umbilical Doppler for high-risk pregnant women; these interventions appeared to reduce women's chance of experiencing baby death.

Clear harm

We found no intervention that increased women's risk of baby death.

Possible benefit

We found a possible benefit with cervical stitch (cerclage) for women with singleton pregnancy and high risk of preterm birth.

Possible harm

One review reported possible harm associated with having fewer antenatal visits, even for pregnant women at low risk of pregnancy problems. The pregnant women in this review already received limited antenatal care.

Outcomes: preterm birth and perinatal death

Unknown benefit or harm

For pregnant women at high risk of preterm birth for any reason including multiple pregnancy, home uterine monitoring was of unknown benefit or harm. For high-risk pregnant women with multiple pregnancy: bedrest, prophylactic oral betamimetics, vaginal progesterone and cervical cerclage were all of unknown benefit or harm.

What does this mean?

There is valuable information in the Cochrane Library relevant to women, doctors, midwives and researchers interested in preventing early birth. We have summarised the results of systematic reviews to describe how well different strategies work to prevent early birth and baby death. We organised our information in clear figures with graphic icons to represent how confident we were in the results and to point readers toward promising treatments for specific groups of pregnant women.

Our overview found no up-to-date information in the Cochrane Library for the important treatments of cervical pessary, vaginal progesterone or cervical assessment with ultrasound. We found no high-quality evidence relevant to women at high risk of preterm birth due to multiple pregnancy. It remains important for pregnant women and their healthcare providers to carefully consider whether specific strategies to prevent preterm birth will be of benefit for individual women, or for specific populations of women.

BACKGROUND

Preterm birth (PTB) places a substantial burden on economies, health systems and families worldwide. PTB contributes disproportionately to global rates of neonatal death, and infants who survive early birth may suffer serious complications and long-term disability (Blencowe 2013; Liu 2015). PTB cost the UK economy £2.9 billion in a single year (Mangham 2009), while affected families suffered equally profound financial, social and psychological consequences (Carson 2015; Hodek 2011; Vigod 2010).

The World Health Organization (WHO) identified PTB as a "top ten" research priority to 2025 (Yoshida 2016). The United Nations has also positioned PTB research as central to achieving a reduction in newborn deaths and the Sustainable Development Goals to 2030 (Lawn 2016). Though the PTB rate in countries such as the USA has fallen, early birth contributes to high mortality in low- and middle-income countries (Blencowe 2012; Gyamfi-Bannerman 2014). The March of Dimes and the International Federation of Gynecology and Obstetrics Working Group urge researchers to think beyond current clinical and preventive strategies and to expand basic research to develop new interventions for PTB (Ferrero 2016; Martin 2017).

Description of the condition

PTB is usually defined as birth before 37 weeks + 0 days' gestation (or 259 days) (Anonymous 1977). The Global Alliance to Prevent Prematurity and Stillbirth (GAPPS) defined 'provider-initiated' PTB as early births related to: severe maternal, fetal or placental conditions, maternal request, breech presentation, cephalopelvic disproportion, or previous caesarean section. In contrast, 'spontaneous' preterm births (sPTB) were not related to severe clinical conditions (Villar 2012). When applied to 60,000+ births, the GAPPS classification system found 22% of all early births to occur spontaneously and without association to any severe condition (Barros 2015).

This overview is concerned with sPTB rather than provider-initiated PTB, but in practice the early births reported in included SRs may be a mix of the two. The United States Vital Statistics and the Office of National Statistics in the UK do not distinguish spontaneous from provider-initiated PTB when reporting national rates. A recent US study makes the distinction, but the PTB Core Outcome Set does not (Gyamfi-Bannerman 2014; van't Hooft 2016).

Description of the interventions

Many interventions to prevent PTB target all pregnant women; these may begin before or during pregnancy, such as folate supplementation or improved access to antenatal care (Iams 2008). Other interventions target specific populations of pregnant women considered to be at higher risk of PTB, such as women with multiple pregnancy or women who smoke. This overview will include all interventions that could be applied during pregnancy to prevent PTB, regardless of women's risk factors. In this overview, we included interventions for women with co-morbid conditions such as diabetes or hypertension, for women with recognised risk factors such as short cervix or multiple pregnancy, and for pregnant women without any known risk.

How the intervention might work

We do not fully understand the pathways leading to PTB, which limits implementation of appropriate clinical and public health prevention strategies. Interventions may reduce PTB via diverse pathways. Nutritional supplements may modify low body mass index (BMI) to reduce risk, while screening for asymptomatic bacteriuria may reduce the risk of infection-related PTB. We included SRs of interventions regardless of whether the mechanism of action on PTB was known.

Why it is important to do this overview

We aim to improve the care of pregnant women. There is a wealth of information in the Cochrane Library relevant to PTB prevention. Initial searches returned over 800 SRs, and yet there is no efficient way for researchers, funding organisations, clinicians or pregnant women themselves to quickly access this evidence. This overview extends previous efforts to catalogue SRs of PTB interventions (Iams 2008; Piso 2014) and applies the agreed core outcome set for PTB (van't Hooft 2016).

OBJECTIVES

We aimed to summarise all evidence for interventions relevant to the prevention of PTB as reported in Cochrane systematic reviews (SRs). We intended to highlight promising interventions and to identify SRs in need of an update.

METHODS

Criteria for considering reviews for inclusion

We applied the inclusion and exclusion criteria below when assessing SRs for this overview.

Eligibility		Inclusion	Exclusion
Participant level	Population	Pregnant women including those with risk factors or co-morbidities	Pregnant women with acute signs of preterm labour
		Pregnant women with singleton or multiple pregnancy	Pregnant women with a previous episode of preterm labour in current pregnancy
		Healthcare providers	Pregnant women with preterm prelabour ruptured membranes (PPROM)
			Pregnant women with major fetal anomalies

			Pregnant women undergoing induction of labour
Review level	Review	Cochrane systematic reviews published from 2014	We listed reviews with a relevant population and intervention published before 2014 or with a search date prior to 2014
	Trial design	Randomised clinical trials (including cluster- or quasi-randomised trials) Any setting, language or year	Non-randomised study designs
	Intervention	Behavioural, clinical or health systems interventions	Interventions for preconception only Interventions to prevent or treat miscarriage Interventions comprising fertility treatment
	Comparison	One main comparison (designated as such by review authors) Alternative intervention or no intervention (including placebo)	Additional review comparisons do not contribute to data summaries
	Outcome	Preterm birth outcome (any gestational age (GA) value) specified or reported (prespecified or post-hoc)	Preterm birth outcome (any GA value) not specified or reported

The inclusion and exclusion criteria define a pregnant population without signs of preterm labour. We excluded SRs of interventions for women with PPRM because these women receive individualised management with interventions with a short timeframe to delivery and, often, the goal of improving neonatal outcomes (e.g. magnesium sulphate for neuro protection). We excluded reviews of interventions targeting the neonate.

Search methods for identification of reviews

Only Cochrane systematic reviews were eligible for inclusion in this overview. We searched the *Cochrane Database of Systematic Reviews* (2 November 2017) with a broad strategy using all possible keywords to capture reviews relevant to preterm birth (see [Appendix 1](#)).

Data collection and analysis

We followed standard methods for overviews of reviews as described below.

Selection of reviews

We conducted eligibility in duplicate with all conflicts resolved through discussion with a third overview author.

We staged eligibility assessment. First, two overview authors assessed titles and abstracts to ensure that SRs targeted appropriate populations of pregnant women and summarised relevant interventions. Second, we set aside all relevant Cochrane protocols; we will check the status of these in future updates of this overview. We set aside potentially relevant titles published before 2014. Finally, we assessed the full text of remaining reviews

to explore further whether the population and intervention met our inclusion criteria and to confirm that the review included randomised trial evidence. We list characteristics of reviews excluded at full-text assessment.

Data extraction and management

For each included SR one overview author (NM) independently extracted data; a second overview author checked the data for accuracy. We resolved disagreements by discussion. We extracted the following characteristics from each included SR.

1. Search date
2. Number of trials included in the review
3. Number of participants included in the review
4. Overview outcomes with data
5. Authors' conclusions

Two overview authors piloted the data synthesis tables to improve the content and presentation of SR results. When the final format was agreed amongst all overview authors, one overview author created figures to present outcome data and a second overview author checked the figures.

Assessment of methodological quality of included reviews

Methodological quality of included systematic reviews

Several recent publications have explored problems with the AMSTAR tool for assessing systematic review (SR) quality ([Burda 2016](#); [Faggion 2015](#); [Pollock 2017](#); [Shea 2007](#); [Wegewitz 2016](#)). Because of the identified shortcomings of AMSTAR, we decided

against using it and explored using the Risk of Bias in Systematic Reviews (ROBIS) quality checklist instead.

ROBIS users first establish the relevance of the systematic review (SR) by checking that the population, intervention, comparator and outcomes align between the review and overview. Next, researchers use ROBIS to consider review methods: study eligibility criteria; identification and selection of studies; data collection and study appraisal; and synthesis and findings. Key questions within each of these four domains sum to an overall risk of bias for the review. Throughout, the tool prompts assessors to flag specific concerns. Finally, ROBIS requires the assessors to weigh their concerns with SR methods alongside SR conclusions for a judgement of low, high or unclear risk of bias (Whiting 2016).

Quality of trial evidence included in reviews

We did not reassess the risks of bias for the individual trials included in eligible SRs. 'Risk of bias' assessment is a component of all Cochrane Reviews; review authors consider aspects of trial conduct and reporting and note whether bias may have impacted the trial's reported effect estimates (Higgins 2011a). Further, when we evaluated pooled outcome data from a particular SR, we made use of the review authors' GRADE assessments. GRADE incorporates the review authors' 'Risk of bias' judgements for individual trials comprising pooled estimates. Where review authors did not undertake GRADE, we ourselves reviewed pooled summary statistics and trials' risks of bias according to criteria described in the GRADE Handbook (Schünemann 2013).

Data synthesis

Types of outcomes

We included Cochrane systematic reviews that specified or reported preterm birth (PTB) as one of the outcome measures. We planned to include all core outcome set domains for effectiveness, safety, maternal satisfaction and economic costs (van't Hooft 2016). Due to variation in outcome definition and reporting, and to the large number of included SRs, we present outcome data for PTB (any gestational age (GA) value) and perinatal death (PD), only (PD includes stillbirth and neonatal death). Formal consensus work with diverse stakeholders (including parents, midwives and other health professionals) confirmed GA at delivery and offspring mortality as the top two clinical outcomes for evaluating interventions to prevent preterm birth (van't Hooft 2016). Updates of this overview will explore adding additional outcome domains and/or specific outcome measures. As stated above, we expect most SR reporting of PTB outcomes may include both spontaneous and provider-initiated early births.

Data synthesis and presentation

We planned to structure data synthesis as in other Cochrane overviews (Farquhar 2018; Jones 2012; Lassi 2015; Welsh 2015). Due to the large number of included SRs, we instead created figures to present review data visually. We assigned graphic icons to communicate the direction of review effect estimates and our confidence in the available data. The graphic icons indicate mutually exclusive assessment categories: clear evidence of benefit; clear evidence of harm; clear evidence of no effect or equivalence; possible benefit; possible harm; or unknown benefit or harm. Our presentation of data is modelled on graphics produced by the WHO to describe different types of workers and

their roles in maternal and newborn care (<http://optimizemnh.org/optimizing-health-worker-roles-maternal-newborn-health/>).

We separated SRs of interventions to prevent PTB from SRs of interventions targeting other problems during pregnancy. We then organised SR outcome data in figures according to type of intervention: device, health systems, lifestyle, medicines, multiple interventions, nutrition, screening, psychosocial, and surgical. The category of 'multiple' interventions includes SRs where different types of interventions were tested for a particular problem; for example, the SR of interventions to prevent nausea in pregnancy included both drug and acupuncture interventions.

To assign a graphic icon we first considered the pooled summary statistic and confidence interval, any GRADE judgements made by the review author team and the amount of information present: the number of trials, women and events contributing to the meta-analysis. GRADE criteria include: study design limitations in trials contributing data; imprecision of effects (wide confidence intervals, sparse data or both); inconsistency (unexplained heterogeneity as measured by the I^2 statistic); indirectness (when trials test a different population, intervention, comparison or outcome than is specified in the review); and publication bias. All randomised trial evidence begins as of high certainty but may be downgraded by one for a 'serious' problem or by two for 'very serious' problem with any of these criteria. The GRADEpro Guideline Development Tool sums downgrading decisions into a quality rating of high, moderate, low, or very low (Schünemann 2013).

There is a large literature on conducting, interpreting and communicating GRADE assessment (Guyatt 2011a; Santesso 2015a; Santesso 2015b). Researchers continue to explore ways to express SR effect estimates and to delineate the scope of their interpretation (Hultcrantz 2017). Our graphic icons communicate the GRADE quality assessment found in SR 'Summary of findings' tables. Where SR authors did not GRADE relevant trial evidence for PTB or PD, two overview authors independently applied GRADE criteria and resolved disagreement through discussion (GRADE assessments not shown).

For all included SRs, two overview authors independently assessed data for the outcomes of PD and PTB to assign a graphic icon. We resolved disagreements through discussion or by consultation with a third overview author.

Limitations of the overview and bias in the review process

One limitation of our overview is its restriction to Cochrane systematic reviews. Including all relevant, published SRs may have captured more current evidence on important topics. Cochrane Reviews are also all published in the English language. Finally, our search strategy did not include unpublished, industry and governmental SRs.

To minimise bias in the overview process, we followed standard review methods including duplication of effort. Where an overview author was also an author of a relevant SR, they were excluded from eligibility assessment and data synthesis for that SR.

RESULTS

Search results

We searched the *Cochrane Database of Systematic Reviews* (2 November 2017) with a list of key words to capture all systematic reviews (SRs) specifying or reporting any preterm birth (PTB) outcome (See [Appendix 1](#) for the search strategy). We documented

assessment of titles and full-text reviews in a study flow diagram (See [Figure 1](#)). We used [Covidence](#) software to manage eligibility assessment for 898 titles. Two overview authors independently assessed titles and full texts against prespecified inclusion and exclusion criteria. We resolved discrepancies between overview authors by discussion with a third author. No overview author assessed his or her own SR (several eligible reviews were authored by members of the overview team).

Figure 1. Study flow diagram. Search date 2 November 2017

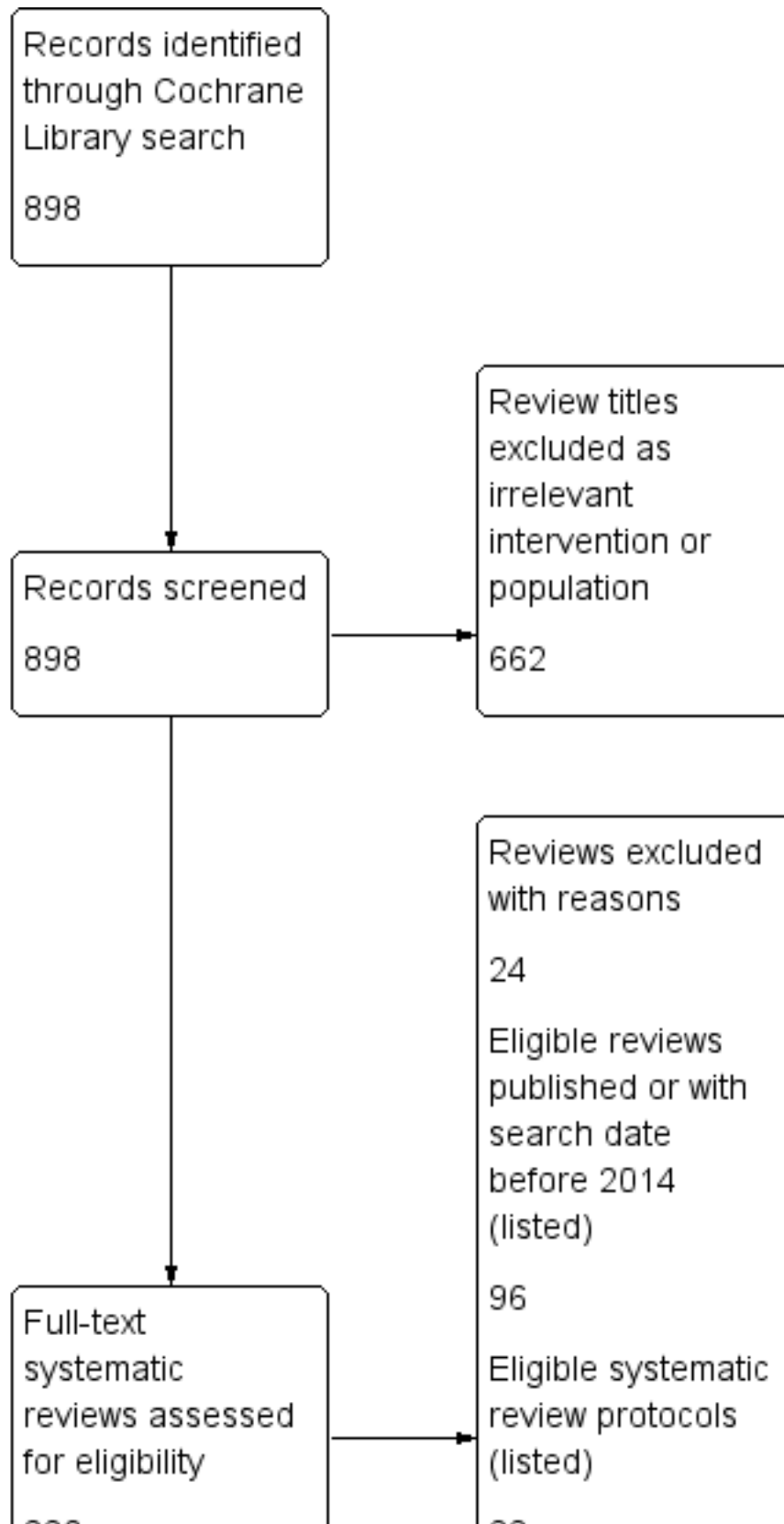
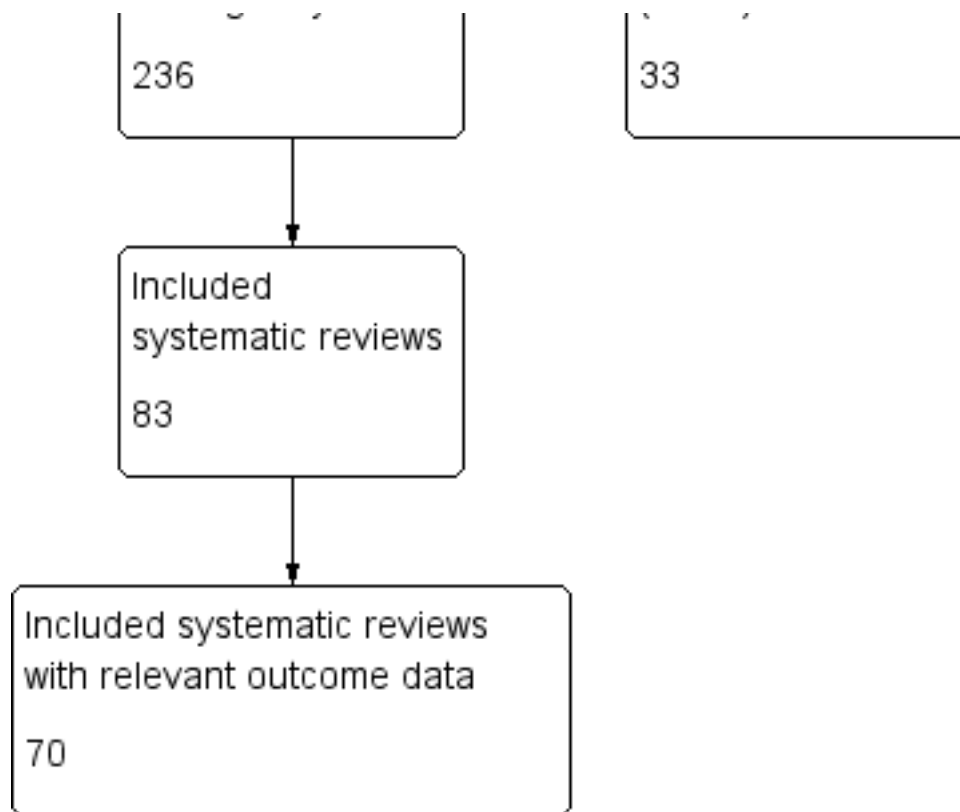


Figure 1. (Continued)



We included 83 SRs; their characteristics appear in [Table 1](#); 70 included reviews reported relevant outcome data. We documented reasons for exclusion of 24 reviews at full-text assessment in [Table 2](#). We identified 33 protocols for SRs potentially relevant to future versions of this overview; these titles will be assessed for future versions of the overview.

Out-of-date systematic reviews

Our protocol stated that we would apply a decision tree to potentially eligible but out-of-date reviews - SRs published before 2014 or with a search date before 2014. We found 96 reviews in this category. It was not feasible to conduct new searches for all of these reviews. Some reviews in this category are currently in the editorial process, while other reviews may summarise evidence for interventions no longer in use. We regret that we were unable to complete further assessment of these SR titles. As one peer reviewer pointed out - there may be interventions associated with harm in the list of out-of-date reviews, because authors would be less likely to complete new trials or to update reviews of interventions where there was known evidence of harm. We listed the titles in [Appendix 2](#).

Description of included reviews

We documented key characteristics of 83 included studies in [Table 1](#), including the search date, number of trials, number of women and the authors' conclusions about the effectiveness of the interventions. The language of authors' conclusions is directly quoted from the original Cochrane published reviews. Of 83 included SRs, seven had no randomised trials (i.e. these are 'empty reviews'). Six further SRs reported no trial data for either

overview outcome of PTB or perinatal death (PD). All remaining 70 SRs contributed effect estimates for at least one overview outcome.

Methodological quality of included reviews

As noted above, researchers identified problems with the AMSTAR tool to assess SR quality ([Burda 2016](#); [Pollock 2017](#)). We decided not to use AMSTAR and explored using the ROBIS tool instead. We decided against using ROBIS as well, because all SRs included in this overview followed standard methodology recommended in the *Cochrane Handbook for Systematic Reviews of Interventions* ([Higgins 2011b](#)). For more recent SR updates, review authors also followed methods set out by the Cochrane Editorial Unit (i.e. MECIR standards [Higgins 2016](#)). We are confident that all included SRs are of low risk of methods bias.

Effect of interventions

Notes for Interpreting Graphic Icons

Graphic Icons

We summarised systematic review (SR) evidence for the outcomes of PTB and PD by assigning one of six graphic icons to indicate our confidence in and interpretation of the available evidence. [Figure 2](#) shows the icons and explains the mutually exclusive assessment categories. The first three categories represent GRADE moderate- or high-quality evidence for which we found either clear benefit, clear harm or clear evidence of no effect (i.e. equivalence with a comparator). These categories are identified by a green tick, a red-cross and a green equal-sign icon, respectively. For 'clear' benefit or harm, the confidence interval associated with the effect size did not cross the line of no effect. For 'clear evidence of no

effect or equivalence' we considered a confidence interval within the range of risk ratio (RR) 0.75 to 1.25 as sufficiently narrow to indicate a minimal effect relative to the comparator; these are

thresholds recommended by GRADE (Guyatt 2011b). The overview author team individually discussed the handful of cases where an SR reported the odds ratio rather than relative risk.

Figure 2. All icons by Freepik at www.flaticon.com.



Continuing down Figure 2, two yellow icons represent evidence for which we had less confidence due to risks of bias in trials, imprecision or other GRADE criteria for downgrading evidence quality (Schünemann 2013). The yellow icons indicate possible benefit (the plus sign) or possible harm (the minus sign). Yellow icons represent GRADE moderate- or high-quality evidence when the confidence interval crossed the line of no effect. Yellow icons also represent GRADE low-quality evidence when the finding was of clear benefit or harm (i.e. the confidence interval did not cross RR 1.0). We must emphasise the reduced confidence and caution we intended to communicate when assigning the yellow graphic icons.

Finally, we assigned a blue question mark icon to indicate 'unknown harm or benefit' associated with the relative effect of an intervention versus a comparator. The blue question-mark icon indicates GRADE low-quality evidence with a confidence interval crossing the line of no effect as well as all GRADE very low-quality evidence.

Evidence 'Quality' or 'Certainty'?

Our categories of assessment follow GRADE ratings of high, moderate, low or very low to indicate the quality of evidence rather than its certainty. Though evidence quality and certainty are often used interchangeably, 'certainty' represents newer language that extends GRADE methods to the development of formal clinical guidance (Hultcrantz 2017). This overview does not constitute formal clinical guidance. The SR evidence summarised here would be considered "non-contextualised" rather than "contextualised" in the newer GRADE framework that aims to situate evidence in clinical, economic and geographic contexts.

SRs included in this overview applied GRADE criteria to a single outcome independently rather than to a set of critical outcomes valued together, as guideline developers might. SRs did not pre-specify thresholds for interpreting effect size or confidence intervals, and, though most SRs identified trial countries in their

'Summary of findings' tables, none reported this geographic information *per outcome*. No SR calculated a minimum information size required for meta-analysis. Most SRs did not incorporate formal analyses of the economic cost or of the acceptability of interventions to pregnant women. Finally, included Cochrane SRs do not make clinical recommendations (see Higgins 2011b, section 12.7.4 "Common errors in reaching conclusions").

Therefore, a GRADE of moderate or high quality in this overview reflects our confidence that the pooled estimate is correct (i.e. its 'quality') rather than an endorsement of the intervention's use in a specific clinical setting or population of pregnant women. Following this, the overview graphic icons do not represent clinical recommendations for the use of interventions. The graphic icons represent the quality of the evidence found in reviews and direct readers to the full text of SRs for further scrutiny.

Presentation of results

Due to the large number of reviews we wanted to included in this overview, and because we wanted to avoid presenting 50 pages of data tables, we took decisions to limit the information presented in tables and figures. Figures 3 to 19 summarise evidence from the 83 included reviews.

We grouped SRs specifically targeting preterm birth prevention (Figures 3 to 10) and SRs targeting either diabetes or other problems women might face during pregnancy (Figures 11 to 19). Interventions targeting PTB had this indication stated in the SR title or had PTB as a primary review outcome. We also grouped SRs according to intervention type: device, health systems, lifestyle, medicines, multiple interventions, nutrition, screening, psychosocial, and surgical. In all figures below 'Not reported' means that no trial included in the SR reported data for that particular outcome. 'Not a review outcome' means that the SR did not search eligible trials for that outcome because the outcome was not a review priority. Finally, 'No trials' indicates an empty SR.

Results from all SRs

We included 83 SRs; seven of these were empty reviews (i.e. included no randomised trials), and a further six SRs had no trial data for either of the overview outcomes of PD or PTB. We categorised available outcome effect estimates as of clear benefit, clear harm, clear evidence of no effect or equivalence, possible

benefit, possible harm or unknown benefit or harm as explained in Figure 2.

The figures of graphic icons are not intended to replace close inspection of individual SRs and the often complicated results found there. Rather, our overview intends to assess the quality of evidence for the key outcomes of preterm birth and perinatal death and to signpost readers interested in specific effect estimates to full SRs. Individual SRs report complete details of the relevant clinical populations and geographic settings, trials' risks of bias, and forest plots indicating the relative contribution of individual trials to a pooled estimate. The Characteristics of included studies table (Table 1) may be read in tandem with the figures of graphic icons; this table includes information on the review search date, the number of clinical trials and pregnant women included in the review, overview outcome measures reported and review authors' conclusions.

Not all included SRs reported on both overview outcomes of preterm birth and perinatal death. For some SRs, our figures summarise effect estimates for more than one comparison. No SR reported spontaneous PTB as distinct from PTB, and we expect most results below represent a mix of spontaneous and provider-initiated PTB.

We report results for preterm birth and perinatal death separately. For preterm birth, we summarised any outcome measure of preterm birth (any gestational age (GA) value) and followed individual SR decisions to prioritise specific time points (e.g. if we found PTB < 34 weeks' gestation in the 'Summary of findings' table and other time points in the text, we took the measure used in the 'Summary of findings' table). We note that perinatal death as reported in SRs refers to stillbirths and neonatal deaths counted together.

Results from reviews of interventions targeting PTB prevention

Thirty-six SRs summarised evidence for interventions with the aim of preterm birth prevention. The following figures present graphic icons assigned to SR evidence for the outcomes of preterm birth and perinatal death: Figure 3 Device; Figure 4 Health systems; Figure 5 Lifestyle; Figure 6 Medicines; Figure 7 Nutrition; Figure 8 Psychosocial; Figure 9 Screening; and Figure 10 Surgical. Because these interventions were intended as preterm birth prevention, we include details of any preterm birth-related risk factors in the target populations.

Figure 3.

Device interventions for preterm birth prevention	Risk factor	Main Comparison	Perinatal death	Preterm birth
Home uterine monitoring for detecting preterm labour (Urquhart 2017)	Women considered to be of high risk of PTB for any reason including multiple pregnancy	Home uterine monitoring vs standard care		

Figure 4.








Health systems interventions for preterm birth prevention	Risk factor	Main comparison	Perinatal death	Preterm birth
Alternative versus standard packages of antenatal care for low-risk pregnancy (Dowswell 2015)	Women at low risk of developing pregnancy or labour complications	Reduced visits vs standard antenatal care		
Group versus conventional antenatal care for women (Catling 2015)	All pregnant women	Group vs individual antenatal care		
Incentives for increasing prenatal care use by women in order to improve maternal and neonatal outcomes (Till 2015)	All pregnant women	Incentive vs no incentive	Not reported	Not reported
Midwife-led continuity models versus other models of care for childbearing women (Sandall 2016)	All pregnant women	Midwife-led vs other models of care		
Specialised antenatal clinics for women with a multiple pregnancy for improving maternal and infant outcomes (Dodd 2015b)	Multiple pregnancy	Specialised antenatal clinic vs standard antenatal care		Not reported

Figure 5.





Lifestyle interventions for preterm birth prevention	Risk factor	Main comparison	Perinatal death	Preterm birth
Bed rest in singleton pregnancies for preventing preterm birth (Sosa 2015)	Women at high risk of PTB due to prior PTB; threatened PTL; positive screening test (fetal fibronectin or short cervical length on ultrasound); low pre-pregnancy BMI; women identified through risk scoring systems Women without PPRM or multiple pregnancy	Bed rest vs placebo or no treatment	Not reported	
Bed rest with and without hospitalisation in multiple pregnancy for improving perinatal outcomes (da Silva Lopes 2017)	Multiple pregnancy	Strict bed rest in hospital vs no activity restriction at home		
Effects of restricted caffeine intake by mother on fetal, neonatal and pregnancy outcomes (Jahanfar 2015)	All pregnant women	Caffeinated vs decaffeinated coffee	Not reported	
Nutritional advice for improving outcomes in multiple pregnancies (Bricker 2015b)	Multiple pregnancy	No trials		

Figure 6.













Medicines for preterm birth prevention	Risk factor	Main comparison	Perinatal death	Preterm birth
Antibiotic prophylaxis during the second and third trimester to reduce adverse pregnancy outcomes and morbidity (Thinkhamrop 2015)	All pregnant women and women with a history of PTB or LBW infant, BV in the current pregnancy, or pre-pregnancy weight of < 50 kg	Prophylactic antibiotics vs placebo		
Antibiotics for asymptomatic bacteriuria in pregnancy (Smaill 2015)	Bacteriuria (asymptomatic)	Antibiotic vs no treatment	Not reported	
Duration of treatment for asymptomatic bacteriuria during pregnancy (Widmer 2015)	Bacteriuria (asymptomatic)	Single dose vs short course (4-7 days)	Not reported	
Interventions for treating genital Chlamydia trachomatis infection in pregnancy (Cluver 2017)	Chlamydia trachomatis infection	Erythromycin vs placebo		
Prenatal administration of progestogens for preventing spontaneous preterm birth in women with a multiple pregnancy (Dodd 2017)	Multiple pregnancy	Intra-muscular Progesterone vs placebo		
		Vaginal Progesterone vs placebo		
Prophylactic oral betamimetics for reducing preterm birth in women with a twin pregnancy (Yamasmit 2015)	Multiple pregnancy	Oral betamimetic vs placebo		

Figure 7.

Nutrition interventions for preterm birth prevention	Risk factor	Main comparison	Perinatal death	Preterm birth
Calcium supplementation (other than for preventing or treating hypertension) for improving pregnancy and infant outcomes (Buppasiri 2015)	All pregnant women	Calcium supplementation vs placebo or no treatment		
Daily oral iron supplementation during pregnancy (Peña-Rosas 2015a)	All pregnant women	Any iron vs supplements without iron or no treatment/placebo		
Intermittent oral iron supplementation during pregnancy (Peña-Rosas 2015b)	Pregnant women without HIV or TB	Intermittent vs daily regimen		
Iodine supplementation for women during the preconception, pregnancy and postpartum period (Harding 2017)	Pregnant women without thyroid disorders, HIV, or TB	Any iodine vs no iodine or no treatment		
Multiple-micronutrient supplementation for women during pregnancy (Haider 2017)	Pregnant women without HIV	MMN with iron and folic acid vs iron with or without folic acid		
Pyridoxine (vitamin B6) supplementation during pregnancy or labour for maternal and neonatal outcomes (Salam 2015c)	All pregnant women	Pyridoxine B6 vs control	Not a review outcome	Not reported
Vitamin C supplementation in pregnancy (Rumbold 2015a)	All pregnant women	Vitamin C alone or with other supplements vs placebo or control		
Vitamin D supplementation for women during pregnancy (De-Regil 2016)	Pregnant women without pre-existing conditions (i.e. diabetes)	Vitamin D alone vs placebo Vitamin D, calcium and other vs other	 	
Vitamin E supplementation in pregnancy (Rumbold 2015b)	All pregnant women	Vitamin E vs placebo, no treatment or other supplements		
Zinc supplementation for	Pregnant	Zinc vs no zinc		

Figure 7. (Continued)



Zinc supplementation for improving pregnancy and infant outcome (Ota 2015a)	Pregnant women without systemic illness	supplements Zinc vs no zinc		
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Figure 8.


Psychosocial interventions for preterm birth prevention	Risk factor	Main Comparison	Perinatal death	Preterm birth
Psychosocial interventions for pregnant women in outpatient illicit drug treatment programs compared to other interventions (Terplan 2015)	Women in illicit drug treatment programmes for any substance abuse or drug dependency including methadone	Motivational-interviewing based or contingency-management based interventions vs usual care	Unclear if review outcome	

Figure 9.












Screening for preterm birth prevention	Risk factor	Main comparison	Perinatal death	Preterm birth
Antenatal lower genital tract infection screening and treatment programs for preventing preterm delivery (Sangkomkamhang 2015)	Women not in PTL, < 37 weeks' gestation, no vaginal bleeding, asymptomatic of lower genital tract infection	Lower genital tract infection screening vs no screening	Not reported	
Risk-scoring systems for predicting preterm birth with the aim of reducing associated adverse outcomes (Davey 2015)	Women not in PTL, < 37 week's gestation, regardless of medical history or other risk factors. Multiple pregnancy included.	No trials		
Routine ultrasound in late pregnancy (after 24 weeks' gestation) (Bricker 2015a)	Women after 24 weeks' gestation in both unselected and designated low-risk populations	Routine ultrasound > 24 weeks vs no/concealed/selective ultrasound		
Screening and subsequent management for thyroid dysfunction pre-pregnancy and during pregnancy for improving maternal and infant health (Spencer 2015)	Women before or during pregnancy without a pre-existing diagnosis of thyroid dysfunction. Single or multiple pregnancy.	Universal screening vs case finding in pregnancy for any thyroid dysfunction		
Screening for genital chlamydia infection (Low 2016)	Women over the age of 13. No specific pregnancy risk stated.	No trials in pregnancy		

Figure 10.

Surgical interventions for preterm birth prevention	Risk factor	Main comparison	Perinatal death	Preterm birth
Cervical stitch (cerclage) for preventing preterm birth in multiple pregnancy (Rafael 2014)	Multiple pregnancy	Cerclage vs no cerclage		
Cervical stitch (cerclage) for preventing preterm birth in singleton pregnancy (Alfirevic 2017a)	Women of high risk of PTB due to prior PTB, prior cervical surgery or short cervix on ultrasound	Cerclage vs no cerclage		
Reduction of the number of fetuses for women with a multiple pregnancy (Dodd 2015a)	Multiple pregnancy	No trials		
Treating periodontal disease for preventing adverse birth outcomes in pregnant women (Iheozor-Ejiofor 2017)	Women with periodontal disease	Periodontal treatment vs no treatment		

Systematic reviews with no included trials

The following SRs targeted preterm birth prevention but included no randomised clinical trials: nutritional advice in multiple pregnancy (Bricker 2015b); risk-scoring systems for predicting preterm birth (Davey 2015); screening for genital Chlamydia (Low 2016); and reduction in the number of fetuses in multiple pregnancy (Dodd 2015a).

Impact on preterm birth rates

Clear evidence of preterm birth reduction

Clear evidence of benefit signals evidence of GRADE moderate or high quality with a confidence interval that does not cross the line of no effect. We found four SRs reporting clear evidence of benefit to prevent specific populations of pregnant women from giving birth early. These reviews were: midwife-led continuity models of care versus other models of care for all women (Sandall 2016); cervical cerclage for women at high risk of preterm birth and with singleton pregnancy (Alfirevic 2017a); zinc supplementation for pregnant women without systemic illness (Ota 2015a); and antenatal lower genital tract infection screening for pregnant women less than 37 weeks' gestation and without signs of labour, bleeding or infection (Sangkomkhang 2015).

Clear evidence of harm

Clear evidence of harm refers to evidence of GRADE moderate or high quality with a confidence interval that does not cross the line of no effect. No included SR reported clear evidence of harm

associated with an intervention; that is, we found no intervention that increased women's chances of giving birth preterm.

Clear evidence of no effect or equivalence

The following SRs reported evidence of GRADE moderate or high quality with a narrow confidence interval crossing the line of no effect. These interventions we categorised to be of clear evidence of no effect or equivalence with a comparator: alternative versus standard packages of antenatal care for women at low risk of developing pregnancy complications (Dowswell 2015); bed rest for women with high risk of preterm birth and singleton pregnancy (Sosa 2015); antibiotic prophylaxis during the second and third trimester for all women or for women with high risk of PTB (Thinkhamrop 2015); calcium supplementation (other than for hypertension) for all pregnant women (Buppasiri 2015); daily oral iron supplementation for all pregnant women (Peña-Rosas 2015a); vitamin C supplementation for all pregnant women (Rumbold 2015a); vitamin E supplementation for all pregnant women (Rumbold 2015b); routine ultrasound in later pregnancy for women in unselected or low-risk populations (Bricker 2015a); and screening and subsequent management for thyroid dysfunction for women before or during pregnancy without pre-existing thyroid problems (Spencer 2015).

Possible benefit

Possible benefit refers to GRADE low-quality evidence with clear benefit (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide

confidence intervals crossing the line of no effect. We found evidence of possible benefit to specific populations of women reported in the following SRs: group antenatal care for all pregnant women (Catling 2015); antibiotics for pregnant women with asymptomatic bacteriuria (Smaill 2015); pharmacological interventions for smoking cessation for pregnant women who smoke (Coleman 2015); and vitamin D supplements alone for women without pre-existing conditions such as diabetes (De-Regil 2016).

Possible harm

Possible harm refers to GRADE low-quality evidence with clear harm (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide confidence intervals crossing the line of no effect. One SR reported possible harm for the effects of intramuscular progesterone for women at high risk of preterm birth with multiple pregnancy (Dodd 2017). Another review found possible harm with vitamin D supplements, calcium and other minerals for pregnant women without pre-existing conditions (De-Regil 2016).

Unknown harm or benefit

Ten SRs reported evidence for the outcome of PTB that we categorised to be of unknown harm or benefit due to effect estimates with wide confidence intervals and evidence of GRADE low or very low quality: home uterine monitoring for high-risk pregnant women (Urquhart 2017); bed rest for women with multiple pregnancy (da Silva Lopes 2017); restricted caffeine intake for all women (Jahanfar 2015); duration of treatment for women with asymptomatic bacteriuria (Widmer 2015); treating genital *Chlamydia trachomatis* infection in pregnancy (Cluver 2017); vaginal progesterone for women with multiple pregnancy (Dodd 2017); prophylactic oral betamimetics for women with multiple pregnancy (Yamasmit 2015); intermittent oral iron supplementation for women without tuberculosis or HIV (Peña-Rosas 2015a); iodine supplementation for pregnant women without thyroid disorders, tuberculosis or HIV (Harding 2017); and multiple micronutrient supplements for pregnant women without HIV (Haider 2017).

Systematic reviews not reporting data for a preterm birth outcome

The following SRs included trials that did not report data for any preterm birth outcome measure: increasing prenatal care use (Till 2015); specialised antenatal clinics for multiple pregnancy (Dodd 2015b); and pyridoxine (vitamin B6) supplementation (Salam 2015c).

Impact on perinatal death in preterm birth prevention reviews

Clear evidence of reduction in perinatal deaths

Two SRs reported evidence of GRADE moderate or high quality with clear evidence of benefit (the confidence interval does not cross the line of no effect): midwife-led continuity models of care for all pregnant women (Sandall 2016); and fetal and umbilical Doppler for pregnant women at high risk (Alfirevic 2017b).

Clear evidence of harm

No SR reported clear evidence of harm associated with the intervention; no intervention summarised here increased a woman's chance of PD.

Clear evidence of no effect or equivalence

The following SRs reported GRADE moderate- or high-quality evidence with narrow confidence intervals crossing the line of no effect, suggesting the intervention had no effect or equivalence to a comparator: antibiotic prophylaxis during the second and third trimester for all pregnant women and women at high risk of PTB (Thinkhamrop 2015); calcium supplementation (other than for preventing or treating hypertension) for all pregnant women (Buppasiri 2015); vitamin C for all pregnant women (Rumbold 2015a); vitamin E supplementation for all pregnant women (Rumbold 2015b); routine ultrasound in late pregnancy (after 24 weeks' gestation) for unselected pregnant women and women of low risk (Bricker 2015a); and screening and subsequent management for thyroid dysfunction before or during pregnancy, for women without pre-existing thyroid disorders (Spencer 2015).

Possible benefit

Possible benefit refers to GRADE low-quality evidence with clear benefit (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide confidence intervals crossing the line of no effect. One SR reported possible benefit of cervical stitch (cerclage) for women with singleton pregnancy and high risk of preterm birth (Alfirevic 2017a).

Possible harm

Possible harm refers to GRADE low-quality evidence with clear harm (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide confidence intervals crossing the line of no effect. One SR reported possible harm associated with a reduced schedule of antenatal visits for pregnant women at low risk of pregnancy complications; these pregnant women already received antenatal care in settings with limited resources (Dowswell 2015).

Unknown harm or benefit

Evidence summarised in the following SRs was of GRADE low or very low quality with a wide confidence interval crossing the line of no effect: home uterine monitoring for high-risk pregnant women (Urquhart 2017); group versus conventional antenatal care for all pregnant women (Catling 2015); specialised antenatal clinics for women with multiple pregnancy (Dodd 2015b); bedrest for women with multiple pregnancy (da Silva Lopes 2017); treating genital *Chlamydia trachomatis* infection in pregnancy (Cluver 2017); intramuscular progesterone for women with multiple pregnancy (Dodd 2017); vaginal progesterone for women with multiple pregnancy (Dodd 2017); prophylactic oral beta mimetics for women with multiple pregnancy (Yamasmit 2015); daily oral iron supplementation for all pregnant women (Peña-Rosas 2015a); intermittent oral iron supplements for women without tuberculosis or HIV (Peña-Rosas 2015b); iodine supplementation for pregnant women without thyroid disorders, tuberculosis or HIV (Harding 2017); multiple micronutrient supplements for pregnant women without HIV (Haider 2017); vitamin D supplementation for pregnant women without pre-existing conditions such as diabetes (De-Regil 2016); zinc supplementation for pregnant women without systemic illness (Ota 2015a); Cervical stitch (cerclage) for women with multiple pregnancy (Rafael 2014); and Treating periodontal disease (Iheozor-Ejiofor 2017).

Systematic reviews not reporting data for the outcome perinatal death

The following SRs included trials that did not report the outcome of perinatal death: incentives for increasing prenatal care for all pregnant women (Till 2015); bed rest for women with singleton pregnancy and high risk of PTB (Sosa 2015); restricted caffeine for all pregnant women (Jahanfar 2015); antibiotics for asymptomatic bacteriuria (Smaill 2015); duration of treatment for asymptomatic bacteriuria (Widmer 2015); and antenatal lower genital tract infection screening for women without signs of preterm labour (Sangkomkamhang 2015).

Results from reviews of interventions targeting diabetes or other problems during pregnancy

Forty-seven SRs summarised relevant evidence for interventions targeting diabetes or other problems women might face during

pregnancy. All reviews were included in this overview because they included an eligible population of pregnant women and prespecified or reported data for an overview priority outcome (perinatal death or any measure of preterm birth). The following figures present the graphic icons assigned to SR evidence for these outcomes: Figure 11 Health systems; Figure 12 Lifestyle; Figure 13 Medicines targeting diabetes; Figure 14 Medicines targeting other pregnancy conditions; Figure 15 Multiple interventions; Figure 16 Nutrition; Figure 17 Psychosocial; Figure 18 Screening; Figure 19 Surgical. We found no SRs of Device interventions, so there is no figure for this category.

Figure 11.





		Perinatal death	Preterm birth
Health systems interventions for diabetes	Main comparison		
Interconception care for women with a history of gestational diabetes for improving maternal and infant outcomes (Tieu 2017a)	No trials		
Health systems interventions for other pregnancy conditions	Main comparison	Perinatal death	Preterm birth
Directed preconception health programs and interventions for improving pregnancy outcomes for women who are overweight or obese (Opray 2015)	No trials		
Health system and community level interventions for improving antenatal care coverage and health outcomes (Mbuagbaw 2015)	One intervention vs none		
	Combined intervention vs none		

Figure 12.















Lifestyle interventions targeting diabetes	Main comparison	Perinatal death	Preterm birth
Diet and exercise interventions for preventing gestational diabetes mellitus (Bain 2015)	Combined diet and exercise vs control		
Diet or exercise, or both, for preventing excessive weight gain in pregnancy (Muktabhant 2015)	All diet and/or exercise interventions vs standard/other care	Not reported	
Dietary advice interventions in pregnancy for preventing gestational diabetes mellitus (Tieu 2017b)	Dietary advice interventions vs standard care		Not reported
Different types of dietary advice for women with gestational diabetes mellitus (Han 2017)	Low-moderate GI diet vs moderate-high GI diet	Not reported	
	Energy-restricted diet vs no energy-restricted diet		
Exercise for pregnant women with gestational diabetes for improving maternal and fetal outcomes (Brown 2017b)	Exercise vs control		
Lifestyle interventions for the treatment of women with gestational diabetes (Brown 2017a)	Lifestyle intervention vs usual care or diet alone		
Lifestyle interventions targeting other pregnancy conditions	Main comparison	Perinatal death	Preterm birth
Antenatal dietary education and supplementation to increase energy and protein intake (Ota 2015b)	Nutritional education vs no counselling or education		
Interventions for preventing or reducing domestic violence against pregnant women (Jahanfar 2014)	Any intervention to prevent violence (all interventions) vs standard care	Not reported	

Figure 13.











Medicines targeting diabetes	Main comparison	Perinatal death	Preterm birth
Continuous subcutaneous insulin infusion versus multiple daily injections of insulin for pregnant women with diabetes (Farrar 2016)	Continuous subcutaneous infusion vs multiple daily injections of insulin		
Different insulin types and regimens for pregnant women with pre-existing diabetes (O'Neill 2017)	Lispro vs regular insulin	Not reported	
Different intensities of glycaemic control for women with gestational diabetes mellitus (Martis 2016)	Strict glycaemic control vs Less strict glycaemic control	Not reported	
Different intensities of glycaemic control for pregnant women with pre-existing diabetes (Middleton 2016)	Very tight vs tight-moderate glycaemic control targets		
Impact of Haemophilus influenzae type B (Hib) and viral influenza vaccinations in pregnancy for improving maternal, neonatal and infant health outcomes (Salam 2015b)	Viral influenza vaccine vs placebo vaccine		
Oral anti-diabetic agents for women with established diabetes/impaired glucose tolerance or previous gestational diabetes planning pregnancy, or pregnant women with pre-existing diabetes (Tieu 2017c)	Metformin vs insulin		
Oral anti-diabetic pharmacological therapies for the treatment of women with gestational diabetes (Brown 2017c)	Oral anti-diabetic pharmacological therapies vs placebo	Not reported	Not reported

Figure 14.







Medicines targeting other pregnancy conditions	Main comparison	Perinatal death	Preterm birth
Drugs for preventing malaria in pregnant women in endemic areas (Radeva-Petrova 2014)	Malaria chemoprevention (any regimen) vs placebo or no intervention		
Effect of administration of antihelminthics for soil-transmitted helminths during pregnancy (Salam 2015a)	Antihelminthics vs control		
Pharmacological interventions for promoting smoking cessation during pregnancy (Coleman 2015)	Nicotine replacement therapy vs control		

Figure 15.







Multiple interventions targeting other pregnancy conditions	Main comparison	Perinatal death	Preterm birth
Interventions for managing asthma in pregnancy (Bain 2014)	Inhaled corticosteroid vs control		
Interventions for heartburn in pregnancy (Phupong 2015)	<ul style="list-style-type: none"> • Drug vs placebo or no treatment • Drug vs diet/ lifestyle intervention • Acupuncture vs no treatment 	Not a review outcome	Not reported for any comparison
Interventions for leg cramps in pregnancy (Zhou 2015)	Oral Magnesium vs placebo/no treatment	Not reported	Not reported
Interventions for treating hyperemesis gravidarum (Boelig 2016)	<ul style="list-style-type: none"> • Acupuncture vs placebo • Corticosteroid vs promethazine • Corticosteroid vs placebo 	All comparisons 	All comparisons 
Interventions for nausea and vomiting in early pregnancy (Matthews 2015)	Comparisons not prioritised by review authors. <ul style="list-style-type: none"> • Ginger vs Vitamin B6 	Not reported	
Interventions for preventing recurrent urinary tract infection during pregnancy (Schneeberger 2015)	Nitrofurantoin and close surveillance vs close surveillance only	Not reported	
Interventions for varicose veins and leg oedema in pregnancy (Smyth 2015)	Rutoside vs placebo	Not reported	Not reported

Figure 16.








Nutrition interventions targeting diabetes	Main comparison	Perinatal death	Preterm birth
Dietary supplementation with myo-inositol in women during pregnancy for treating gestational diabetes (Brown 2016)	Myo-inositol vs placebo	Not reported	
Nutrition interventions targeting other pregnancy conditions	Main comparison	Perinatal death	Preterm birth
Calcium supplementation commencing before or early in pregnancy, or food fortification with calcium, for preventing hypertensive disorders of pregnancy (Hofmeyr 2017)	Calcium plus other supplements vs placebo		
Effects and safety of periconceptual oral folate supplementation for preventing birth defects (De-Regil 2015)	Folate vs no treatment, placebo or MMN without folate	Not reported	
Multiple micronutrient powders for home (point-of-use) fortification of foods in pregnant women (Suchdev 2015)	MMN powders vs iron and folic acid supplements	Not reported	Not reported
Vitamin A supplements for reducing mother-to-child HIV transmission (Wiysonge 2017)	Vitamin A vs placebo or no treatment	Not a review outcome	
Vitamin A supplementation during pregnancy for maternal and newborn outcomes (McCauley 2015)	Vitamin A alone vs placebo or no treatment		

Figure 17.



Psychosocial interventions targeting other pregnancy conditions	Main comparison	Perinatal death	Preterm birth
Psychosocial interventions for supporting women to stop smoking in pregnancy (Chamberlain 2017)	Any psychosocial intervention vs control		

Figure 18.






















Screening for diabetes	Main comparison	Perinatal death	Preterm birth
Different methods and settings for glucose monitoring for gestational diabetes during pregnancy (Raman 2017)	Telemedicine versus standard care Self-monitoring versus periodic monitoring	 	 Not reported
Different strategies for diagnosing gestational diabetes to improve maternal and infant health (Farrar 2017)	Glucose monomer vs glucose polymer	Not reported	
Screening for gestational diabetes mellitus based on different risk profiles and settings for improving maternal and infant health (Tieu 2017d)	Universal vs risk factor screening Primary care vs secondary care screening	Not reported 	 
Techniques of monitoring blood glucose during pregnancy for women with pre-existing diabetes (Moy 2017)	Self-monitoring vs standard care Self-monitoring vs hospitalisation	 	 
Screening for other pregnancy conditions	Main comparison	Perinatal death	Preterm birth
Antenatal cardiotocography for fetal assessment (Grivell 2015)	Traditional CTG vs no CTG Computerised CTG vs Traditional CTG	 	 
Customised versus population-based growth charts as a screening tool for detecting small for gestational age infants in low-risk pregnant women (Carberry 2014)	No trials		
Fetal and umbilical Doppler ultrasound in normal pregnancy (Alfirevic 2015)	Any Doppler vs no Doppler		
Fetal and umbilical Doppler ultrasound in high-risk pregnancies (Alfirevic 2017b)	Any Doppler vs no Doppler		
Routine blood cultures in the management of pyelonephritis in pregnancy for improving outcomes (Gomi 2015)	No trials		
Use of biochemical tests of placental function for improving pregnancy	Tests of placental function vs standard care		

Figure 18. (Continued)







function for improving pregnancy outcome (Heazell 2015)	standard care		
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Figure 19.

Surgical interventions targeting other pregnancy conditions	Main Comparison	Perinatal death	Preterm birth
Amnioinfusion for chorioamnionitis (Hofmeyr 2016)	Transcervical amnioinfusion vs no amnioinfusion		
External cephalic version (ECV) for breech presentation before term (Hutton 2015)	ECV before term vs ECV at term		

For the sake of brevity, below we do not summarise individual SRs with evidence of unknown harm or benefit for overview outcomes as symbolised by the blue question mark graphic - this task would reiterate what is already apparent in the figures - i.e. there are many interventions for which we have little trial evidence summarised in SRs. Neither do we list the SRs that did not report outcome data - this information may also be found directly in the figures. As above, we direct readers to consider evidence presented in the figures together with the 'Characteristics of included systematic reviews' table (Figure 1).

Systematic reviews with no included trials

The following SRs targeted diabetes or other problems during pregnancy and prespecified an overview priority outcome but included no randomised clinical trials: customised versus population-based growth charts as a screening tool for detecting small for gestational age babies (Carberry 2014); routine blood cultures for the management of pyelonephritis in pregnancy (Gomi 2015); and directed preconception health programs and interventions for women who are overweight or obese (Opray 2015).

Impact on preterm birth

Clear evidence of benefit

One SR reported evidence of GRADE moderate or high quality with clear evidence of benefit (the confidence interval does not cross the line of no effect): lifestyle interventions for the treatment of women with gestational diabetes (Brown 2017a).

Clear evidence of harm

No SR reported clear evidence of harm associated with the intervention; no intervention summarised here increased a woman's chance of PTB.

Clear evidence of no effect or equivalence

The following SRs reported GRADE moderate- or high-quality evidence with narrow confidence intervals crossing the line of no

effect, suggesting the intervention had no effect or equivalence to a comparator: combined health systems interventions to improve antenatal care uptake (Mbuagbaw 2015); diet or exercise or both for preventing excessive weight gain in pregnancy (Muktabhant 2015); vitamin A supplementation (McCauley 2015); antihelminthics during pregnancy (Salam 2015a); psychosocial interventions to stop smoking in pregnancy (Chamberlain 2017); and fetal and umbilical Doppler ultrasound in normal pregnancy (Alfirevic 2015).

Possible benefit

Possible benefit refers to GRADE low-quality evidence with clear benefit (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide confidence intervals crossing the line of no effect. Three SRs reported such evidence for: diet and exercise interventions for preventing gestational diabetes mellitus (Bain 2015); antenatal dietary education and supplementation to increase energy and protein intake (Ota 2015b); and pharmacological interventions for promoting smoking cessation during pregnancy (Coleman 2015).

Possible harm

Possible harm refers to GRADE low-quality evidence with clear harm (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide confidence intervals crossing the line of no effect. One SR reported such evidence for the effects of external cephalic version for breech presentation before term (Hutton 2015).

Unknown harm or benefit

Evidence summarised in many SRs was of GRADE low or very low quality with a wide confidence interval crossing the line of no effect (see Figures 11 to 19).

Impact on perinatal death

Clear evidence of benefit

Two SRs reported evidence of GRADE moderate- or high-quality with clear evidence of benefit (the confidence interval does not

cross the line of no effect): combined health systems interventions to improve antenatal care uptake (Mbuagbaw 2015); and fetal and umbilical Doppler ultrasound in high-risk pregnancy (Alfirevic 2017b).

Clear evidence of harm

No SR reported clear evidence of harm associated with the intervention; no intervention summarised here increased a woman's chance of PD.

Clear evidence of no effect or equivalence

The following SRs reported GRADE moderate- or high-quality evidence with narrow confidence intervals crossing the line of no effect, suggesting the intervention had no effect or equivalence to a comparator: combined health systems interventions to improve antenatal care uptake (Mbuagbaw 2015); vitamin A supplementation (McCauley 2015); and antihelminthics during pregnancy (Salam 2015a).

Possible benefit

Possible benefit refers to GRADE low-quality evidence with clear benefit (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide confidence intervals crossing the line of no effect. One SR reported such evidence for computerised cardiotocography in the SR Antenatal cardiotocography for fetal assessment (Grivell 2015).

Possible harm

Possible harm refers to GRADE low-quality evidence with clear harm (the confidence interval does not cross the line of no effect) or GRADE moderate- or high-quality evidence with wide confidence intervals crossing the line of no effect. No SR reported such evidence.

Unknown harm or benefit

Evidence summarised in many SRs was of GRADE low or very low quality with a wide confidence interval crossing the line of no effect (see Figures 11 to 19).

Network meta-analysis

We found no eligible Cochrane Reviews with network meta-analyses (NMA) for inclusion in this overview. If in future updates of the overview we identify relevant reviews, we will apply the methodologies outlined in our protocol to pilot GRADE for important network comparisons and outcomes and to present key network results visually.

Interactive 'Summary of findings' tables

We did not pilot an interactive 'Summary of findings' table for this version of the overview, but we aim to pilot ideas for presenting layered data tables in online versions of the overview for the next update.

DISCUSSION

There is little consensus or guidance on the appropriate rationale, methodology or outputs for overviews of systematic reviews (SRs) (Ballard 2017; Pieper 2012; Pollock 2016). We have summarised the evidence from relevant Cochrane Reviews using graphic icons to communicate both our confidence in, and the relative effectiveness

of diverse interventions with possible impact on preterm birth (PTB). We stress that preterm birth rates reported in included SRs are likely to be a mix of provider-initiated and spontaneous PTB. Perinatal deaths include both stillbirth and neonatal deaths.

Summary of main results

We found 36 SRs evaluating interventions with PTB prevention as the main focus. Most reviews evaluated interventions related to health systems or screening, or tested the impact of medicines and vitamin supplements.

Four reviews reported clear evidence of benefit, with three relevant to a general pregnant population: midwife-led continuity models of care versus other models of care for all women (Sandall 2016); zinc supplementation for pregnant women without systemic illness (Ota 2015a); and antenatal lower genital tract infection screening for women without signs of preterm labour (Sangkomkhang 2015). Cervical cerclage was the only surgical intervention found to be of benefit, and is relevant only for women with high risk of PTB and singleton pregnancy (Alfirevic 2017a).

It is worth noting that only midwife-led continuity models of care for all pregnant women (Sandall 2016) showed both reduction in PTB and improvement in perinatal survival. The review authors called for further investigation into how a model of care might prevent PTB. They stressed that this is a complex intervention that requires theoretical modelling of the relationships between processes and outcomes in the trial. Authors cautioned against generalising results to a population with health complications and noted that no included trial took place in a resource-constrained setting.

Authors of the zinc supplementation review qualified their finding of benefit to say that most women in included trials were of low income and may have had poor nutrition (Ota 2015a). Infection screening programs were relevant to a general population of pregnant women (Sangkomkhang 2015), whilst the benefit of cervical cerclage was only relevant to women at high risk of PTB and singleton pregnancy (Alfirevic 2017a).

No included reviews reported clear evidence of harm, which is very reassuring.

Reviews targeting diabetes or other pregnancy conditions

Forty-seven included reviews described various interventions targeting diabetes and other conditions in pregnancy. Each specified or reported a PTB outcome, but PTB prevention was not their primary focus. The review by Brown 2017a reported clear benefit of lifestyle interventions for women with gestational diabetes; lifestyle interventions usually involved healthy eating, exercise and self-monitoring of glucose concentrations. Review authors cautioned against generalising this result to women in low- or middle-income countries.

No review targeting diabetes or other health problems during pregnancy reported benefit for both PTB and perinatal death outcomes.

Combined community and health systems interventions to improve antenatal care coverage reduced pregnant women's chances of experiencing perinatal death but had no clear impact on PTB rates (Mbuagbaw 2015). However, this review finding is from

a single trial investigating augmented care for African-American women with multiple pregnancy risks (Klerman 2001).

Fetal and umbilical Doppler for high-risk pregnant women also showed a reduction in perinatal death but no clear impact on PTB rates (Alfirevic 2017b). As most trials did not have a clearly-defined management protocol that followed abnormal Doppler test, it is difficult to be certain which aspect of subsequent care is the most important – the obvious candidates are better timing of delivery, more appropriate mode of labour (vaginal versus caesarean), and better targeting of antenatal corticosteroids.

Reassuringly, no included reviews reported clear evidence of harm.

Overall completeness and applicability of evidence

One of our main aims was to highlight strategies to prevent spontaneous preterm birth (sPTB), but none of the included Cochrane Reviews reported sPTB as an outcome distinct from provider-initiated PTB. Despite their focus on PTB prevention, none of the 36 relevant Cochrane Reviews referenced the Global Alliance to Prevent Prematurity and Stillbirth (GAPPS) classification system for PTB (Villar 2012). We would encourage researchers to try to apply these distinctions to enable more nuanced analysis of PTB-related interventions.

What is missing from the overview?

We identified 96 SRs that may have been eligible during title assessment but were out-of-date (these are listed in Appendix 2). We regret that we were unable to look further into these SRs.

Our peer reviewers noted that the overview misses out three widely-used interventions to prevent PTB in high-risk pregnant women: cervical pessary, cervical length screening and vaginal progesterone. Certainly there are other interventions that are absent because they are not the subject of a Cochrane Review. For these three specific interventions, we direct readers to several other SRs. Saccone 2017a and Saccone 2017b reviewed trials of cervical pessary for women with singleton and multiple pregnancy and short cervix; pessary did not reduce the rate of sPTB or improve outcomes for the neonate. Berghella 2017 found that cervical length screening reduced women's risk of preterm before 37 weeks' gestation, but led to no improvements in other outcomes for women or neonates. Romero 2018 reported that vaginal progesterone reduced risk of preterm births < 33 weeks' gestation for women with singleton pregnancy and a short cervix mid-trimester; progesterone also improved outcomes for the neonate. We also look forward to the EPPIC collaboration's individual participant data (IPD) meta-analysis of progestogens (Stewart 2017), and to an updated Cochrane Review on progestogens for singleton pregnancy (Dodd and colleagues, forthcoming). A network meta-analysis comparing progesterone, cerclage and cervical pessary was also relevant; Jarde 2017a found that progesterone (natural and 17-OHPC were both included here) was the most effective for preventing PTB in singleton pregnancy. No intervention was effective to prevent PTB in twin pregnancy (Jarde 2017b).

Quality of the evidence

We did not conduct formal evaluation of the risks of bias in included SRs with either the AMSTAR or ROBIS tool, due to our confidence in Cochrane Review methodology. All Cochrane Review authors

must conduct exhaustive searches for published and unpublished studies; an Information Specialist checks review search results, and editorial topic experts check review authors' eligibility decisions. All review authors apply Cochrane's 'Risk of bias' tool to included trials and report transparent results in tables (Higgins 2011a). From 2016, all Cochrane review authors also assess the quality of review effect estimates using GRADE criteria and report results in 'Summary of findings' tables (Higgins 2018). Cochrane Review methodology alerts readers to potential bias in pooled effect estimates. Our overview further classified review effect estimates to assign a graphic icon and communicate evidence quality (see Figure 2).

To ensure the credible interpretation of SR estimates researchers should consider whether meta-analyses have adequate statistical power (e.g. Schünemann 2013). GRADE criteria consider the imprecision of a pooled estimate, which is related to power, but no SR included in this overview formally reported whether a meta-analysis had adequate data to draw meaningful conclusions. Overview authors may provide such context to improve interpretation of SR evidence.

We also question how far an overview should attempt to interpret SR results in terms of practice recommendations. If individual SRs should not make clinical recommendations (see Higgins 2011b, section 12.7.4 "Common errors in reaching conclusions"), then neither should overviews of reviews, unless overview authors have embarked upon a formal process to vet all available evidence and produce fully-contextualised clinical guidance.

An example of an evidence synthesis project that can endorse specific interventions for clinical practice is a SR of relevant guidelines. Medley 2018 reviewed recommendations in 49 clinical guidelines to identify international consensus for specific interventions to prevent or manage PTB.

We draw attention to a lack of a good evidence base for women with multiple pregnancy. We classified all evidence relevant to multiple pregnancy to be of unknown harm or benefit, and of low- or very-low quality, with the exception of intramuscular progesterone (which may be harmful for women, see Dodd 2017 and Figure 6). We currently have no credible options to offer this population, and this situation must change. Clinicians, researchers and funders must address the lack of high-quality evidence for interventions relevant to women at high risk of preterm birth due to multiple pregnancy.

Potential biases in the overview process

We followed standard methods to minimise bias. Two overview authors independently conducted eligibility assessment and data extraction, with resolution of conflicts through consultation with a third overview author. None of the overview authors appraised his or her own SR for inclusion; likewise, overview authors did not assess outcome data or assign graphic icons to pooled effect estimates from their own SRs.

We changed our methods from those presented in our protocol to accommodate the large number of included SRs and to communicate the evidence we found. Such deviation from protocol may be considered a source of bias.

Agreements and disagreements with other studies or reviews

We found three other publications that assessed SRs relevant to PTB prevention. We found no study that summarised randomised trial evidence for interventions according to a classification scheme similar to ours.

[Piso 2014](#) summarised 56 Cochrane Reviews of interventions to reduce PTB at less than 37 weeks' gestation. Authors found three interventions that increased preterm birth: metronidazole for pregnant women with asymptomatic trichomoniasis ([Gülmezoglu 2011](#)); oestrogen supplementation to prevent miscarriage; and vitamin C ([Bamigboye 2003](#)). The metronidazole and oestrogen Cochrane Reviews are out of date and are not included in this overview. The result for vitamin C in an updated review included here no longer shows harm but rather shows equivalence of vitamin C with the comparator ([Rumbold 2015a](#), see also [Figure 7](#)). [Piso](#) cites 12 additional reviews reporting benefit to women, including interventions such as smoking cessation programs and antenatal genital tract infection screening. Our overview does not include the same set of Cochrane Reviews as [Piso](#) because some of these Cochrane Reviews are now out-of-date.

[Nijman 2016](#) reviewed interventions to diagnose and treat threatened PTB. The authors found that cervical length measurement combined with fetal fibronectin was the most effective way to identify women who may labour within seven days; corticosteroids improved neonatal outcome; tocolytic therapy should be restricted to 48 hours' use; maintenance tocolysis did not work; magnesium sulphate improved neurological outcomes for the baby; and erythromycin improved neonatal outcomes when women had preterm prelabour rupture of membranes.

[Lucaroni 2018](#) summarised SRs of biomarkers to predict spontaneous PTB and concluded that cervical fetal fibronectin, alpha fetoprotein, C-reactive protein and interleukin 6 each had good diagnostic accuracy to identify women at risk of spontaneous PTB.

One of the important challenges we faced was the appropriate use of user-friendly language bearing in mind that a significant proportion of the readership may not have English as their first language. [Jones 2012](#) categorised interventions for pain during labour with “What works”; “What may work”; and “Insufficient evidence.” For [Ciapponi 2017](#), an “effective” intervention had GRADE moderate- or high-certainty evidence showing benefit for a key outcome and no negative effects via other outcomes; similarly, for [Shepherd 2017](#), “effective” meant GRADE high-quality evidence. [Farquhar 2018](#) classified interventions as effective, promising, ineffective, and possibly ineffective or no conclusions, possible due to lack of evidence.

We aimed to create a similarly straight-forward classification scheme able to communicate both review conclusions about an intervention and GRADE criteria and decision-making. We also wanted readers to be able to consider multiple reviews together in the same figure, without encouraging the direct comparison of relative risk estimates across different SRs. To this end, we created the graphic icons to communicate the effects of interventions and our confidence in the estimates. To improve the usefulness of overviews, researchers should agree standard language and

investigate new ways to communicate complex SR evidence via graphic or interactive media.

AUTHORS' CONCLUSIONS

Implications for practice

No clear evidence of harm from any of the included intervention is reassuring, but the possibility that as many as four may be harmful to women and neonates is a concern. For anyone contemplating their implementation, a careful risk/benefit assessment should be made depending on the setting and available alternative. Reduced antenatal visits may have increased women's risk of perinatal death in settings where pregnant women already received limited antenatal care ([Dowswell 2015](#)). Likewise, women's risk of preterm birth (PTB) may have increased with the administration of: intramuscular progesterone for women with multiple pregnancy ([Dodd 2017](#)); vitamin D, calcium and other supplements for women without pre-existing conditions ([De-Regil 2016](#)); and external cephalic version for women with breech presentation before term ([Hutton 2015](#)).

For health systems where the four effective interventions listed in the discussion are not part of clinical care, a rationale for their exclusion should be made available to users. The forums in which such decisions can be justified may not exist, but this can hardly be an excuse today when access to some form of social media is almost universal. The overview summarises no evidence for cervical pessary, cervical length assessment and vaginal progesterone in singleton pregnancy because these Cochrane Reviews were not current. Readers should be aware that these are active areas for PTB research.

The graphic icons assigned to systematic review effect estimates do not constitute clinical guidance or an endorsement of specific interventions for pregnant women. Readers should refer to full systematic reviews for important details on clinical efficacy, target populations, and the geographic setting of clinical trials. Pregnant women and their healthcare providers must carefully consider whether specific strategies to prevent PTB will be of benefit for individual women, or for specific populations of women.

Implications for research

The difficulty of maintaining timeliness of live systematic reviews in such an active research field as PTB prevention cannot be overstated. Researchers associated with Cochrane have proposed strategies to streamline review production in a series of papers ([Elliott 2017](#)). Limiting meta-analyses to priority outcome domains and measures may also make regular updates more feasible and facilitate easier and more accurate data pooling ([van't Hooft 2016](#)). Future updates of Cochrane Reviews should apply the Global Alliance to Prevent Prematurity and Stillbirth (GAPPS) classification system or another to enable more nuanced reporting and analysis of PTB ([Villar 2012](#)).

Formal consensus work is needed to establish standardised language for overviews and to draw limits for their interpretation of review evidence and its (un)certainly for benefit or harm.

For effective interventions, the underlying mechanism of action on PTB or perinatal death was not always clear. Whilst this should not be an excuse for not implementing interventions demonstrating clear benefit, researchers must be encouraged to find the links

that would enable further improvements and open other areas of scientific enquiry.

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ADDITIONAL TABLES
Table 1. Characteristics of included systematic reviews

Review ID and title	Search date (CPC register)	No. included trials	No. included participants	Overview outcomes reported	Review authors' conclusions (quoted directly from published Cochrane Review)
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Interventions during pregnancy to prevent preterm birth: an overview of Cochrane systematic reviews (Review)

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Table 1. Characteristics of included systematic reviews *(Continued)*

			Participants (women)		
Alfirevic 2015	28 February 2015	5	14,185	<ol style="list-style-type: none"> 1. Perinatal death with anomalies 2. Any death after randomisation 3. Neonatal death (to 28 days) 4. Stillbirth 5. Potentially preventable perinatal death 6. Preterm birth < 37 weeks 7. Gestational age at birth 	Existing evidence does not provide conclusive evidence that the use of routine umbilical artery Doppler ultrasound, or combination of umbilical and uterine artery Doppler ultrasound in low-risk or unselected populations benefits either mother or baby. Future studies should be designed to address small changes in perinatal outcome, and should focus on potentially preventable deaths.
Alfirevic 2017a	30 June 2016	15	3490	<ol style="list-style-type: none"> 1. Perinatal death 2. Stillbirth 3. Neonatal death 4. Preterm birth < 37, < 34 and < 28 weeks 	Cervical cerclage reduces the risk of preterm birth in women at high risk of preterm birth and probably reduces risk of perinatal deaths. There was no evidence of any differential effect of cerclage based on previous obstetric history or short cervix indications, but data were limited for all clinical groups. The question of whether cerclage is more or less effective than other preventative treatments, particularly vaginal progesterone, remains unanswered.
Alfirevic 2017b	31 March 2017	19	10,667	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm labour 	Current evidence suggests that the use of Doppler ultrasound on the umbilical artery in high-risk pregnancies reduces the risk of perinatal deaths and may result in fewer obstetric interventions. The results should be interpreted with caution, as the evidence is not of high quality. Serial monitoring of Doppler changes in duc-

Table 1. Characteristics of included systematic reviews (Continued)

				< 37 weeks	tus venosus may be beneficial, but more studies of high quality with follow-up including neurological development are needed for evidence to be conclusive.
Bain 2014	2 June 2014	8	1181	1. Perinatal death 2. Still-birth 3. Neonatal death 4. Preterm birth 5. Gestational age at birth	Based on eight included trials, of moderate quality overall, no firm conclusions about optimal interventions for managing asthma in pregnancy can be made. Five trials assessing pharmacological interventions did not provide clear evidence of benefits or harms to support or refute current practice.
Bain 2015	11 February 2014	13	4983	1. Perinatal death 2. Still-birth 3. Neonatal death 4. Preterm birth 5. Gestational age at birth	Based on the data currently available, conclusive evidence is not available to guide practice.
Boelig 2016	20 December 2015	25	2052 across 18 comparisons	1. Still-birth and neonatal death 2. Preterm birth	On the basis of this review, there is little high-quality and consistent evidence supporting any one intervention, which should be taken into account when making management decisions.
Bricker 2015a	31 May 2015	13	34,980	1. Perinatal death 2. Still-birth 3. Neonatal death 4. Preterm birth 5. Gestational age at birth	Based on existing evidence, routine late pregnancy ultrasound in low-risk or unselected populations does not confer benefit on mother or baby. There was no difference in the primary outcomes of perinatal mortality, preterm birth less than 37 weeks, caesarean section rates, and induction of labour rates if ultrasound in late pregnancy was performed routinely versus not performed routinely.

Table 1. Characteristics of included systematic reviews (Continued)

Bricker 2015b	15 June 2015	0	0	None	There is no robust evidence from randomised trials to indicate whether specialised diets or nutritional advice for women with multiple pregnancies do more good than harm. There is a clear need to undertake a randomised controlled trial.
Nutritional advice for improving outcomes in multiple pregnancies.					
Brown 2016	7 April 2016	2	142	1. Preterm birth	There are insufficient data to evaluate the effect of myo-inositol for the treatment of gestational diabetes, with no data to examine the majority of outcomes in this review. There do not appear to be any benefits for the infant associated with exposure to myo-inositol
Dietary supplementation with myo-inositol in women during pregnancy for treating gestational diabetes.					
Brown 2017a	14 May 2016	15	4501	1. Perinatal death 2. Preterm birth < 37 and < 32 weeks	Lifestyle interventions are the primary therapeutic strategy for women with GDM. Women receiving lifestyle interventions were less likely to have postnatal depression and were more likely to achieve postpartum weight goals. Exposure to lifestyle interventions was associated with a decreased risk of the baby being born LGA and decreased neonatal adiposity. Long-term maternal and childhood/adulthood outcomes were poorly reported. The value of lifestyle interventions in low- and middle-income countries or for different ethnicities remains unclear. The longer-term benefits or harms of lifestyle interventions remains unclear due to limited reporting.
Lifestyle interventions for the treatment of women with gestational diabetes.					
Brown 2017b	27 August 2016	11	638	1. Perinatal death 2. Preterm birth < 37 and < 32 weeks	Short- and long-term outcomes of interest for this review were poorly reported. Current evidence is confounded by the large variety of exercise interventions. There was insufficient high-quality evidence to be able to determine any differences between exercise and control groups for our outcomes of interest.
Exercise for pregnant women with gestational diabetes for improving maternal and fetal outcomes.					
Brown 2017c	14 May 2016	11	1487	None	There were insufficient data comparing oral anti-diabetic pharmacological therapies with placebo/standard care (lifestyle advice) to inform clinical practice. There was insufficient high-quality evidence to be able to draw any meaningful conclusions as to the benefits of one oral anti-diabetic pharmacological therapy over another due to limited reporting of data for the primary and secondary outcomes in this review. Short- and long-term clinical outcomes for this review were inadequately reported or not reported. Current choice of oral anti-diabetic pharmacological therapy appears to be based on clinical preference, availability and national clinical practice guidelines.
Oral anti-diabetic pharmacological therapies for the treatment of women with gestational diabetes.					
Buppasiri 2015	30th September 2014	25	18587	1. Perinatal death 2. Still-birth or	This review indicates that there are no clear additional benefits to calcium supplementation in prevention of preterm birth or low infant birthweight.
Calcium supplementation (other than for preventing or treating hypertension) for improv-					

Table 1. Characteristics of included systematic reviews (Continued)

ing pregnancy and infant outcomes.				fetal death 3. Preterm birth	
Carberry 2014	12 March 2014	0	0	None	There is no randomised trial evidence currently available. Further randomised trials are required to accurately assess whether the improvement in detection shown is secondary to customised charts alone or an effect of the policy change.
Customised versus population-based growth charts as a screening tool for detecting small for gestational age infants in low-risk pregnant women.					
Catling 2015	31 October 2014	4	2350	1. Perinatal death 2. Preterm birth 3. Gestational age at birth	Available evidence suggests that group antenatal care is acceptable to women and is associated with no adverse outcomes for them or for their babies. No differences in the rate of preterm birth were reported when women received group antenatal care. This review is limited because of the small numbers of studies and women, and because one study contributed 42% of the women. Most of the analyses are based on a single study. Additional research is required to determine whether group antenatal care is associated with significant benefit in terms of preterm birth or birthweight.
Group versus conventional antenatal care for women.					
Chamberlain 2017	13 November 2015	102	28,000	1. Perinatal death 2. Preterm birth < 37 weeks	Psychosocial interventions to support women to stop smoking in pregnancy can increase the proportion of women who stop smoking in late pregnancy and the proportion of infants born low birthweight.
Psychosocial interventions for supporting women to stop smoking in pregnancy.					
Cluver 2017	26 June 2017	15	1754	1. Perinatal death 2. Preterm birth	Treatment with antibacterial agents achieves microbiological cure from <i>C. trachomatis</i> infection during pregnancy. There was no apparent difference between assessed agents (amoxicillin, erythromycin, clindamycin, azithromycin) in terms of efficacy (microbiological cure and repeat infection) and pregnancy complications (preterm birth, preterm rupture of membranes, low birthweight). Azithromycin and clindamycin appear to result in fewer side effects than erythromycin.
Interventions for treating genital Chlamydia trachomatis infection in pregnancy.					
Coleman 2015	11 July 2015	9	2210	1. Stillbirth 2. Neonatal death 3. Preterm birth	There is no evidence that NRT used for smoking cessation in pregnancy has either positive or negative impacts on birth outcomes.
Pharmacological interventions for promoting smoking cessation during pregnancy.					
da Silva Lopes 2017	30 May 2016	6	636	1. Preterm birth	The evidence to date is insufficient to inform a policy of routine bed rest in hospital or at home for women with a multiple pregnancy.
Bed rest with and without hospitalisation in multiple pregnancy for im-					

Table 1. Characteristics of included systematic reviews (Continued)

<p>proving perinatal outcomes.</p> <p>Davey 2015</p> <p>Risk-scoring systems for predicting preterm birth with the aim of reducing associated adverse outcomes.</p>	30 June 2015	0	0	None	The role of risk-scoring systems in the prevention of preterm birth is unknown.
<p>De-Regil 2015</p> <p>Effects and safety of periconceptional oral folate supplementation for preventing birth defects.</p>	31 August 2015	5	7391	<ol style="list-style-type: none"> 1. Still-birth 2. Neonatal death 3. Preterm birth 	Folic acid, alone or in combination with vitamins and minerals, prevents neural tube defects, but does not have a clear effect on other birth defects.
<p>De-Regil 2016</p> <p>Vitamin D supplementation for women during pregnancy.</p>	23 February 2015	15	2388	<ol style="list-style-type: none"> 1. Still-birth 2. Neonatal death 3. Preterm birth 	New studies have provided more evidence on the effects of supplementing pregnant women with vitamin D alone or with calcium on pregnancy outcomes. Supplementing pregnant women with vitamin D in a single or continued dose increases serum 25-hydroxyvitamin D at term and may reduce the risk of pre-eclampsia, low birthweight and preterm birth. However, when vitamin D and calcium are combined, the risk of preterm birth is increased. The clinical significance of the increased serum 25-hydroxyvitamin D concentrations is still unclear. In light of this, these results need to be interpreted with caution. Data on adverse effects were lacking in all studies.
<p>Dodd 2015a</p> <p>Reduction of the number of fetuses for women with a multiple pregnancy.</p>	31 July 2015	0	0	None	We found no available data from randomised trials to inform the risks and benefits of pregnancy reduction procedures for women with a multiple pregnancy. While randomised controlled trials will provide the most reliable evidence about the risks and benefits of fetal reduction procedures, reduction in the number of fetuses by selective termination may not be acceptable to women, particularly couples with a past history of infertility. The acceptability of this option, and willingness to undergo randomisation will depend on the couple's social background and beliefs, and consequently, recruitment to such a trial may prove exceptionally difficult.
<p>Dodd 2015b</p> <p>Specialised antenatal clinics for women with a multiple pregnancy for improving maternal and infant outcomes.</p>	31 May 2015	1	162	<ol style="list-style-type: none"> 1. Perinatal death 2. Still-birth 3. Neonatal death 	There is currently limited information available from randomised controlled trials to assess the role of 'specialised' antenatal clinics for women with a multiple pregnancy compared with 'standard' antenatal care in improving maternal and infant health outcomes.

Table 1. Characteristics of included systematic reviews (Continued)

Dodd 2017	1 November 2016	17	4773	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth < 37, < 34 and < 28 weeks 	Overall, for women with a multiple pregnancy, treatment with progesterone (either intramuscular or vaginal) does not appear to reduce the likelihood of preterm birth or improve outcomes for babies.
Dowswell 2015	23 March 2015	7	60,000+	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth 	In settings with limited resources where the number of visits is already low, reduced visits programmes of antenatal care are associated with an increase in perinatal mortality compared to standard care, although admission to neonatal intensive care may be reduced. Women prefer the standard visits schedule. Where the standard number of visits is low, visits should not be reduced without close monitoring of fetal and neonatal outcome.
Farrar 2016	31 March 2016	5	153	<ol style="list-style-type: none"> 1. Perinatal death 2. Gestational age at birth 3. Preterm birth 	There is no evidence to support the use of one particular form of insulin administration over another for pregnant women with diabetes. There are only a small number of trials appropriate for meta-analysis, a small number of women included and questionable generalisability of the trial population.
Farrar 2017	9 January 2017	7	1420	<ol style="list-style-type: none"> 1. Gestational age at birth 	There is insufficient evidence to suggest which strategy is best for diagnosing GDM.
Gomi 2015	31 December 2014	0	0	None	There are no large-scale randomised controlled trials to assess outcomes in the management of pyelonephritis in pregnancy with or without blood cultures.
Grivell 2015	26 June 2015	6	2105	<ol style="list-style-type: none"> 1. Perinatal death 2. Any potentially preventable perinatal death 	There is no clear evidence that antenatal CTG improves perinatal outcome, but further studies focusing on the use of computerised CTG in specific populations of women with increased risk of complications are warranted.

Table 1. Characteristics of included systematic reviews *(Continued)*

				3. GA at birth	
Haider 2017	11 March 2015	19	138,538	1. Perinatal death 2. Preterm birth	Our findings support the effect of MMN supplements with iron and folic acid in improving some birth outcomes. Overall, pregnant women who received MMN supplementation had fewer low birthweight babies and small-for-gestational-age babies. The findings, consistently observed in several systematic evaluations of evidence, provide a basis to guide the replacement of iron and folic acid with MMN supplements containing iron and folic acid for pregnant women in low and middle-income countries where MMN deficiencies are common among women of reproductive age. Efforts could focus on the integration of this intervention in maternal nutrition and antenatal care programs in low- and middle-income countries.
Han 2017	8 March 2016	19	1398	1. Perinatal death 2. Preterm birth	Evidence from 19 trials assessing different types of dietary advice for women with GDM suggests no clear differences for primary outcomes and secondary outcomes assessed using GRADE, except for a possible reduction in caesarean section for women receiving a DASH diet compared with a control diet. Few differences were observed for secondary outcomes.
Harding 2017	14 November 2016	11	2700	1. Perinatal death 2. Preterm birth	There were insufficient data to reach any meaningful conclusions on the benefits and harms of routine iodine supplementation in women before, during or after pregnancy.
Heazell 2015	31 July 2015	3	740	1. Perinatal death (still-birth or neonatal death together) 2. Still-birth 3. Neonatal death 4. Preterm birth	There is insufficient evidence to support the use of biochemical tests of placental function to reduce perinatal mortality or increase identification of small-for-gestational-age infants. However, we were only able to include data from two studies that measured oestrogens and hPL. The quality of the evidence was low or very low.
Hofmeyr 2016	6 July 2016	1	34	None	There is insufficient evidence to fully evaluate the effectiveness of using transcervical amnioinfusion for chorioamnionitis and to assess the safety of this intervention or women's satisfaction. We did not identify any trials that used transabdominal amnioinfusion.

Table 1. Characteristics of included systematic reviews (Continued)

<p>Hofmeyr 2017</p> <p>Calcium supplementation commencing before or early in pregnancy, or food fortification with calcium, for preventing hypertensive disorders of pregnancy.</p>	10 August 2017	1	60	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth 	<p>Calcium supplementation (≥ 1 g/day) is associated with a significant reduction in the risk of pre-eclampsia, particularly for women with low-calcium diets. The treatment effect may be overestimated due to small-study effects or publication bias. It also reduces preterm birth and the occurrence of the composite outcome 'maternal death or serious morbidity'. We considered these benefits to outweigh the increased risk of HELLP syndrome, which was small in absolute numbers. The World Health Organization recommends calcium 1.5 g to 2 g daily for pregnant women with low dietary calcium intake.</p>
<p>Hutton 2015</p> <p>External cephalic version for breech presentation before term.</p>	31 March 2015	5	2187	<ol style="list-style-type: none"> 1. Perinatal death 2. Stillbirth 3. Preterm labour 4. Stillbirth or neonatal death < 7 days 	<p>Compared with no ECV attempt, ECV commenced before term reduces non-cephalic presentation at birth. Compared with ECV at term, beginning ECV at between 34 to 35 weeks may have some benefit in terms of decreasing the rate of non-cephalic presentation, and risk of vaginal breech birth. However, early ECV may increase risk of late preterm birth, and it is important that any future research reports infant morbidity outcomes. Results of the review suggest that there is a need for careful discussion with women about the timing of the ECV procedure so that they can make informed decisions.</p>
<p>Ihezor-Ejiofor 2017</p> <p>Treating periodontal disease for preventing adverse birth outcomes in pregnant women</p>	7 October 2016	15	7161	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth < 37 weeks 	<p>It is not clear if periodontal treatment during pregnancy has an impact on preterm birth (low-quality evidence). There is low-quality evidence that periodontal treatment may reduce low birth weight (< 2500 g), however, our confidence in the effect estimate is limited</p>
<p>Jahanfar 2014</p> <p>Interventions for preventing or reducing domestic violence against pregnant women.</p>	31 July 2014	10	3417	<ol style="list-style-type: none"> 1. Preterm birth 	<p>There is insufficient evidence to assess the effectiveness of interventions for domestic violence on pregnancy outcomes. There is a need for high-quality, RCTs with adequate statistical power to determine whether intervention programs prevent or reduce domestic violence episodes during pregnancy, or have any effect on maternal and neonatal mortality and morbidity outcomes.</p>
<p>Jahanfar 2015</p> <p>Effects of restricted caffeine intake by mother on fetal, neonatal and pregnancy outcomes.</p>	16 January 2016	1	1197	<ol style="list-style-type: none"> 1. Preterm birth 	<p>There is insufficient evidence to confirm or refute the effectiveness of caffeine avoidance on birthweight or other pregnancy outcomes. There is a need to conduct high-quality, double-blinded RCTs to determine whether caffeine has any effect on pregnancy outcome.</p>
<p>Low 2016</p> <p>Screening for genital chlamydia infection.</p>	14 February 2016	6	359,078	<p>None. No included RCT measured the effects of screen-</p>	<p>There is an absence of RCT evidence about the effects of chlamydia screening in pregnancy. Future RCTs of chlamydia screening interventions should determine the effects of chlamydia screening in pregnancy, of repeated rounds of screening on the incidence of chlamy-</p>

Table 1. Characteristics of included systematic reviews *(Continued)*

				ing in pregnant women	dia-associated PID and chlamydia reinfection in general and high-risk populations.
Martis 2016	31 January 2016	1	180	1. Gestational age at birth	There is insufficient evidence to guide clinical practice for targets for glycaemic control for women with GDM to minimise adverse effects on maternal and fetal health. Glycaemic target recommendations from international professional organisations for maternal glycaemic control vary widely and are reliant on consensus given the lack of high-quality evidence. Further high-quality trials are needed, and these should compare different glycaemic targets for guiding treatment of women with GDM, assess both short-term and long-term health outcomes for women and their babies, include women's experiences and assess health services costs. Four studies are ongoing.
Matthews 2015	19 January 2015	41	5449	1. Spontaneous abortion 2. Still-birth 3. Perinatal death 4. Preterm birth	Given the high prevalence of nausea and vomiting in early pregnancy, women and health professionals need clear guidance about effective and safe interventions, based on systematically reviewed evidence. There is a lack of high-quality evidence to support any particular intervention.
Mbuagbaw 2015	7 June 2015	34	400,000	1. Perinatal death 2. Preterm labour	Single interventions may improve ANC coverage (at least one visit and four or more visits) and deliveries in health facilities. Combined interventions may improve ANC coverage (at least one visit), reduce perinatal mortality and reduce the occurrence of low birth-weight. The effects of the interventions are unrelated to whether they are community or health system interventions.
McCauley 2015	30 March 2015	19	310,000	1. Perinatal death 2. Still-birth 3. Neonatal mortality 4. Preterm birth	The pooled results of three large trials in Nepal, Ghana and Bangladesh (with over 153,500 women) do not currently suggest a role for antenatal vitamin A supplementation to reduce maternal or perinatal mortality. However, the populations studied were probably different with regard to baseline vitamin A status and there were problems with follow-up of women. There is good evidence that antenatal vitamin A supplementation reduces maternal night blindness, maternal anaemia for women who live in areas where vitamin A deficiency is common or who are HIV-positive. In addition, the available evidence suggests a reduction in maternal infection, but these data are not of a high quality.
Middleton 2016	31 January 2016	3	223	1. Perinatal death 2. Gestational	In a very limited body of evidence, few differences in outcomes were seen between very tight and tight-moderate glycaemic control targets in pregnant women with pre-existing type 1 diabetes, including actual glycaemic control achieved. There is evidence of harm (in-

Table 1. Characteristics of included systematic reviews (Continued)

women with pre-existing diabetes.				age at birth	creased pre-eclampsia, caesareans and birthweights greater than 90th centile) for 'loose' control (FBG above 7 mmol/L). Future trials comparing interventions, rather than glycaemic control targets, may be more feasible. Trials in pregnant women with pre-existing type 2 diabetes are required.
Moy 2017	30 November 2016	10	538	1. Perinatal death 2. Preterm birth	This review found no evidence that any glucose monitoring technique is superior to any other technique among pregnant women with pre-existing type 1 or type 2 diabetes. The evidence base for the effectiveness of monitoring techniques is weak and additional evidence from large well-designed randomised trials is required to inform choices of glucose monitoring techniques.
Muktabhant 2015	5 November 2014	65 (49 with data)	11,444	1. Preterm birth	High-quality evidence indicates that diet or exercise, or both, during pregnancy can reduce the risk of excessive GWG. Other benefits may include a lower risk of caesarean delivery, macrosomia, and neonatal respiratory morbidity, particularly for high-risk women receiving combined diet and exercise interventions. Maternal hypertension may also be reduced. Exercise appears to be an important part of controlling weight gain in pregnancy and more research is needed to establish safe guidelines. Most included studies were carried out in developed countries and it is not clear whether these results are widely applicable to lower-income settings.
O'Neill 2017	17 October 2016	5	554	1. Preterm birth	With limited evidence and no meta-analyses, as each trial looked at a different comparison, no firm conclusions could be made about different insulin types and regimens in pregnant women with pre-existing type 1 or 2 diabetes. Further research is warranted to determine who has an increased risk of adverse pregnancy outcome. This would include larger trials, incorporating adequate randomisation and blinding, and key outcomes that include macrosomia, pregnancy loss, pre-eclampsia, caesarean section, fetal anomalies, and birth trauma.
Opray 2015	31 December 2014	0	0	None	We found no randomised controlled trials that assessed the effect of preconception health programs and interventions in overweight and obese women with the aim of improving pregnancy outcomes. Until the effectiveness of preconception health programs and interventions can be established, no practice recommendations can be made. Further research is required in this area.
Ota 2015a	31 October 2014	21	17,000	1. Preterm birth 2. Still-birth or neonatal death	The evidence for a 14% relative reduction in preterm birth for zinc compared with placebo was primarily represented by trials involving women of low income and this has some relevance in areas of high-perinatal mortality. There was no convincing evidence that zinc supplementation during pregnancy results in other useful and important benefits. Since the preterm associa-

Table 1. Characteristics of included systematic reviews (Continued)

				(re-ported as a single out-come)	tion could well reflect poor nutrition, studies to address ways of improving the overall nutritional status of populations in impoverished areas, rather than focusing on micronutrient and or zinc supplementation in isolation, should be an urgent priority.
				3. Gestational age at birth	
Ota 2015b	31 January 2015	17	9030	1. Still-birth 2. Neonatal death 3. Preterm birth	This review provides encouraging evidence that antenatal nutritional education with the aim of increasing energy and protein intake in the general obstetric population appears to be effective in reducing the risk of preterm birth, low birthweight, increasing head circumference at birth, increasing birthweight among undernourished women, and increasing protein intake. There was no evidence of benefit or adverse effect for any other outcome reported.
Peña-Rosas 2015a	10 January 2015	61 (44 with data)	43,274	1. Neonatal death 2. Preterm birth	Supplementation reduces the risk of maternal anaemia and iron deficiency in pregnancy but the positive effect on other maternal and infant outcomes is less clear. Implementation of iron supplementation recommendations may produce heterogeneous results depending on the populations' background risk for low birthweight and anaemia, as well as the level of adherence to the intervention.
Peña-Rosas 2015b	31 July 2015	27 (21 with data)	5490	1. Neonatal death 2. Preterm birth	This review is the most comprehensive summary of the evidence assessing the benefits and harms of intermittent iron supplementation in pregnant women on haematological and pregnancy outcomes. Findings suggest that intermittent regimens produced similar maternal and infant outcomes as daily supplementation but were associated with fewer side effects and reduced the risk of high levels of Hb in mid and late pregnancy, although the risk of mild anaemia near term was increased. While the quality of the evidence was assessed as low or very low, intermittent may be a feasible alternative to daily iron supplementation among those pregnant women who are not anaemic and have adequate antenatal care.
Phupong 2015	30 June 2015	9	725	None	Further RCTs are needed to fully evaluate the effectiveness of interventions for heartburn in pregnancy. Future research should also address other medications such as histamine 2-receptor antagonists, promotility drugs, proton pump inhibitors, and a raft-forming alginate reflux suppressant in treatment of heartburn in pregnancy. More research is needed on acupuncture and other complimentary therapies as treatments for heartburn in pregnancy. Future research should also evaluate any adverse outcomes, maternal satisfaction with treatment and measure pregnant women's quality of life in relation to the intervention.
Radeva-Petrova 2014	1 June 2014	17	14,481	1. Still-birth	Routine chemoprevention to prevent malaria and its consequences has been extensively tested in RCTs,

Table 1. Characteristics of included systematic reviews *(Continued)*

Drugs for preventing malaria in pregnant women in endemic areas: any drug regimen versus placebo or No treatment.				<ol style="list-style-type: none"> 2. Perinatal death 3. Neonatal and Infant death 4. Preterm birth 	with clinically important benefits on anaemia and parasitaemia in the mother, and on birthweight in infants.
Rafael 2014 Cervical stitch (cerclage) for preventing preterm birth in multiple pregnancy.	30 June 2014	5	1577	<ol style="list-style-type: none"> 1. Perinatal death 2. Stillbirth 3. Neonatal death 4. Preterm birth 5. Gestational age at birth 	This review is based on limited data from five small studies of average to above average quality. For multiple gestations, there is no evidence that cerclage is an effective intervention for preventing preterm births and reducing perinatal deaths or neonatal morbidity.
Raman 2017 Different methods and settings for glucose monitoring for gestational diabetes during pregnancy.	30 September 2016	11	1272	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth 	Evidence from 11 RCTs assessing different methods or settings for glucose monitoring for GDM suggests no clear differences for the primary outcomes or other secondary outcomes assessed in this review.
Rumbold 2015a Vitamin C supplementation in pregnancy.	31 March 2015	21	22,192	<ol style="list-style-type: none"> 1. Perinatal death 2. Stillbirth 3. Neonatal death 4. Preterm birth 5. Gestational age at birth 	The data do not support routine vitamin C supplementation alone or in combination with other supplements for the prevention of fetal or neonatal death, poor fetal growth, preterm birth or pre-eclampsia. Further research is required to elucidate the possible role of vitamin C in the prevention of placental abruption and prelabour rupture of membranes. There was no convincing evidence that vitamin C supplementation alone or in combination with other supplements results in other important benefits or harms.
Rumbold 2015b Vitamin E supplementation in pregnancy.	31 March 2015	29	24,300	<ol style="list-style-type: none"> 1. Stillbirth 2. Neonatal death 3. Perinatal death 	The data do not support routine vitamin E supplementation in combination with other supplements for the prevention of stillbirth, neonatal death, preterm birth, pre-eclampsia, preterm or term PROM or poor fetal growth. Further research is required to elucidate the possible role of vitamin E in the prevention of placental abruption. There was no convincing evidence that vitamin E supplementation in combination with other supplements results in other important benefits or harms.

Table 1. Characteristics of included systematic reviews *(Continued)*

				4. Preterm birth 5. Gestational age at birth	
Salam 2015a	31 January 2015	4	4265	1. Perinatal death 2. Preterm birth	The evidence to date is insufficient to recommend use of antihelminthic for pregnant women after the first trimester of pregnancy. More well-designed, large-scale randomised controlled trials are needed to establish the benefit of antihelminthic treatment during pregnancy.
Salam 2015b	29 January 2015	2	2262	1. Perinatal death 2. Infant death 3. Preterm birth	There is limited evidence (from one small trial at a high risk of bias) on the effectiveness on Hib during pregnancy for improving maternal, neonatal and infant health outcomes. Evidence from one large high-quality trial on the effectiveness of viral influenza vaccine during pregnancy suggests reduced RT-PCR confirmed influenza among women and their babies, suggesting the potential of this strategy for scale up but further evidence from varying contexts is required. Further trials for both Hib and viral influenza vaccines with appropriate study designs and suitable comparison groups are required.
Salam 2015c	31 March 2015	4	1646	None	There were few trials, reporting few clinical outcomes and mostly with unclear trial methodology and inadequate follow-up. There is not enough evidence to detect clinical benefits of vitamin B6 supplementation in pregnancy and/or labour other than one trial suggesting protection against dental decay. Future trials assessing this and other outcomes such as orofacial clefts, cardiovascular malformations, neurological development, preterm birth, pre-eclampsia and adverse events are required.
Sandall 2016	31 May 2015	15	17,674	1. Overall fetal loss and neonatal death 2. Preterm birth	This review suggests that women who received midwife-led continuity models of care were less likely to experience intervention and more likely to be satisfied with their care with at least comparable adverse outcomes for women or their infants than women who received other models of care. Further research is needed to explore findings of fewer preterm births and fewer fetal deaths less than 24 weeks, and all fetal loss/neonatal death associated with midwife-led continuity models of care.
Sangkomkhang 2015	30 November 2014	1	4155	1. Neonatal death 2. Preterm birth	There is evidence from one trial that infection screening and treatment programs for pregnant women before 20 weeks' gestation reduce preterm birth and preterm low birthweight. Infection screening and treatment programs are associated with cost savings when used for the prevention of preterm birth. Future trials should evaluate the effects of different types of infection screening programs.

Table 1. Characteristics of included systematic reviews (Continued)

Review title	Date	Number of studies	Number of participants	Number of outcomes	Description of outcomes
<p>Schneeberger 2015</p> <p>Interventions for preventing recurrent urinary tract infection during pregnancy.</p>	20 May 2015	1	200	1. Miscarriage 2. Preterm birth	A daily dose of nitrofurantoin and close surveillance has not been shown to prevent RUTI compared with close surveillance alone. A significant reduction of ASB was found in women with a high clinic attendance rate and who received nitrofurantoin and close surveillance. There was limited reporting of both primary and secondary outcomes for both women and infants. No conclusions can be drawn regarding the optimal intervention to prevent RUTI in women who are pregnant. Randomised controlled trials comparing different pharmacological and non-pharmacological interventions are necessary to investigate potentially effective interventions to prevent RUTI in women who are pregnant.
<p>Smaill 2015</p> <p>Antibiotics for asymptomatic bacteriuria in pregnancy.</p>	19 March 2015	14	2000	1. Preterm birth	While antibiotic treatment is effective in reducing the risk of pyelonephritis in pregnancy, the estimate of the effect is very uncertain because of the very low quality of the evidence. The reduction in low birthweight and preterm birth with antibiotic treatment is consistent with theories about the role of infection in adverse pregnancy outcomes, but this association should be interpreted with caution given the very poor quality of the included studies.
<p>Smyth 2015</p> <p>Interventions for varicose veins and leg oedema in pregnancy.</p>	31 May 2015	7	326	None	There is moderate-quality evidence to suggest that rutosides appear to help relieve the symptoms of varicose veins in late pregnancy. However, this finding is based on one study (69 women) and there are not enough data presented in the study to assess its safety in pregnancy. Reflexology or water immersion appears to help improve symptoms for women with leg oedema, but again this is based on two small studies (43 and 32 women, respectively).
<p>Sosa 2015</p> <p>Bed rest in singleton pregnancies for preventing preterm birth.</p>	18 December 2014	2	1266	1. Preterm birth	There is no evidence, either supporting or refuting the use of bed rest at home or in hospital, to prevent preterm birth. Although bed rest in hospital or at home is widely used as the first step of treatment, there is no evidence that this practice could be beneficial. Due to the potential adverse effects that bed rest could have on women and their families, and the increased costs for the healthcare system, clinicians should discuss the pros and cons of bed rest to prevent preterm birth. Potential benefits and harms should be discussed with women facing an increased risk of preterm birth. Appropriate research is mandatory. Future trials should evaluate both the effectiveness of bed rest, and the effectiveness of the prescription of bed rest, to prevent preterm birth.
<p>Spencer 2015</p> <p>Screening and subsequent management for thyroid dysfunction pre-pregnancy and dur-</p>	14 July 2015	2	26,408	1. Fetal and neonatal death (reported)	Based on the existing evidence, though universal screening for thyroid dysfunction in pregnancy increases the number of women diagnosed with hypothyroidism who can be subsequently treated, it does not clearly impact (benefit or harm) maternal and infant outcomes. More evidence is needed to assess the benefits or harms of different screening methods for thyroid

Table 1. Characteristics of included systematic reviews (Continued)

ing pregnancy for improving maternal and infant health.				as a single outcome)	dysfunction in pregnancy, on maternal, infant and child health outcomes. Future trials should assess impacts on use of health services and costs, and be adequately powered to evaluate the effects on short- and long-term outcomes.
				2. Miscarriage	
				3. Preterm birth	
				4. Preterm labour	
Suchdev 2015	31 January 2015	2	1172	None	Limited evidence suggests that micronutrient powders for point-of-use fortification of foods have no clear difference as multiple micronutrient supplements on maternal anaemia (very low-quality evidence) and Hb at or near term. There is limited evidence to suggest that women were more likely to adhere to taking tablets than using micronutrient powders.
Multiple micronutrient powders for home (point-of-use) fortification of foods in pregnant women.					
Terplan 2015	January 2015	14	1298	1. Preterm birth	The present evidence suggests that there is no difference in treatment outcomes to address drug use in pregnant women with use of psychosocial interventions, when taken in the presence of other comprehensive care options. However, few studies evaluated obstetrical or neonatal outcomes and rarely did so in a systematic way, making it difficult to assess the effect of psychosocial interventions on these clinically important outcomes. It is important to develop a better evidence base to evaluate psychosocial modalities of treatment in this important population.
Psychosocial interventions for pregnant women in outpatient illicit drug treatment programs compared to other interventions.					
Thinkhamrop 2015	31 July 2014	7	2100	1. Perinatal death 2. Preterm birth 3. Gestational age at birth	Antibiotic prophylaxis did not reduce the risk of preterm prelabour rupture of membranes or preterm delivery (apart from in the subgroup of women with a previous preterm birth who had bacterial vaginosis). Antibiotic prophylaxis given during the second or third trimester of pregnancy reduced the risk of postpartum endometritis, term pregnancy with prelabour rupture of membranes and gonococcal infection when given routinely to all pregnant women. Substantial bias possibly exists in the review's results because of a high rate of loss to follow-up and the small numbers of studies included in each of our analyses. There is also insufficient evidence on possible harmful effects on the baby. Therefore, we conclude that there is not enough evidence to support the use of routine antibiotics during pregnancy to prevent infectious adverse effects on pregnancy outcomes.
Antibiotic prophylaxis during the second and third trimester to reduce adverse pregnancy outcomes and morbidity.					
Tieu 2017a	7 April 2017	0	0	None	The role of interconception care for women with a history of GDM remains unclear. Randomised controlled trials are required evaluating different forms and protocols of interconception care for these women on perinatal and long-term maternal and infant health outcomes, acceptability of such interventions and cost-effectiveness.
Interconception care for women with a history of gestational diabetes for improving maternal and infant outcomes.					

Table 1. Characteristics of included systematic reviews (Continued)

<p>Tieu 2017b</p> <p>Dietary advice interventions in pregnancy for preventing gestational diabetes.</p>	3 January 2016	11	2786	<ol style="list-style-type: none"> 1. Perinatal death 2. Gestational age at birth 	<p>Very low-quality evidence from five trials suggests a possible reduction in GDM risk for women receiving dietary advice versus standard care, and low-quality evidence from four trials suggests no clear difference for women receiving low- versus moderate- to high-GI dietary advice. A possible reduction in pregnancy-induced hypertension for women receiving dietary advice was observed and no clear differences were seen for other reported primary outcomes. There were few outcome data for secondary outcomes. More high-quality evidence is needed to determine the effects of dietary advice interventions in pregnancy.</p>
<p>Tieu 2017c</p> <p>Oral anti-diabetic agents for women with established diabetes/impaired glucose tolerance or previous gestational diabetes planning pregnancy, or pregnant women with pre-existing diabetes.</p>	31 October 2016	3	241	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth < 37 and < 32 weeks 	<p>There are insufficient RCT data to evaluate the use of oral anti-diabetic agents in women with established diabetes, impaired glucose tolerance or previous gestational diabetes who are planning a pregnancy, or in pregnant women with pre-existing diabetes.</p>
<p>Tieu 2017d</p> <p>Screening for gestational diabetes mellitus based on different risk profiles and settings for improving maternal and infant health.</p>	31 January 2017	2	4523	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth 	<p>Further, high-quality randomised controlled trials are needed to assess the value of screening for GDM, which may compare different protocols, guidelines or programmes for screening (based on different risk profiles and settings), with the absence of screening, or with other protocols, guidelines or programmes. There is a need for future trials to be sufficiently powered to detect important differences in short- and long-term maternal and infant outcomes, such as those important outcomes pre-specified in this review. As only a proportion of women will be diagnosed with GDM in these trials, large sample sizes may be required.</p>
<p>Till 2015</p> <p>Incentives for increasing prenatal care use by women in order to improve maternal and neonatal outcomes.</p>	31 January 2015	5	11,935	None	<p>The included studies did not report on this review's main outcomes: preterm birth, small-for-gestational age, or perinatal death. There is limited evidence that incentives may increase utilisation and quality of prenatal care, but may also increase caesarean rate. Overall, there is insufficient evidence to fully evaluate the impact of incentives on prenatal care initiation. There is a need for high-quality RCTs to determine whether incentive program increase prenatal care use and improve maternal and neonatal outcomes. Incentive programs, in particular cash-based programs, as suggested in this review and in several observational studies may improve the frequency and ensure adequate quality of prenatal care.</p>
<p>Urquhart 2017</p> <p>Home uterine monitoring for detecting preterm labour.</p>	30 June 2016	15	6008	<ol style="list-style-type: none"> 1. Perinatal death 2. Preterm birth 	<p>Home uterine monitoring may result in fewer admissions to a neonatal intensive care unit, but in more unscheduled antenatal visits and tocolytic treatment; the level of evidence is generally low to moderate. Important group differences were not evident when we un-</p>

Table 1. Characteristics of included systematic reviews (Continued)

					dertook sensitivity analysis using only trials at low risk of bias. There is no impact on maternal and perinatal outcomes such as perinatal mortality or incidence of preterm birth.
Widmer 2015	31 August 2015	13	1622	1. Preterm birth	A single-dose regimen of antibiotics may be less effective than a short course (four- to seven-day) regimen, but more evidence is needed from large trials measuring important outcomes, such as cure rate. Women with asymptomatic bacteriuria in pregnancy should be treated by the standard regimen of antibiotics until more data become available testing seven-day treatment compared with shorter courses of three- or five-day regimens.
Duration of treatment for asymptomatic bacteriuria during pregnancy.					
Wiysonge 2017	7 September 2017	5	7298	1. Preterm birth	Antepartum or postpartum vitamin A supplementation, or both, probably has little or no effect on mother-to-child transmission of HIV in women living with HIV infection and not on antiretroviral drugs. The intervention has largely been superseded by ART which is widely available and effective in preventing vertical transmission.
Vitamin A supplements for reducing mother-to-child HIV transmission.					
Yamasmit 2015	21 September 2015	6	374	1. Neonatal death 2. Preterm labour 3. Preterm birth	There is insufficient evidence to support or refute the use of prophylactic oral betamimetics for preventing preterm birth in women with a twin pregnancy.
Prophylactic oral betamimetics for reducing preterm birth in women with a twin pregnancy.					
Zhou 2015	31 March 2015	6	390	None	It is unclear from the evidence reviewed whether any of the interventions (oral magnesium, oral calcium, oral vitamin B or oral vitamin C) provide an effective treatment for leg cramps. This is primarily due to outcomes being measured and reported in different, incompatible ways, and design limitations compromising the quality of the evidence (the level of evidence was graded low or very low). This was mainly due to poor study design and trials being too small to address the question satisfactorily.
Interventions for leg cramps in pregnancy.					

ANC: antenatal care
 ART: antiretroviral therapy
 ASB: asymptomatic bacteriuria
 CTG: cardiotocography
 DASH: Dietary Approaches to Stop Hypertension
 ECV: external cephalic version
 FBG: fasting blood glucose
 GA: gestational age
 GDM: gestational diabetes mellitus
 GWG: gestational weight gain
 Hb: haemoglobin
 HELLP: haemolysis, elevated liver enzymes, low platelets
 hpl: human placental lactogen
 LGA: large-for-gestational age
 MMN: multiple-micronutrient
 NRT: nicotine replacement therapy

PROM: prelabour rupture of membranes
 RCT: randomised controlled trial
 RT-PCR: reverse-transcriptase–polymerase-chain-reaction
 RUTI: recurrent urinary tract infections
 UTI: urinary tract infection

Table 2. Excluded systematic reviews

Review ID	Review title	Reason for exclusion
Amaya-Guio 2016	Antibiotic treatment for the sexual partners of women with bacterial vaginosis.	The review does not report results for pregnant women; the intervention does not target health providers.
Balogun 2016a	Vitamin supplementation for preventing miscarriage.	Reviews targets miscarriage in early pregnancy.
Balogun 2016b	Interventions for initiation of breastfeeding.	Overview outcomes of preterm birth and perinatal death are not specified as review outcomes. Review concerned with post-pregnancy outcomes.
Chi 2015	Safety of topical corticosteroids in pregnancy.	All evidence is from cohort studies
Demicheli 2015	Vaccines for women for preventing neonatal tetanus.	The intervention targets the neonate.
Dickinson 2014	Creatine for women in pregnancy for neuroprotection of the fetus.	The intervention targets the neonate.
Frazer 2016	Legislative smoking bans for reducing harms from secondhand smoke exposure, smoking prevalence and tobacco consumption.	Evidence for the effectiveness of smoking bans to impact preterm birth comes from non-randomised study designs.
Grivell 2014	Prenatal versus postnatal repair procedures for spina bifida for improving infant and maternal outcomes.	Intervention targets pregnant women with major fetal abnormality.
Hemmingsen 2017	Dipeptidyl-peptidase (DPP)-4 inhibitors and glucagon-like peptide (GLP)-1 analogues for prevention or delay of type 2 diabetes mellitus and its associated complications in people at increased risk for the development of type 2 diabetes mellitus.	Pregnant women are excluded from many of the trials included in this review.
Heslop 2016	Interventions for men and women with their first episode of genital herpes.	Data are not reported for pregnant women. Pregnancy was an exclusion criteria for several included clinical trials. The intervention does not target healthcare providers.
Jahanfar 2017	Ultrasound for diagnosis of birthweight discordance in twin pregnancies (Protocol).	This is a DTA review and no outcome data relevant to the overview are planned.
Karanth 2015	Desmopressin acetate (DDAVP) for preventing and treating acute bleeds during pregnancy in women with congenital bleeding.	Preterm birth is not a review outcome.
Khattri 2017	Adjunctive systemic antimicrobials for the non-surgical treatment of chronic and aggressive periodontitis (Protocol).	Pregnant women are excluded from this review protocol.

Table 2. Excluded systematic reviews (Continued)

Li 2014	Periodontal therapy for the management of cardiovascular disease in patients with chronic periodontitis.	Pregnant women were not eligible for this review. The intervention does not target healthcare providers.
Li 2016	Chinese herbal medicines for unexplained recurrent miscarriage.	This review is concerned with recurrent miscarriage in early pregnancy.
Liddle 2015	Interventions for preventing and treating low-back and pelvic pain during pregnancy.	Preterm birth and perinatal death are not specified as review outcomes.
Martineau 2016	Vitamin D for the management of asthma.	This review included children and adults. There is no indication that pregnant women were included or excluded from the review.
O'Doherty 2015	Screening women for intimate partner violence in healthcare settings.	This review is not specific to a pregnant population and no pregnancy outcomes are specified.
Okusanya 2016	Prophylactic versus selective blood transfusion for sickle cell disease in pregnancy.	Preterm birth mentioned in the text; only non-randomised data presented in the discussion.
Stock 2016	Immediate versus deferred delivery of the preterm baby with suspected fetal compromise for improving outcomes.	Pregnant women included in this review were expected to deliver preterm.
Tieu 2017f	Preconception care for diabetic women for improving maternal and infant health.	Intervention limited to the preconception period and targeted women who were not pregnant. PTB is a review outcome.
Weston 2016	Monotherapy treatment of epilepsy in pregnancy: congenital malformation outcomes in the child.	This review is concerned with outcomes for the neonate and child.
Whitworth 2015	Ultrasound for fetal assessment in early pregnancy.	This review does not cover the use of ultrasound to identify women at risk of preterm birth; that topic is covered in another review.
Wilkinson 2016	Melatonin for women in pregnancy for neuro-protection of the fetus.	The intervention in this review targets the health of the neonate.

DTA: diagnostic test accuracy

PTB: preterm birth

APPENDICES

Appendix 1. Search strategy

Date Run: 02/11/17 15:23:15.642

#1 preterm near birth* 2798

#2 preterm near lab*r 1164

#3 preterm near delivery 1213

#4 pre-term near birth* 161

#5 pre-term near delivery 89

#6 premature near birth* 2067

#7 pre-term near lab*r 55

#8 premature near delivery 319

#9 premature near lab*r 2029

#10 MeSH descriptor: [Premature Birth] explode all trees 582

#11 MeSH descriptor: [Obstetric Labor, Premature] explode all trees 1329

#12 #1 or #2 or #3 or #4 or #5 or #6 or #7 or #8 or #9 or #10 or #11 5913

Limit to Cochrane reviews and protocols

898 Cochrane reviews or protocols

Appendix 2. Eligible but out of date systematic reviews

1. Abalos E, Duley L, Steyn DW. Antihypertensive drug therapy for mild to moderate hypertension during pregnancy. *Cochrane Database of Systematic Reviews* 2014, Issue 2. [DOI: 10.1002/14651858.CD002252.pub3]
2. Abdel-Aleem H, Shaaban OM, Abdel-Aleem MA. Cervical pessary for preventing preterm birth. *Cochrane Database of Systematic Reviews* 2013, Issue 5. [DOI: 10.1002/14651858.CD007873.pub3]
3. Alexander S, Boulvain M, Ceysens G, Haelterman E, Zhang W-H. Repeat digital cervical assessment in pregnancy for identifying women at risk of preterm labour. *Cochrane Database of Systematic Reviews* 2010, Issue 6. [DOI: 10.1002/14651858.CD005940.pub2]
4. Barrett HL, Dekker NM, Conwell LS, Callaway LK. Probiotics for preventing gestational diabetes. *Cochrane Database of Systematic Reviews* 2014, Issue 2. [DOI: 10.1002/14651858.CD009951.pub2]
5. Bergamaschi DP, Mariath AB, Abbade JF, Grillo LP, Diniz CS, Hinnig PF. Selenium supplementation during pregnancy for improving maternal and newborn outcomes. *Cochrane Database of Systematic Reviews* 2012, Issue 3. [DOI: 10.1002/14651858.CD009673]
6. Berghella V, Baxter JK, Hendrix NW. Cervical assessment by ultrasound for preventing preterm delivery. *Cochrane Database of Systematic Reviews* 2013, Issue 1. [DOI: 1002/14651858.CD007235.pub3]
7. Berghella V, Hayes E, Visintine J, Baxter JK. Fetal fibronectin testing for reducing the risk of preterm birth. *Cochrane Database of Systematic Reviews* 2008, Issue 4. [DOI: 1002/14651858.CD006843.pub2]
8. Brocklehurst P, Gordon A, Heatley E, Milan SJ. Antibiotics for treating bacterial vaginosis in pregnancy. *Cochrane Database of Systematic Reviews* 2013, Issue 1. [DOI: 10.1002/14651858.CD000262.pub4]
9. Carlin AJ, Alfrevic Z, Gyte GM. Interventions for treating peripartum cardiomyopathy to improve outcomes for women and babies. *Cochrane Database of Systematic Reviews* 2010, Issue 9. [DOI: 10.1002/14651858.CD008589.pub2]
10. Churchill D, Beevers GD, Meher S, Rhodes C. Diuretics for preventing pre-eclampsia. *Cochrane Database of Systematic Reviews* 2007, Issue 1. [DOI: 10.1002/14651858.CD004451.pub2]
11. Churchill D, Duley L, Thornton JG, Jones L. Interventionist versus expectant care for severe pre-eclampsia between 24 and 34 weeks' gestation. *Cochrane Database of Systematic Reviews* 2013, Issue 7. [DOI: 1002/14651858.CD003106.pub2]
12. Dennis C-L, Dowswell T. Interventions (other than pharmacological, psychosocial or psychological) for treating antenatal depression. *Cochrane Database of Systematic Reviews* 2013, Issue 7. [DOI: 10.1002/14651858.CD006795.pub3]
13. Dennis C-L, Ross LE, Grigoriadis S. Psychosocial and psychological interventions for treating antenatal depression. *Cochrane Database of Systematic Reviews* 2007, Issue 3. [DOI: 10.1002/14651858.CD006309.pub2]
14. Dodd JM, McLeod A, Windrim RC, Kingdom J. Antithrombotic therapy for improving maternal or infant health outcomes in women considered at risk of placental dysfunction. *Cochrane Database of Systematic Reviews* 2013, Issue 7. [DOI: 10.1002/14651858.CD006780.pub3]
15. Dowswell T, Middleton P, Weeks A. Antenatal day care units versus hospital admission for women with complicated pregnancy. *Cochrane Database of Systematic Reviews* 2009, Issue 4. [DOI: 10.1002/14651858.CD001803.pub2]

16. Duley L, Gülmezoglu AM, Henderson-Smart DJ, Chou D. Magnesium sulphate and other anticonvulsants for women with pre-eclampsia. *Cochrane Database of Systematic Reviews* 2010, Issue 11. [DOI: 1002/14651858.CD000025.pub2]
17. Duley L, Henderson-Smart DJ, Chou D. Magnesium sulphate versus phenytoin for eclampsia. *Cochrane Database of Systematic Reviews* 2010, Issue 10. [DOI: 1002/14651858.CD000128.pub2]
18. Duley L, Henderson-Smart DJ, Meher S. Altered dietary salt for preventing pre-eclampsia, and its complications. *Cochrane Database of Systematic Reviews* 2005, Issue 4. [DOI: 1002/14651858.CD005548]
19. Duley L, Henderson-Smart DJ, Meher S, King JF. Antiplatelet agents for preventing pre-eclampsia and its complications. *Cochrane Database of Systematic Reviews* 2007, Issue 2. [DOI: 1002/14651858.CD004659.pub2]
20. Duley L, Henderson-Smart DJ, Walker GJ, Chou D. Magnesium sulphate versus diazepam for eclampsia. *Cochrane Database of Systematic Reviews* 2010, Issue 12. [DOI: 1002/14651858.CD000127.pub2]
21. Duley L, Matar HE, Almerie MQ, Hall DR. Alternative magnesium sulphate regimens for women with pre-eclampsia and eclampsia. *Cochrane Database of Systematic Reviews* 2010, Issue 8. [DOI: 1002/14651858.CD007388.pub2]
22. Duley L, Meher S, Jones L. Drugs for treatment of very high blood pressure during pregnancy. *Cochrane Database of Systematic Reviews* 2013, Issue 7. [DOI: 1002/14651858.CD001449.pub3]
23. Duley L, Williams J, Henderson-Smart DJ. Plasma volume expansion for treatment of pre-eclampsia. *Cochrane Database of Systematic Reviews* 1999, Issue 4. [DOI: 1002/14651858.CD001805]
24. Eames AJ, Grivell RM, Deussen AR, Hague W, Dodd JM. Metformin for women who are obese during pregnancy for improving maternal and infant outcomes. *Cochrane Database of Systematic Reviews* 2013, Issue 6. [DOI: 1002/14651858.CD010564]
25. Earl R, Crowther CA, Middleton P. Interventions for hyperthyroidism pre-pregnancy and during pregnancy. *Cochrane Database of Systematic Reviews* 2013, Issue 11. [DOI: 1002/14651858.CD008633.pub3]
26. Furber CM, McGowan L, Bower P, Kontopantelis E, Quenby S, Lavender T. Antenatal interventions for reducing weight in obese women for improving pregnancy outcome. *Cochrane Database of Systematic Reviews* 2013, Issue 1. [DOI: 1002/14651858.CD009334.pub2]
27. Gagnon AJ, Sandall J. Individual or group antenatal education for childbirth or parenthood, or both. *Cochrane Database of Systematic Reviews* 2007, Issue 3. [DOI: 1002/14651858.CD002869.pub2]
28. Gamble CL, Ekwaru JP, ter KFO. Insecticide-treated nets for preventing malaria in pregnancy. *Cochrane Database of Systematic Reviews* 2006, Issue 2. [DOI: 1002/14651858.CD003755.pub2]
29. Grivell RM, Wong L, Bhatia V. Regimens of fetal surveillance for impaired fetal growth. *Cochrane Database of Systematic Reviews* 2012, Issue 6. [DOI: 1002/14651858.CD007113.pub3]
30. Guinto VT, De Guia B, Festin MR, Dowswell T. Different antibiotic regimens for treating asymptomatic bacteriuria in pregnancy. *Cochrane Database of Systematic Reviews* 2010, Issue 9. [DOI: 1002/14651858.CD007855.pub2]
31. Gülmezoglu AM, Azhar M. Interventions for trichomoniasis in pregnancy. *Cochrane Database of Systematic Reviews* 2011, Issue 5. [DOI: 1002/14651858.CD000220.pub2]
32. Gurung V, Stokes M, Middleton P, Milan SJ, Hague W, Thornton JG. Interventions for treating cholestasis in pregnancy. *Cochrane Database of Systematic Reviews* 2013, Issue 6. [DOI: 1002/14651858.CD000493.pub2]
33. Han S, Crowther CA, Middleton P. Interventions for pregnant women with hyperglycaemia not meeting gestational diabetes and type 2 diabetes diagnostic criteria. *Cochrane Database of Systematic Reviews* 2012, Issue 1. [DOI: 1002/14651858.CD009037.pub2]
34. Han S, Middleton P, Crowther CA. Exercise for pregnant women for preventing gestational diabetes mellitus. *Cochrane Database of Systematic Reviews* 2012, Issue 7. [DOI: 1002/14651858.CD009021.pub2]
35. Henriquez DD, Roos-Hesselink JW, Schaliq MJ, Klautz RJ, Helmerhorst FM, de Groot CJ. Treatment of valvular heart disease during pregnancy for improving maternal and neonatal outcome. *Cochrane Database of Systematic Reviews* 2011, Issue 5. [DOI: 1002/14651858.CD008128.pub2]
36. Hodnett ED, Fredericks S, Weston J. Support during pregnancy for women at increased risk of low birthweight babies. *Cochrane Database of Systematic Reviews* 2010, Issue 6. [DOI: 1002/14651858.CD000198.pub2]
37. Jefferys AE, Siassakos D, Draycott T, Akande VA, Fox R. Deflation of gastric band balloon in pregnancy for improving outcomes. *Cochrane Database of Systematic Reviews* 2013, Issue 4. [DOI: 1002/14651858.CD010048.pub2]

38. Jepson RG, Williams G, Craig JC. Cranberries for preventing urinary tract infections. *Cochrane Database of Systematic Reviews* 2012, Issue 10. [DOI: 1002/14651858.CD001321.pub5]
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41. Khunpradit S, Tavender E, Lumbiganon P, Laopaiboon M, Wasiak J, Gruen RL. Non-clinical interventions for reducing unnecessary caesarean section. *Cochrane Database of Systematic Reviews* 2011, Issue 6. [DOI: 1002/14651858.CD005528.pub2]
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48. Lindegren ML, Kennedy CE, Bain-Brickley D, Azman H, Creanga AA, Butler LM, et al. Integration of HIV/AIDS services with maternal, neonatal and child health, nutrition, and family planning services. *Cochrane Database of Systematic Reviews* 2012, Issue 9. [DOI: 1002/14651858.CD010119]
49. Lui S, Terplan M, Smith EJ. Psychosocial interventions for women enrolled in alcohol treatment during pregnancy. *Cochrane Database of Systematic Reviews* 2008, Issue 3. [DOI: 1002/14651858.CD006753.pub2]
50. Macdonald G, Bennett C, Higgins JP, Dennis JA. Home visiting for socially disadvantaged mothers. *Cochrane Database of Systematic Reviews* 2010, Issue 10. [DOI: 1002/14651858.CD008784]
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52. Makrides M, Crosby DD, Bain E, Crowther CA. Magnesium supplementation in pregnancy. *Cochrane Database of Systematic Reviews* 2014, Issue 4. [DOI: 1002/14651858.CD000937.pub2]
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55. Martí-Carvajal AJ, Peña-Martí GE, Comunián-Carrasco G, Martí-Peña AJ. Interventions for treating painful sickle cell crisis during pregnancy. *Cochrane Database of Systematic Reviews* 2009, Issue 1. [DOI: 1002/14651858.CD006786.pub2]
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58. Meher S, Duley L. Progesterone for preventing pre-eclampsia and its complications. *Cochrane Database of Systematic Reviews* 2006, Issue 4. [DOI: 1002/14651858.CD006175]

59. Meher S, Duley L. Rest during pregnancy for preventing pre-eclampsia and its complications in women with normal blood pressure. *Cochrane Database of Systematic Reviews* 2006, Issue 2. [DOI: 1002/14651858.CD005939]
60. Meher S, Duley L. Exercise or other physical activity for preventing pre-eclampsia and its complications. *Cochrane Database of Systematic Reviews* 2006, Issue 2. [DOI: 1002/14651858.CD005942]
61. Meher S, Duley L. Garlic for preventing pre-eclampsia and its complications. *Cochrane Database of Systematic Reviews* 2006, Issue 3. [DOI: 1002/14651858.CD006065]
62. Meher S, Duley L. Nitric oxide for preventing pre-eclampsia and its complications. *Cochrane Database of Systematic Reviews* 2007, Issue 2. [DOI: 1002/14651858.CD006490]
63. Miller BJ, Murray L, Beckmann MM, Kent T, Macfarlane B. Dietary supplements for preventing postnatal depression. *Cochrane Database of Systematic Reviews* 2013, Issue 10. [DOI: 1002/14651858.CD009104.pub2]
64. Minozzi S, Amato L, Bellisario C, Ferri M, Davoli M. Maintenance agonist treatments for opiate-dependent pregnant women. *Cochrane Database of Systematic Reviews* 2013, Issue 12. [DOI: 1002/14651858.CD006318.pub3]
65. Mujzinovic F, Alfirevic Z. Technique modifications for reducing the risks from amniocentesis or chorionic villus sampling. *Cochrane Database of Systematic Reviews* 2012, Issue 8. [DOI: 10.1002/14651858.CD008678.pub2]
66. Nabhan AF, Elsedawy MM. Tight control of mild-moderate pre-existing or non-proteinuric gestational hypertension. *Cochrane Database of Systematic Reviews* 2011, Issue 7. [DOI: 1002/14651858.CD006907.pub2]
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68. Orton LC, Omari AA. Drugs for treating uncomplicated malaria in pregnant women. *Cochrane Database of Systematic Reviews* 2008, Issue 4. DOI: 10.1002/14651858.CD004912.pub3]
69. Othman M, Alfirevic Z, Neilson JP. Probiotics for preventing preterm labour. *Cochrane Database of Systematic Reviews* 2007, Issue 1. [DOI: 1002/14651858.CD005941.pub2]
70. Pasricha S-R, De-Regil LM, Garcia-Casal MN, Burford BJ, Gwartz JA, Peña-Rosas JP. Fortification of maize flour with iron for preventing anaemia and iron deficiency in populations. *Cochrane Database of Systematic Reviews* 2012, Issue 11. [DOI: 1002/14651858.CD010187]
71. Quijano CE, Abalos E. Conservative management of symptomatic and/or complicated haemorrhoids in pregnancy and the puerperium. *Cochrane Database of Systematic Reviews* 2005, Issue 3. [DOI: 1002/14651858.CD004077.pub2]
72. Rayment R, Brunskill SJ, Soothill PW, Roberts DJ, Bussell JB, Murphy MF. Antenatal interventions for fetomaternal alloimmune thrombocytopenia. *Cochrane Database of Systematic Reviews* 2011, Issue 5. [DOI: 1002/14651858.CD004226.pub3]
73. Raynes-Greenow CH, Roberts CL, Bell JC, Peat B, Gilbert GL, Parker S. Antibiotics for ureaplasma in the vagina in pregnancy. *Cochrane Database of Systematic Reviews* 2011, Issue 9. [DOI: 1002/14651858.CD003767.pub3]
74. Reid SM, Middleton P, Cossich MC, Crowther CA, Bain E. Interventions for clinical and subclinical hypothyroidism pre-pregnancy and during pregnancy. *Cochrane Database of Systematic Reviews* 2013, Issue 5. [DOI: 1002/14651858.CD007752.pub3]
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76. Rumbold A, Duley L, Crowther CA, Haslam RR. Antioxidants for preventing pre-eclampsia. *Cochrane Database of Systematic Reviews* 2008, Issue 1. [DOI: 1002/14651858.CD004227.pub3]
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78. Say L, Gülmezoglu AM, Hofmeyr GJ. Bed rest in hospital for suspected impaired fetal growth. *Cochrane Database of Systematic Reviews* 1996, Issue 1. [DOI: 1002/14651858.CD000034]
79. Self JL, Serdula M, Dowswell T, De-Regil LM. Fortification of condiments and seasonings with iron for preventing anaemia and improving health. *Cochrane Database of Systematic Reviews* 2012, Issue 2. [DOI: 1002/14651858.CD009604]
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81. Sibley LM, Sipe TA, Barry D. Traditional birth attendant training for improving health behaviours and pregnancy outcomes. *Cochrane Database of Systematic Reviews* 2012, Issue 8. [DOI: 1002/14651858.CD005460.pub3]
82. Siegfried N, Irlam JH, Visser ME, Rollins NN. Micronutrient supplementation in pregnant women with HIV infection. *Cochrane Database of Systematic Reviews* 2012, Issue 3. [DOI: 1002/14651858.CD009755]
83. Siegfried N, van der Merwe L, Brocklehurst P, Sint TT. Antiretrovirals for reducing the risk of mother-to-child transmission of HIV infection. *Cochrane Database of Systematic Reviews* 2011, Issue 7. [DOI: 1002/14651858.CD003510.pub3]
84. Stade BC, Bailey C, Dzendoletas D, Sgro M, Dowswell T, Bennett D. Psychological and/or educational interventions for reducing alcohol consumption in pregnant women and women planning pregnancy. *Cochrane Database of Systematic Reviews* 2009, Issue 2. [DOI: 1002/14651858.CD004228.pub2]
85. Stampalija T, Gyte GM, Alfirevic Z. Utero-placental Doppler ultrasound for improving pregnancy outcome. *Cochrane Database of Systematic Reviews* 2010, Issue 9. [DOI: 1002/14651858.CD008363.pub2]
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87. Turnbull C, Osborn DA. Home visits during pregnancy and after birth for women with an alcohol or drug problem. *Cochrane Database of Systematic Reviews* 2012, Issue 1. [DOI: 1002/14651858.CD004456.pub3]
88. Vazquez JC, Abalos E. Treatments for symptomatic urinary tract infections during pregnancy. *Cochrane Database of Systematic Reviews* 2011, Issue 1. [DOI: 1002/14651858.CD002256.pub2]
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92. Webb RT, Howard L, Abel KM. Antipsychotic drugs for non-affective psychosis during pregnancy and postpartum. *Cochrane Database of Systematic Reviews* 2004, Issue 2. [DOI: 1002/14651858.CD004411.pub2]
93. Whitworth M, Dowswell T. Routine pre-pregnancy health promotion for improving pregnancy outcomes. *Cochrane Database of Systematic Reviews* 2009, Issue 4. [DOI: 1002/14651858.CD007536.pub2]
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CONTRIBUTIONS OF AUTHORS

ZA, JV, NM and AC designed the study and wrote the protocol. All authors contributed to eligibility assessment, creation of data tables and writing the text for the overview.

DECLARATIONS OF INTEREST

NM: Nancy Medley's work was financially supported by the University of Liverpool's Harris-Wellbeing of Women Preterm Birth Centre research award. NM was an author on a Cochrane systematic review included in the overview. Assessments for this review was made by another member of the overview team.

JV: none known.

AC: none known.

ZA: Zarko Alfirevic was an author on a Cochrane systematic review included in the overview and also an author on a clinical trial included in an eligible Cochrane systematic review. Assessments for these were made by another member of the overview team. My employer (University of Liverpool) has received grants from UK National Institute of Health Research, Wellbeing of Women charity and Perkin Elmer to support my research group's work related to preterm birth prevention and my Cochrane editorial work.

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INDEX TERMS

Medical Subject Headings (MeSH)

*Systematic Reviews as Topic; Anti-Bacterial Agents [therapeutic use]; Bed Rest; Premature Birth [*prevention & control]; Smoking Cessation; Vitamin D [administration & dosage]; Vitamins [administration & dosage]

MeSH check words

Female; Humans; Pregnancy