Treatments for gestational diabetes: a systematic review and meta-analysis

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ABSTRACT

Objective To investigate the effectiveness of different treatments for gestational diabetes mellitus (GDM).

Design Systematic review, meta-analysis and network meta-analysis.

Methods Data sources were searched up to July 2016 and included MEDLINE and Embase. Randomised trials comparing treatments for GDM (packages of care (dietary and lifestyle interventions with pharmacological treatments as required), insulin, metformin, glibenclamide (glyburide)) were selected by two authors and double checked for accuracy. Outcomes included large for gestational age, shoulder dystocia, neonatal hypoglycaemia, caesarean section and pre-eclampsia. We pooled data using random-effects meta-analyses and used Bayesian network meta-analysis to compare pharmacological treatments (ie, including treatments not directly compared within a trial).

Results Forty-two trials were included, the reporting of which was generally poor with unclear or high risk of bias. Packages of care varied in their composition and reduced the risk of most adverse perinatal outcomes compared with routine care (eg, large for gestational age: relative risk 0.58 (95% CI 0.49 to 0.68; I²=0%; trials 8; participants 3462). Network meta-analyses suggest that metformin had the highest probability of being the most effective treatment in reducing the risk of most outcomes compared with insulin or glibenclamide.

Conclusions Evidence shows that packages of care are effective in reducing the risk of most adverse perinatal outcomes. However, trials often include few women, are poorly reported with unclear or high risk of bias and report few outcomes. The contribution of each treatment within the packages of care remains unclear. Large well-designed and well-conducted trials are urgently needed.

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INTRODUCTION

Treatment of gestational diabetes mellitus (GDM) aims to reduce hyperglycaemia and in turn reduce the risk of adverse perinatal outcomes including large for gestational age (LGA), macrosomia, shoulder dystocia, neonatal hypoglycaemia and the need for caesarean section. Diet modification is often used as first-line treatment, and if partly or wholly unsuccessful or where women have substantially elevated glucose at diagnosis, pharmacological treatments (metformin, glibenclamide (glyburide) and/or insulin) are offered.

Previous systematic reviews have investigated the effectiveness of treatments for GDM.1-15 Although results from these reviews generally indicate that treatment reduces the risk of adverse perinatal outcomes, the searches have variable inclusion criteria and were undertaken between 20091 5 and 201414 15 with three reviews with searches in 2015,9 14 15 and since then, several trials have been published and recommended criteria for GDM diagnosis have changed. Some reviews have included observational studies, and most do not review all treatments, with the exception of the Cochrane treatments review1 (which is now out of date and has been divided for future updates) and the UK National Institute for Health and Care Excellence (NICE) guideline.16 Consequently, most previous reviews do not provide an assessment of all available treatments, and most have not used a network meta-analysis to determine the most effective pharmacological treatment across all alternatives included in any randomised controlled trial (RCT).

The aim of this study was to systematically review and, where appropriate, pool all results from RCTs of the effect of any treatment on GDM and to determine which treatment is the most effective.
METHODS
We conducted a systematic review, meta-analysis and network meta-analysis to evaluate whether treatments for GDM reduce the risks of adverse perinatal outcomes and to compare the effectiveness of these treatments.

This review and meta-analysis was conducted in accordance with Cochrane systematic reviews and the Centre for Reviews and Dissemination recommendations; we have reported our findings following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) reporting guidelines (see online supplementary research checklist). This review forms part of a larger health technology assessment report of the diagnosis and management of GDM.20

Patient involvement
The outcomes we included were from the Cochrane Pregnancy and Childbirth Group’s standardised outcomes for reviews of diabetes in pregnancy. Women who had experienced or had the potential to experience GDM contribute to the design and appraisal of this group’s methods and reviews and therefore have influenced the design of this review and outcomes examined.21

Search methods
The search strategies were designed to identify records of RCTs of treatment of women with GDM, added to search sources since the search date (July 2011, trials awaiting classification) of the Cochrane ‘treatments for GDM’ review. The bibliographic databases searched were MEDLINE and MEDLINE in Process, Embase and the Cochrane Central Register of Controlled Trials. Strategies were not restricted by language and were developed using a combination of subject indexing terms and free text search terms in the title and abstract fields. Searches were first conducted in September 2013 and updated in October 2014 and 6 July 2016, using the same search strategies. Information on studies in progress was sought by searching relevant trial registers including ClinicalTrials.gov.

We also searched previously published systematic reviews to ensure any eligible RCTs from these were included in our review if eligible. In addition, we checked the references of included journal articles. An example of search terms for MEDLINE is included in online supplementary file 1.

Study selection
Inclusion and exclusion criteria
We included RCTs in which women with diagnosed GDM or impaired glucose tolerance (IGT) (using any definition) were randomised to a treatment designed to lower blood glucose (pharmacological or dietary modification) compared with routine antenatal care (however defined by the trial) or another treatment. Trials including women with pre-existing diabetes were excluded. Trials had to report effects on adverse perinatal outcomes. Included
outcomes (defined in any way by the trials) were gestational age at birth, birth weight (BW), macrosomia, LGA, shoulder dystocia, preterm birth (less than 37 weeks gestation), neonatal hypoglycaemia, admission to neonatal intensive care unit (NICU), caesarean section (elective or emergency), pre-eclampsia, pregnancy-induced hypertension (PIH), induction of labour, instrumental birth (forceps or ventouse), Apgar score at 5 min and negative treatment effects (eg, gastrointestinal upset, well-being). Data on side effects and quality of life measures were also examined. Conference abstracts and letters to journals were eligible for inclusion if they reported sufficient outcome data.

**Data extraction and risk of bias assessment**

Title and abstract screening and then full-text screening were performed by two reviewers (DF, MS, MB or SG) with disagreements resolved by consensus or by the third reviewer. The risk of bias of the included trials was assessed using the Cochrane risk of bias tool, which considers sequence generation, allocation concealment, blinding of participants and medical staff to treatment allocation, blinding of assessors, loss to follow-up, selective reporting of outcomes and other sources of bias. Each criterion was classified as being at low or high risk of bias or unclear. Two reviewers independently assessed all criteria (DF, MS or SG).

**Statistical analysis**

Trials were divided into categories according to the following included treatments: (1) insulin versus metformin; (2) insulin versus glibenclamide (glyburide); (3) metformin versus glibenclamide; (4) packages of care: diet or dietary advice with or without exercise or glucose monitoring, with or without supplemental metformin, glibenclamide or insulin, compared with routine antenatal care; and (5) comparisons of different dietary modifications.

For dichotomous outcomes, the relative risk (RR) comparing each group, with its 95% CI, was calculated from the numbers of outcome events in each randomised group and the number randomised to each group. For continuous outcomes, the difference in means between groups was calculated from the mean and SD of the outcome. For each outcome and within each of the treatment categories, RRs or differences in means were pooled in random-effects DerSimonian-Laird meta-analyses. Heterogeneity was assessed using $I^2$. Analyses were performed to investigate differences in risk of outcomes across varying degrees of hyperglycaemia (defined by a positive/negative GDM screening and diagnostic test). Because of the large number of treatments and outcome comparisons, pooled estimates only are presented in the main paper. Tests for publication bias were considered, but not performed, because there were insufficient trials in any meta-analysis for such tests to be reliable.

We also conducted a network meta-analysis to combine information across multiple treatments simultaneously; this combines direct and indirect data to improve the estimation of the effect of treatments and specifically to try to estimate which is the most effective of a number of different treatment options. Analyses were undertaken for each dichotomous outcome using a Bayesian approach, based on the models originally created by Lu and Ades, using the OpenBUGS software. The model has a ‘binominal-normal’ structure; that is, events were assumed to follow a binomial distribution, with log odds and random effects being normally distributed. Vague normal priors (mean 0, variance 10000) were used except for heterogeneity, where an inverse-gamma (0.1, 0.1) distribution was used. The model fit and consistency were assessed by comparing the results to the meta-analyses comparing each treatment directly.

Each model generated a comparison between treatments, expressed as an OR and as a percentage indicating the probability that the treatment was the best treatment to reduce the incidence of the adverse outcome. ORs were used to ensure model stability because log ORs more closely follow a normal distribution than RRs. The probabilities of being most effective treatment were calculated from the posterior odds as part of the Bayesian model developed by Lu and Ades. This approach was not possible for continuously measured outcomes and so was not undertaken for gestational age, BW and Apgar score. As there were no trials comparing diet modification to pharmacological treatments, diet modification could not be included in the network meta-analyses.

**RESULTS**

**Details of included and excluded trials**

A total of 12234 citations were identified by the original and the two update searches. These citations were combined with three additional citations identified by previous systematic reviews conducted prior to our first searches. Following de-duplication and inclusion of additional records, 6437 citations were reviewed. Of these, 214 were judged potentially eligible based on title and abstract. After obtaining the full-text publications and assessing eligibility, 42 trials were included, and 35 of these were combined in at least one meta-analysis (figure 1).

Having extracted data from the RCTs assessing packages of care and dietary intervention comparisons (table 1), we decided that it was not appropriate to pool results from trials comparing dissimilar dietary modification interventions (table 1). Packages of care included various combinations of interventions; however, all packages of care compared with routine care trial results were pooled in meta-analyses.

We included eight publications not included in any previous published review. One compared metformin and insulin; one, glibenclamide and insulin; four, packages of care with routine care; and two compared different dietary modification interventions. Six of these trials were reported after the
### Table 1: Trials comparing a package of care starting with dietary modification to routine care and trials comparing a dietary modification with another dietary modification

| First author | Year | Location | No | Screening strategy used to determine need for diagnostic test | Diagnostic test and glucose thresholds used to diagnose GDM (mmol/L) | Intervention group | Control group | Insulin use in diet group | In meta-analyses | Meta-analysis outcome |
|--------------|------|----------|----|-------------------------------------------------------------|----------------------------------------------------------------|-------------------|---------------|------------------------|----------------|-----------------------|
| Bevier       | 1999 | USA      | 103| 50 g OGCT >7.8                                             | Positive OGCT, negative 100 g OGTT, levels not reported           | Dietary counselling and home monitoring                        | Routine care     | If needed              | Yes           | Apgar 5 min, BW, C-section, GA at birth, induction, instrumental birth, macrosomia, pre-eclampsia, shoulder dystocia |
| Bonomo       | 2005 | Italy    | 300| Risk factors and 50 g OGCT                                 | Positive OGCT >7.8, negative 100 g OGTT ‘C&C criteria’           | Dietary advice and monitoring                                  | Routine care     | Not reported            | Yes           | Apgar 5 min, BW, C-section, GA at birth, LGA, macrosomia, NN hypoglycaemia, NICU admission |
| Crowther     | 2005 | UK/Australia | 1000| Risk factors or 50 g OGCT                                 | 75 g OGTT fasting <7.8 and 2 hours >7.8 and <11.1             | Individualised dietary advice, monitoring and pharmacological treatments | Routine care     | If needed              | Yes           | Apgar 5 min <7, BW, C-section GA at birth, induction, macrosomia, NN hypoglycaemia, NICU admission, pre-eclampsia, shoulder dystocia |
| Deveer       | 2013 | Turkey   | 100| Universal 50 g OGCT >7.8 and <10.0                       | Positive OGCT, negative 100 g OGTT fasting <5.3, 1 hour <10.0, 2 hours <8.8 and 3 hours <7.8 | Calorie diet                                               | Routine care     | Not reported            | Yes           | BW, C-section, GA at birth, LGA, macrosomia, NICU admission, pre-eclampsia, preterm birth |
| Elnour       | 2006 | UAE      | 180| Not reported                                               | 100 g OGTT, ‘C&C criteria’                                      | Diet education, exercise, monitoring and pharmacological treatments | Routine care     | If needed              | Yes           | C-section, LGA, macrosomia, NN hypoglycaemia, NICU admission, pre-eclampsia, preterm birth |
| Fad           | 2015 | Sweden   | 66 | Risk factors                                               | 75 g OGTT <7.0, >10.0, <12.2                                    | Diet education, exercise, monitoring and pharmacological treatments | Routine care     | If needed in intervention group only | Yes           | BW, C-section, LGA, GA at birth, macrosomia, pre-eclampsia, instrumental birth, induction, NICU admission |

Continued
| First author | Year | Location | No  | Screening strategy used to determine need for diagnostic test | Diagnostic test and glucose thresholds used to diagnose GDM (mmol/L) | Intervention group | Control group | Insulin use in diet group | In meta-analyses | Meta-analysis outcome |
|--------------|------|----------|-----|-------------------------------------------------------------|---------------------------------------------------------------|-------------------|-------------|--------------------------|----------------|----------------------|
| Garner³⁹     | 1997 | Canada   | 299 | 75 g OGCT >8.0                                             | 75 g OGTT fasting >7.5 and 2 hours >9.6                       | Dietary counselling, restricted calorie intake, monitoring and insulin if required | Routine care    | If needed    | Yes                      | BW, C-section, GA at birth, macrosomia, NN hypoglycaemia, pre-eclampsia, preterm birth, shoulder dystocia |
| Landon⁵⁰     | 2009 | USA      | 958 | 50 g OGCT >7.5 to <11.1                                    | 100 g OGTT fasting <5.3, 2 or more, 1 hour >8.6 or 2 hours >8.6 | Individualised dietary advice, monitoring and insulin         | Routine care    | If needed    | Yes                      | BW, C-section, GA at birth, induction, macrosomia, NN hypoglycaemia, NICU admission, pre-eclampsia, preterm birth, shoulder dystocia |
| Li⁵¹         | 1987 | Hong Kong| 58  | Risk factors                                               | 100 g OGTT, two or more: fasting >5.8, 1 hour >10.6, 2 hours >9.2, 3 hours >8.1, then 75 g OGTT fasting <8.0 or 2 hours <11.0 | 30–35g/kg carbohydrate diet and monitoring                       | Routine care    | Not reported | Yes                      | BW, C-section, GA at birth, macrosomia |
| O’Sullivan³² | 1966 | USA      | 615 | OGCT or risk factors                                       | 100 g OGTT two or more fasting >6.1, or 1 hour >9.1 or 2 hours >6.7 or 3 hours >6.1 | Low-calorie diabetic diet                                      | Standard diabetic diet | Only in intervention group | Yes                      | Macroemia, preterm birth |
| Yang³⁵       | 2003 | China    | 150 | Not reported                                               | Not reported                                                 | ‘Intensive’ diabetes management                                 | Routine care    | If needed    | Yes                      | C-section, shoulder dystocia |
| Yang³⁶       | 2014 | China    | 700 | 75 g OGTT fasting 5.1, 1 hour 10.0, 2 hours 8.5            | Individual and group dietary/physical intervention             | Routine care                                                 | If needed    | Yes                      | BW, C-section, GA at birth, induction, macrosomia, NN hypoglycaemia, PH, pre-eclampsia, preterm birth, shoulder dystocia |

**Trials comparing a dietary modification with another dietary modification**

| First author | Year | Location | No  | Screening strategy used to determine need for diagnostic test | Diagnostic test and glucose thresholds used to diagnose GDM (mmol/L) | Intervention group | Control diet | Insulin use in diet group | In meta-analyses | Meta-analysis outcome |
|--------------|------|----------|-----|-------------------------------------------------------------|---------------------------------------------------------------|-------------------|-------------|--------------------------|----------------|----------------------|
| Asemi⁵³      | 2014 | Iran     | 52  | 50 g OGCT                                                  | OGCT >7.8, 75 g OGTT fasting >5.1, 1 hour >10.0, 2 hours >8.5 | DASH diet                                    | Control diet   | Women with GDM excluded, therefore insulin not required | No             | –                     |
| Cypryk⁵⁴     | 2007 | Poland   | 30  | Not reported                                               | Levels not reported only that the WHO criteria were used       | High-carbohydrate diet                                | Low-carbohydrate diet | If needed    | No                      | –              | –                     |

Continued
search dates of the previous reviews and were published in 2014 or 2015; the remaining two trials (dietary modification interventions or packages of care) did not fulfil other review’s inclusion criteria. Few trials reported side effects or measures of participant satisfaction or well-being.

Trials generally included women with GDM diagnosed following a 75 or 100 g oral glucose tolerance test (OGTT) using a variety of international39–41 and locally42 43 recommended thresholds; although some included women with ‘mild or borderline’ GDM (positive oral glucose challenge test (OGCT), negative OGTT) and others included women with IGT, current diagnostic criteria16 44, however, may now consider these women as having GDM rather than a separate and milder condition.

**Quality – risk of bias assessment**

Overall, reporting of and many aspects of trial quality were poor with the result that risk of bias was generally unclear or high (online supplementary table 1). The randomisation procedure and group allocation were rarely described, although all trials reported that participants were ‘randomised’. Blinding of participants, medical staff and outcome assessors was generally not reported, but as most trials include some additional intervention above routine care such as diet advice or a pharmacological treatment, it is probable that participants and most clinicians could not be blinded, although outcome assessment could have been. Most trials had reasonably complete outcome data and loss to follow-up was low, although for some trials, analysis was not conducted on an intention-to-treat basis (so the analysis did not include all women randomised). Selective reporting was assessed as minimal, as the majority of trials presented results for all prespecified outcomes (the possibility that some trials collected data on outcomes but did not report them cannot be ruled out however).

Generally, women were eligible for inclusion in trials evaluating pharmacological treatments if they were unable to achieve adequate glycaemic control with dietary and lifestyle management. Therefore, there is the possibility that those included may have had more severe or refractory hyperglycaemia or may adhere less well to lifestyle interventions than those women who did not require pharmacological treatments to control hyperglycaemia. The specific criteria for the addition of supplemental insulin in trials were often not reported, although some trials did report that supplemental insulin was prescribed if ‘glycaemic control was not achieved by participants’. It is probable that thresholds for what is defined as ‘good’ control differed between trial centres (if multisite) and trials.

**Packages of care and dietary modification trials**

Twelve trials evaluated a package of care (a combination of treatments starting with dietary modification and/or exercise and/or monitoring and/or supplemental pharmacological treatments) (table 1)33–36 45–52 compared with

| First author Year | Location | Diagnostic test and glucose thresholds used to determine need for diagnostic test | Intervention group | Control group |
|-------------------|----------|---------------------------------|-------------------|--------------|
| Louie55 2011 Australia | OGTT >5.5, 1 hour >10.0, or 2 hours >8.0 | No reported | No reported |
| Ma56 2013 China | OGCT >7.8 | 50 g OGCT | High-fibre diet |
| Moreno-Castilla57 2000 Spain | OGTT fasting >7.8 | 50 g OGCT >7.8 | Low GI diet |
| Rae57 2015 China | OGTT fasting >7.8 | 100 g OGTT >5.8, 1 hour >10.6 | Low-GI diet |
| Yao56 2015 China | OGTT fasting >7.8 | 100 g OGTT >5.8, 1 hour >10.6 | USual diet |

*Women who required insulin were excluded from the trial’s analyses. BW, birth weight; C-section, caesarean section; DASH diet, dietary approaches to stop hypertension; GA, gestational age; GDM, gestational diabetes mellitus; LGA, large for gestational age; Meta-analysis outcome, in meta-analyses; NN, neonatal; OGCT, oral glucose challenge test; OGTT, oral glucose tolerance test; PIH, pregnancy-induced hypertension.

**Table 1 Continued**

| Screening strategy used to determine need for diagnostic test | Meta-analysis analyseres: outcome | 55 g OGTT | 75 g OGTT | 100 g OGCT |
|------------------------------------------------------------|----------------------------------|-----------|-----------|------------|
| OGTT >5.5, 1 hour >10.0, or 2 hours >8.0                  | No reported                      | No reported | No reported | No reported |
| OGCT >7.8                                                  | 50 g OGCT                       | 50 g OGCT | Low-GI diet | Low-GI diet |
| OGTT fasting >7.8                                          | 100 g OGTT                       | 100 g OGTT | Low-GI diet | Low-GI diet |

**Table 1**

| First author Year | Location | Screening strategy used to determine need for diagnostic test | Meta-analysis analyseres: outcome | 55 g OGTT | 75 g OGTT | 100 g OGCT |
|-------------------|----------|---------------------------------------------------------------|----------------------------------|-----------|-----------|------------|
| Louie55 2011 Australia | OGTT >5.5, 1 hour >10.0, or 2 hours >8.0 | No reported                          | No reported                      | No reported | No reported | No reported |
| Ma56 2013 China | OGCT >7.8 | 50 g OGCT | No reported | No reported |
| Moreno-Castilla57 2000 Spain | OGTT fasting >7.8 | 50 g OGCT | Low GI diet | No reported |
| Rae57 2015 China | OGTT fasting >7.8 | 100 g OGTT | Low-GI diet | No reported |
| Yao56 2015 China | OGTT fasting >7.8 | 100 g OGTT | USual diet | No reported |
Seven trials \cite{37,38,53-57} evaluated a variety of dietary modifications and compared them to other dietary modifications (table 1). The composition of each dietary modification was generally well reported; however, the interventions and comparisons were too diverse to allow pooling of data. There was no evidence that one type of dietary modification was superior over another, although trials included few women (online supplementary figures 1 and 2). None of these seven trials reported side effects or quality of life measures.

The composition of the dietary modification was poorly reported in the ‘packages of care’ trials (the 12 trials included in the meta-analyses). Overall (in all packages of care and dietary modification trials), 10 out of 19 trials reported that insulin was provided if required; in one trial, insulin was only provided if needed in the intervention group; and for the remainder, it was unclear or not reported if supplemental insulin was provided. The screening and diagnostic tests, criteria and glucose
thresholds used to define GDM (and included/exclude women in the trials) varied across the trials (table 1). For the meta-analysis, the varying forms of dietary modification and/or pharmacological treatment use were not examined.

Packages of care (starting with dietary modification and possibly including monitoring and pharmacological interventions) reduced the risk of shoulder dystocia by 60%, LGA and macrosomia by around 50%, pre-eclampsia by 20% and the incidence of caesarean section by 10% compared with routine care (figure 2A), although for pre-eclampsia and caesarean section, the CIs included the null value. BW was reduced by approximately 110 g in the packages of care compared with routine care group (figure 2B). The degree of heterogeneity (I²) varied by outcome from 0% to 77%. No ‘packages of care trial’ reported side effects; two trials reported quality of life scores47 48 indicating higher (better) quality of life scores for women in the intervention compared with the routine care group.

**Trials comparing metformin with insulin**

Eleven trials compared metformin with insulin (table 2).31 45 58–66 However, most trials reported supplemental insulin use in the metformin group with the exception of two trials.31 64 The risk of most outcomes, including LGA, macrosomia, NICU admission, neonatal hypoglycaemia, pre-eclampsia, PIH and induction of labour, was lower in those randomised to metformin rather than insulin; instrumental delivery was greater in those randomised to insulin (figure 2C). BW, gestational age and Apgar score as continuous measurements did not differ notably between the two treatments (figure 2D). Six trials reported the proportion of women with metformin-associated gastrointestinal upset (between 4% and 46%).58–60 63 65 66 No trial reported quality of life measures.

**Trials comparing glibenclamide (glyburide) with insulin**

Nine trials compared glibenclamide with insulin (table 3).72 67–74 Figure 2E shows the RRs of dichotomous outcomes, suggesting that insulin may be relatively more effective than glibenclamide in reducing the risk of several adverse outcomes; CIs are wide and include the null value however. There was no difference between insulin and glibenclamide for continuous outcomes (figure 2F). One trial reported that glibenclamide was associated with side effects in 3/48 (6%) of women.72 No trial reported quality of life measures.

**Trials comparing glibenclamide (glyburide) with metformin**

Only three trials were identified that directly compared glibenclamide with metformin, and these were relatively small trials including between 149 and 200 women (table 4).75–77 Figure 2G shows the risk of dichotomous, and figure 2H shows continuous outcomes. These suggest that metformin is more effective at reducing risk of LGA and possibly macrosomia. However, for several of the outcomes (eg, LGA), only data from one of these trials are available; it is therefore not possible to make robust conclusions about the relative benefits of metformin and glibenclamide from these direct comparisons. No trials reported side effects or quality of life measures.

**Network meta-analysis comparing glibenclamide (glyburide), insulin and metformin**

Figure 3 shows the relationship of treatment comparisons, and table 5 shows the estimated probability of a treatment being the most effective at reducing the risk of each dichotomous outcome. Only dichotomous outcomes reported in at least two glibenclamide trials (either in comparison to insulin or metformin) were included in these analyses to ensure that there were sufficient trials (and participants) included. When all three treatments are jointly compared, these analyses suggest that, for all outcomes, with the exception of caesarean section, metformin is most likely to be the most effective treatment, with its probability of being most effective in reducing risk being 96.3%, 94.0%, 92.8%, 84.0% and 61.2%, respectively, for neonatal hypoglycaemia, macrosomia, LGA, pre-eclampsia and admission to NICU (the probability of being most effective for reducing risk of caesarean section was 9.7% for metformin, glibenclamide was most likely to be most effective at reducing the risk of caesarean section (79.9%)). The results of the network meta-analysis (figure 4) are consistent with the direct comparisons between treatments shown in figure 2A–H, suggesting that metformin is more effective than insulin or glibenclamide at reducing the majority of adverse outcomes. However, many of these comparisons are based on small numbers and have wide CIs that sometimes include the null value.

**DISCUSSION**

The key finding of our review is that, despite understanding of hyperglycaemia/GDM and its relationship to adverse perinatal outcomes having existed for at least seven decades78 and 42 RCTs completed on its treatment, trials are still being conducted that are of limited size and of poor quality (with subsequent unclear or high risk of bias), and therefore, which treatment is the most effective remains unclear. Given the changing characteristics of the population and the lower fasting diagnostic threshold (compared with previous criteria)40 recommended by the International Association of Diabetes and Pregnancy Study Groups (IADPSG)44 and UK NICE,16 it is important to understand how treatments affect outcomes for these women. Trials do not always report GDM diagnostic criteria clearly, and this is important considering the potential influence on GDM population size and the magnitude of effect.16 44 Our detailed review, including only evidence from RCTs, provides some support for a ‘step up approach’ in the treatment of hyperglycaemia, from dietary interventions, through addition of metformin (in preference to glibenclamide
Table 2  Trials comparing metformin to insulin

| First author | Year | Location | No | Diagnostic test and glucose thresholds used to diagnose GDM | Screening strategy* | Meta-analysis outcome |
|--------------|------|----------|----|-------------------------------------------------------------|---------------------|----------------------|
| Ainuddin65   | 2014 | Pakistan | 150| 75 g OGTT two or more; fasting 5.3, 1 hour 10.0, 2 hours 8.6 | 50 g OGCT ≥7.8      | PIH, pre-eclampsia, GA at delivery, induction, C-section, LGA, NICU admission, neonatal hypoglycaemia |
| Hague64      | 2003 | Australia| 30 | 75 g OGTT fasting >5.5 or 2 hours >8.0                     | Risk factors        | BW, pre-eclampsia, GA at birth, induction, C-section, macrosomia, hypoglycaemia |
| Hassan65     | 2012 | Pakistan | 150| 75 g OGTT two or more levels fasting >5.3, 1 hour >10.0 or 2 hours >8.6 | 50 g OGCT >7.8      | Apgar 5 min, GA at birth, induction, C-section, BW, macrosomia, hypoglycaemia, NICU admission |
| Ijas63       | 2010 | Finland | 100| 75 g OGTT fasting >5.3, 1 hour >11.0 or 2 hours >9.6       | Risk based          | Apgar 5 min, BW, C-section, GA at birth, induction, instrumental birth, LGA, macrosomia, hypoglycaemia, NICU admission |
| Mesdaghinia62| 2013 | Iran    | 200| 100 g OGTT two or more; fasting >5.3 or 1 hour >10.0 or 2 hours >8.6 or 3 hours >7.8 | 50 g OGCT - levels not reported | BW, macrosomia, LGA, hypoglycaemia, NICU admission, shoulder dystocia, 5 min Apgar <7, preterm birth |
| Moore61      | 2007 | USA     | 63 | 100 g OGTT two or more; fasting >5.8 or 1 hour >10.5 or 2 hours >9.1 or 3 hours >8.0 | 50 g OGCT >7.8      | Apgar 5 min, BW, macrosomia, hypoglycaemia, NICU admission |
| Niromanesh60 | 2012 | Iran    | 160| 100 g OGTT two or more fasting >5.3, 1 hour >10.0, 2 hours >8.6 or 3 hours >7.8 | 50 g OGCT >7.2      | Apgar 5 min, pre-eclampsia, PIH GA at birth, induction, C-section, shoulder dystocia, BW macrosomia, LGA, NICU admission, hypoglycaemia, preterm birth |
| Rowan59      | 2008 | Australia / NZ | 751| 75g OGTT fasting >5.5 or 2 hours >8.0                     | Risk factors        | Apgar 5 min <7, BW, GA at birth, LGA, NICU admission, PIH, pre-eclampsia, preterm birth |
| Spaulonci58  | 2013 | Brazil  | 94 | 75 g or 100 g OGTT fasting >5.3 or 1 hour >10.0 or 2 hours >8.0 and two or more fasting >5.3, 1 hour >10.0, 2 hours >8.0 or 3 hours >7.8, respectively | No screening        | GA at birth, BW, Apgar 5 min, macrosomia, hypoglycaemia, pre-eclampsia, preterm birth, C-section |
| Tertti43     | 2013 | Finland | 217| 75 g OGTT both criteria: fasting ≥4.8, 1 hour ≥10.0, 2 hours ≥8.7 and fasting ≥5.3, ≥10.0 and ≥8.6, respectively | Risk factors        | GA at birth, BW, Apgar at 5 min, induction, instrumental birth, C-section, LGA, macrosomia, preterm birth, PIH, pre-eclampsia, NICU admission, hypoglycaemia |
| Zinnat31     | 2013 | Bangladesh | 450| Not reported†                                           | Not reported†       | Macrosomia, shoulder dystocia, C-section, instrumental birth hypoglycaemia, NICU admission |

*It is assumed unless otherwise reported that the screening strategy advocated by the criteria used was adhered to.
†Conference abstract.
BW, birth weight; C-section, caesarean section; GA, gestational age; GDM, gestational diabetes mellitus; LGA, large for gestational age; NICU, neonatal intensive care unit.
Table 3  Trials comparing glibenclamide (glyburide) to insulin

| First author | Year | Location | No | Diagnostic test and glucose thresholds used to diagnose GDM | Screening strategy* | Outcome |
|--------------|------|----------|----|-----------------------------------------------------------|--------------------|---------|
| Anjalakshi57  | 2007 | India    | 23 | 75 g OGTT 2 hours >7.8                                   | Universal OGTT     | BW      |
| Bertin68      | 2005 | Brazil   | 70 | 75 g OGTT fasting >6.1 or 2 hours >7.8                  | Not reported       | BW, C-section, Apgar 5 min, GA at birth, LGA |
| Lain69        | 2009 | USA      | 99 | 100 g OGTT two or more: fasting >5.3, 1 hour >8.6 or 2 hours >8.6 | 50 g >7.5         | BW, GA at birth, LGA, macrosomia |
| Langer70      | 2000 | USA      | 404| 100 g OGTT fasting >5.3 to <7.8                         | 50 g OGCT >7.3     | BW, C-section, GA at birth, LGA, macrosomia, hypoglycaemia, NICU admission, pre-eclampsia |
| Mirzamoradi32  | 2015 | Iran     | 96 | Glucose load not reported; OGTT two or more: fasting >5.3, 1 hour >10.0, 2 hours >8.3 | Universal OGTT     | BW, C-section, GA at birth, NICU admission, hypoglycaemia, pre-eclampsia |
| Oggunyemi71   | 2007 | USA      | 97 | Not reported                                             | No screening       | BW, GA at birth, LGA, hypoglycaemia |
| Silva73       | 2007 | Brazil   | 68 | 75 g OGTT fasting >6.1 or 2 hours >7.8                  | No screening       | BW, C-section, GA at birth, hypoglycaemia, |
| Tempe74       | 2013 | India    | 64 | 100 g OGTT two or more: fasting >5.3, 1 hour >10.0, 2 hours >8.6 or 3 hours >7.8 | 50 g OGCT >7.2     | BW, GA birth, macrosomia, hypoglycaemia, NICU admission, pre-eclampsia, preterm birth |

*It is assumed unless otherwise reported that the screening strategy advocated by the criteria used was adhered to.

BW, birth weight; C-section, caesarean section; GA, gestational age; GDM, gestational diabetes mellitus; LGA, large for gestational age; NICU, neonatal intensive care unit.
Table 4  Trials comparing glibenclamide to metformin

| First author | Year | Location | No | Diagnostic test and thresholds used to diagnose GDM (mmol/L) | Screening strategy | Outcome |
|--------------|------|----------|----|------------------------------------------------------------|-------------------|---------|
| George       | 2015 | India    | 159| 100 g OGTT two or more; fasting >5.3 or 1 hour >10.0 or 2 hours >8.6 | Not reported       | BW, GA at birth, macrosomia, hypoglycaemia. |
| Moore        | 2010 | USA      | 149| 100 g OGTT two or more; fasting >5.3 or 2 hours >6.7, 50 g OGCT >7.2 | BW, C-section, GA at birth, macrosomia, hypoglycaemia, NICU admission | BW, C-section, GA at birth, macrosomia, hypoglycaemia, NICU admission |
| Silva        | 2012 | Brazil   | 200| 75 g OGTT fasting >5.3 or 1 hour >10.0 or 2 hours >8.0        | No screening      | BW, birth weight, C-section, caesarean section; GA, gestational age; GDM, gestational diabetes mellitus; LGA, large for gestational age; NICU, neonatal intensive care unit. |

BW, birth weight; C-section, caesarean section; GA, gestational age; GDM, gestational diabetes mellitus; LGA, large for gestational age; NICU, neonatal intensive care unit.

adverse outcomes and for women diagnosed using more recently recommended criteria. Hence, we feel that it is important to place a moratorium on further small RCTs in this area and that funders should consider commissioning a multicentre large-scale RCT with adequate power to determine the effect and cost-effectiveness of different packages of care on adverse outcomes in women with GDM.

The evidence to support metformin use, although encouraging, has certain weaknesses. First, although there is a general ‘trend’ in favour of metformin use over insulin and glibenclamide (glyburide), CIs are wide, in both the direct and network meta-analysis comparing each two-way treatment effect. Second, the reporting of trial methods was generally poor with ‘unclear or high risk of bias’, and many trials included relatively few women and reported few outcomes. Third, in most trials directly comparing metformin with insulin, women receiving metformin were also given supplemental insulin ‘if required’; in one of the largest trials, this equated to 46% of the metformin group. Therefore, our results more appropriately relate to metformin’s greater effectiveness as a first-line treatment for GDM rather than a standalone treatment compared with insulin.

In addition to being an effective first-line pharmacological treatment for GDM, metformin may also be preferred by women as it is administered orally and can be stored at room temperature, compared with insulin that requires subcutaneous injection and refrigerated storage. Metformin is sometimes associated with gastrointestinal upset, which may affect compliance and quality of life.

Few trials have reported side effects or measures of participant satisfaction or well-being, all important outcomes that have the potential to impact health and therefore should be evaluated. Recent guidance recommends lower glucose thresholds compared with those previously recommended to diagnose GDM (and used in the included trials). Therefore, it is possible that a greater proportion of women diagnosed with GDM will require only diet modification or less ‘intensive’ management compared with those previously diagnosed with GDM because their hyperglycaemia is less severe. There is a continuum of increasing risk of adverse outcomes across the spectrum of glucose however; interventions to reduce hyperglycaemia even at lower glucose levels are likely to improve outcomes, but this needs confirming by large well-designed RCTs.

Strengths and limitations

This systematic review and meta-analysis includes a large number of trials with varied populations and examines the effectiveness of treatment packages and diets as well as individual pharmacological treatments for reducing the risk of adverse perinatal outcomes.

For some comparisons, trials and numbers of women were few, as were outcomes reported. Trial quality was generally poor with subsequent high or unclear risk of bias. GDM diagnostic criteria varied across trials, and recently
recommended thresholds are lower now compared with when most included trials were conducted.

Lower glucose threshold criteria recommended by the International Association of Diabetes and Pregnancy Study Groups44 and subsequently endorsed by the WHO62 aim to identify offspring at risk of obesity through its association with LGA (birth weight >90th percentile), cord C-peptide >90th percentile and percentage body fat >90th percentile. However, there are no trials that have used these criteria, and the classification of less severe hyperglycaemia when lower glucose thresholds are used to diagnose GDM may reduce the magnitude of the effect of interventions, compared with those reported by earlier trials using higher glucose thresholds. There has also been no longer term follow-up conducted to evaluate the treatment of GDM and the effects on risk of offspring outcomes. Importantly, few of the trials that we reviewed had reported side effects or measures of participant satisfaction or well-being.

Implications for practice
This review provides reassurance that a package of care where a ‘step up’ approach of first providing dietary and lifestyle advice, then adding supplementary metformin or insulin if glucose levels are not adequately controlled, is a reasonable and effective approach compared with providing just routine antenatal care, particularly with regard to reducing the risk of LGA. However, it has also highlighted the general poor quality of recent small RCTs that do not improve the evidence base but subject women with GDM to unnecessary ‘experimentation’ and are a cost to society.

Metformin seems to be an effective alternative to insulin, if diet modification inadequately controls hyperglycaemia; however, supplemental insulin may be required in up to 50% of women.59 There is a need to cease further small RCTs in this area and conduct large well-designed RCTs that clarify the most effective treatment across a range of outcomes, including those that are likely to be important to women such as quality of life measurements and those identified by the Cochrane Pregnancy and Childbirth Group as being essential for trials and reviews of diabetes in pregnancy. These should be incorporated into current diagnostic criteria and ideally look at longer term outcomes in mothers and offspring.

Correction notice This paper has been amended since it was published Online First. Owing to a scripting error, some of the publisher names in the references were replaced with ‘BMJ Publishing Group’. This only affected the full text version, not the PDF. We have since corrected these errors and the correct publishers have been inserted into the references.

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REFERENCES

1. Alwan N, Tuffnell D, West J, et al. Treatments for gestational diabetes. Cochrane Database Syst Rev 2009;3:CD003395.
2. Gui J, Liu Q, Feng L. Metformin vs insulin in the management of gestational diabetes: a meta-analysis. PLoS One 2013;8:e64585.
3. Harting A, Dryden DM, Guthrie A, et al. Benefits and harms of treating gestational diabetes mellitus: a systematic review and meta-analysis for the U.S. Preventive Services Task Force and the National Institutes of Health Office of Medical Applications of Research. Ann Intern Med 2013;159:123–9.
4. Falavigna M, Prestes I, Schmidt MI, et al. Impact of gestational diabetes mellitus screening strategies on perinatal outcomes: a simulation study. Diabetologia 2013;56:358–65.
5. Horvath K, Koch K, Jeitler K, et al. Effects of treatment in women with gestational diabetes mellitus: systematic review and meta-analysis. BMJ 2009;339:b1385–6.
6. Balsells M, Garcia-Patterson A, Sola I, et al. Glibenclamide, metformin, and insulin for the treatment of gestational diabetes: a systematic review and meta-analysis. BMJ 2015;350:h102.
7. Han S, Crowther CA, Middleton P, et al. Different types of dietary advice for women with gestational diabetes mellitus. Cochrane Database Syst Rev 2013;CD009275.
8. Jiang YF, Chen XY, Ding T, et al. Comparative efficacy and safety of OAsD in management of GDM: network meta-analysis of randomized controlled trials. J Clin Endocrinol Metab 2015;100:2071–80.
9. Amin M, Suksoomboon N, Poolspun N, et al. Comparison of glyburide with metformin in treating gestational diabetes mellitus: a systematic review and meta-analysis. Clin Drug Investig 2015;35:433–51.
10. Kittwee P, Limwattananon S, Limwattananon C, et al. Metformin for the treatment of gestational diabetes: an updated meta-analysis. Diabetes Res Clin Pract 2015;109:521–32.
11. Zhao LP, Sheng XY, Zhou S, et al. Metformin versus insulin for gestational diabetes mellitus: a meta-analysis. Br J Clin Pharmacol 2015;80:1224–34.
12. Su DF, Wang XY, Metformin in insulin in the management of gestational diabetes: a systematic review and meta-analysis. Diabetes Res Clin Pract 2014;104:353–7.
13. Zhu B, Zhang L, Fan YY, et al. Metformin versus insulin in gestational diabetes mellitus: a meta-analysis of randomized clinical trials. Int J Med Sci 2016;18:371–81.
14. Feng Y, Yang H, Metformin - a potentially effective drug for gestational diabetes mellitus: a systematic review and meta-analysis. J Matern Fetal Neonatal Med 2016;1:1–8.
15. Butalia S, Gutierrez L, Lodha A, et al. Short- and long-term outcomes of metformin compared with insulin alone in pregnancy: a systematic review and meta-analysis. Diabet Med 2017;34:27–36.
16. National Institute for Health and Care Excellence. Diabetes in pregnancy: management of diabetes and its complications from preconception to the postnatal period. NationalcollaboratingcentreformenandChildrensHealth2015https://www.nice.org.uk/guidance/ng3.
17. Higgins JPT, Green S, Cochrane Handbook for Systematic Reviews of Interventions Version 5.1.0. Cochrane Library, 2011.
18. Centre for Reviews and Dissemination. Systematic reviews: CRD’s guidance for undertaking a systematic review York: CRD, University of York, 2009.
19. PRISMA. Preferred reporting items for systematic reviews and meta-analyses. http://www.prisma-statement.org/
20. Farrar D, Simmonds M, Griffin S, et al. The identification and treatment of women with hyperglycaemia in pregnancy: an analysis of individual participant data, systematic reviews, meta-analyses and an economic evaluation. Health Technol Assess 2016;20:1-348.
21. Sakala C, Gyte G, Henderson S, et al. Consumer-professional partnership to improve research: the experience of the Cochrane Collaboration’s Pregnancy and Childbirth Group. Birth 2001;28:133–7.
22. The Cochrane Collaboration. The Cochrane Collaboration’s tool for assessing risk of bias. http://methods.cochrane.org/bias/assessing-risk-bias-included-studies#The%20Cochrane%20Risk%20of%20Bias%20Tool (accessed Jan 2014).
23. DerSimonian R, Laird N. Meta-analysis in clinical trials. Control Clin Trials 1986;7:177–88.
24. Higgins JP, Thompson SG. Quantifying heterogeneity in a meta-analysis. Stat Med 2002;21:1539–58.
25. Bafeta A, Tiniquart L, Seror R, et al. Reporting of results from network meta-analyses: methodological systematic review. BMJ 2014;348:g1741.
26. Lumley T. Network meta-analysis for indirect treatment comparisons. Stat Med 2002;21:2313–24.
27. Song F, Altman DG, Glenny AM, et al. Validity of indirect comparison for estimating efficacy of competing interventions: empirical evidence from published meta-analyses. BMJ 2003;326:472.
28. Mills EJ, Thorlund K, Ioannidis JP. Demystifying trial networks and network meta-analysis. BMJ 2013;346:f2914.
29. Lu G, Ades AE. Combination of direct and indirect evidence in mixed treatment comparisons. Stat Med 2004;23:3105–24.
30. openbugs. http://www.openbugs.net/w/FrontPage (accessed Jan 2015).
31. Zinnat ANS, Shareen S, Rahaman S. Can metformin be used in place of insulin for the treatment of GDM for low resource countries? BJOG 2013;120:35.
32. Mirzamoradi M, Bakhtiyari M, Kimiaeep et al. Investigating the effects of treatment based on single high blood glucose in gestational diabetes screening on maternal and neonatal complications. Arch Gynecol Obstet 2015;292:687–95.
33. Deveer R, Deveer M, Abkaba E, et al. The effect of diet on pregnancy outcomes among pregnant with abnormal glucose challenge test. Eur Rev Med Pharmacol Sci 2013;17:1258–61.
34. Fadl HE, Gärdefors S, Hjertberg R, et al. Randomized controlled study in pregnancy on treatment of marked hyperglycemia that is short of overt diabetes. Acta Obstet Gynecol Scand 2015;94:1181–7.
35. Yang X, Hsu-Hage B, Dong L, et al. Intensive diabetes management may improve pregnancy. Diabetes Care 2003;26:254–5.
36. Yang X, Tian H, Zhang F, et al. A randomised translational trial of lifestyle intervention using a 3-tier shared care approach on pregnancy outcomes in Chinese women with gestational diabetes mellitus but without diabetes. J Transl Med 2014;12:290.
37. Ma WJ, Huang ZH, Huang BX, et al. Effect of dietary advice for women with gestational diabetes mellitus. Database Syst Rev 2015;99:358–65.
38. Zinnat ANS, Shareen S, Rahaman S. Can metformin be used in place of insulin for the treatment of GDM for low resource countries? BJOG 2013;120:35.
39. Mirzamoradi M, Bakhtiyari M, Kimiaeep et al. Investigating the effects of treatment based on single high blood glucose in gestational diabetes screening on maternal and neonatal complications. Arch Gynecol Obstet 2015;292:687–95.
40. Deveer R, Deveer M, Abkaba E, et al. The effect of diet on pregnancy outcomes among pregnant with abnormal glucose challenge test. Eur Rev Med Pharmacol Sci 2013;17:1258–61.
41. Fadl HE, Gärdefors S, Hjertberg R, et al. Randomized controlled study in pregnancy on treatment of marked hyperglycemia that is short of overt diabetes. Acta Obstet Gynecol Scand 2015;94:1181–7.
42. Yang X, Hsu-Hage B, Dong L, et al. Intensive diabetes management may improve pregnancy. Diabetes Care 2003;26:254–5.
43. Yang X, Tian H, Zhang F, et al. A randomised translational trial of lifestyle intervention using a 3-tier shared care approach on pregnancy outcomes in Chinese women with gestational diabetes mellitus but without diabetes. J Transl Med 2014;12:290.
44. Ma WJ, Huang ZH, Huang BX, et al. Effect of dietary advice for women with gestational diabetes mellitus. Database Syst Rev 2015;99:358–65.
45. Zinnat ANS, Shareen S, Rahaman S. Can metformin be used in place of insulin for the treatment of GDM for low resource countries? BJOG 2013;120:35.
46. Mirzamoradi M, Bakhtiyari M, Kimiaeep et al. Investigating the effects of treatment based on single high blood glucose in gestational diabetes screening on maternal and neonatal complications. Arch Gynecol Obstet 2015;292:687–95.
47. Deveer R, Deveer M, Abkaba E, et al. The effect of diet on pregnancy outcomes among pregnant with abnormal glucose challenge test. Eur Rev Med Pharmacol Sci 2013;17:1258–61.
48. Fadl HE, Gärdefors S, Hjertberg R, et al. Randomized controlled study in pregnancy on treatment of marked hyperglycemia that is short of overt diabetes. Acta Obstet Gynecol Scand 2015;94:1181–7.
49. Yang X, Hsu-Hage B, Dong L, et al. Intensive diabetes management may improve pregnancy. Diabetes Care 2003;26:254–5.
50. Yang X, Tian H, Zhang F, et al. A randomised translational trial of lifestyle intervention using a 3-tier shared care approach on pregnancy outcomes in Chinese women with gestational diabetes mellitus but without diabetes. J Transl Med 2014;12:290.
51. Ma WJ, Huang ZH, Huang BX, et al. Effect of dietary advice for women with gestational diabetes mellitus. Database Syst Rev 2015;99:358–65.
42. Ilija H, Vaääräsmäki M, Morin-Papunen L, et al. Metformin should be considered in the treatment of gestational diabetes: a prospective randomised study. BJOG 2011;118:880–5.

43. Terti K, Ekblad U, Koskinen P, et al. Metformin vs. insulin in gestational diabetes. A randomized study characterizing metformin patients needing additional insulin. Diabetes Obes Metab 2013;15:246–51.

44. Metzger BE, Gabbe SG, Persson B, et al. International Association of Diabetes and Pregnancy Study Groups Consensus Panel. International Association of Diabetes and Pregnancy. Study groups recommendations on the diagnosis and classification of hyperglycemia in Pregnancy. Diabetes Care 2010;33:676–82.

45. Bevier WC, Fischer R, Jovanovic L. Treatment of women with an abnormal glucose challenge test (but a normal oral glucose tolerance test) decreases the prevalence of macrosomia. Am J Perinatol 1999;16:269–75.

46. Bonomo M, Corica D, Mion E, et al. Evaluating the therapeutic approach in pregnancies complicated by borderline glucose intolerance: a randomized clinical trial. Diabet Med 2008;25:176–83.

47. Crowther CA, Hiller JE, Moss JR, et al. Effect of treatment of gestational diabetes mellitus on pregnancy outcomes. *N Engl J Med* 2005;352:2477–86.

48. Elhouri AA, El Mugammar AB, et al. Pharmaceutical care of patients with gestational diabetes mellitus. *J Eval Clin Pract* 2008;14:131–40.

49. Garner P, Okun N, Keely E, et al. A randomized controlled trial of strict glycemic control and tertiary level obstetric care versus routine obstetric care in the management of gestational diabetes: a pilot study. *Am J Obstet Gynecol* 1997;177:190–5.

50. Landon MB, Spong CY, Thom E, et al. A multicenter, randomized trial of antibiotic prophylaxis for women with gestational diabetes mellitus. *N Engl J Med* 2011;364:2162–71.

51. Li DF, Wong VC, O’Hoy KM, et al. Is treatment needed for mild impairment of glucose intolerance in pregnancy? *A randomized controlled trial. Br J Obstet Gynaecol* 1987;94:851–4.

52. O’Sullivan JB, Gellis SS, Dandrow RV, et al. The potential diabetic and her treatment in pregnancy. *Obstet Gynecol* 1986;27:563–9.

53. Asemi Z, Samimi M, Tabassi Z, et al. The effect of DASH diet on pregnancy outcomes in gestational diabetes: a randomized controlled clinical trial. *Eur J Clin Nutr* 2014;68:490–5.

54. Cypryk K, Kamińska P, Kosiński M, et al. A comparison of the effectiveness, tolerability and safety of high and low carbohydrate diets in women with gestational diabetes. *Endokrynol Pol* 2007;58:314–9.

55. Louie JCY, Markovic TP, Perera N, et al. A randomized controlled trial investigating the effects of a low-glycemic index diet on pregnancy outcomes in gestational diabetes mellitus. *Diabetes Care* 2011;34:231–7.

56. Moreno-Castilla C, Hernandez M, Bergua M, et al. Low-carbohydrate diet for the treatment of gestational diabetes mellitus: a randomized controlled trial. *Diabetes Care* 2013;36:2233–8.

57. Rae A, Bond D, Evans S, et al. A randomised controlled trial of dietary energy restriction in the management of obese women with gestational diabetes. *Aust N Z J Obstet Gynaecol* 2000;40:416–22.

58. Spaulonzi CP, Bernardes LS, Trindade TC, et al. Randomized trial of metformin vs insulin in the management of gestational diabetes. *Am J Obstet Gynecol* 2015;213:93.e1–34.e7.

59. Rowan JA, Hague WM, Gao W, et al. Metformin versus insulin for the treatment of gestational diabetes. *N Engl J Med* 2008;358:2003–15.

60. Niromanesh S, Aliav A, Sharbaf FR, et al. Metformin compared with insulin in the management of gestational diabetes mellitus: a randomized clinical trial. *Diabetes Res Clin Pract* 2012;98:422–9.

61. Moore LE, Briery CM, Clokey D, et al. Metformin and insulin in the management of gestational diabetes mellitus: preliminary results of a comparison. *J Reprod Med* 2007;52:1011–5.

62. Mesdaghinia E, Samimi M, Homaei Z, et al. Comparison of newborn outcomes in women with gestational diabetes mellitus treated with metformin or insulin: a randomised blinded trial. *Int J Prev Med* 2013;4:327–33.

63. Ilija H, Vaääräsmäki M, Morin-Papunen L, et al. Metformin should be considered in the treatment of gestational diabetes: a prospective randomised study. *BJOG* 2011;118:880–5.

64. Hague WM, Davoren PM, Oliver J, et al. Contraindications to use of metformin. Metformin may be useful in gestational diabetes. *BMJ* 2003;326:762.

65. Hassan JA, Karim N, Sheikh Z. Metformin prevents macrosomia and neonatal morbidity in gestational diabetes. *Pak J Med Sci* 2012;28:384–9.

66. Anudder J, Karim N, Hasan AA, et al. Metformin versus insulin treatment of gestational diabetes in pregnancy in a developing country. A randomized control trial. *Diabetes Res Clin Pract* 2015;107:290–9.

67. Anjaldwani C, Balaji V, Balaji MS, et al. A prospective study comparing insulin and glibenclamide in gestational diabetes mellitus in Asian Indian women. *Diabetes Res Clin Pract* 2007;76:474–5.

68. Bertini AM, Silva JC, Taborda W, et al. Perinatal outcomes and the use of oral hypoglycemic agents. *J Perinat Med* 2005;33:519–23.

69. Lain KY, Garebaden MJ, Daftary A, et al. Neonatal adiposity following maternal treatment of gestational diabetes with glyburide compared with insulin. *Am J Obstet Gynecol* 2009;200:501.e1–501.e6.

70. Langer O, Conway DL, Berkus MD, et al. A comparison of glyburide and insulin in women with gestational diabetes mellitus. *N Engl J Med* 2000;343:1134–8.

71. Mukhopadhyay P, Bag TS, Kyal A, et al. Oral hypoglycemic glibenclamide: can it be a substitute to insulin in the management of gestational diabetes mellitus? A comparative study. *Journal of SAFOG with DVD* 2012;4:28–31.

72. Ogungbemi J, Jesse M, Davidson M. Comparison of glyburide versus insulin in management of gestational diabetes mellitus. *Endocr Pract* 2007;13:427–8.

73. Silva J, Bertini A, Taborda W FB, et al. Glibenclamide in the treatment for gestational diabetes mellitus in a compared study to insulin. *Arch Endocrinol Metab* 2007;51:541–6.

74. Tempe A, Mayanglambam RD. Glyburide as treatment option for gestational diabetes mellitus. *J Obstet Gynaecol Res* 2013;39:1147–52.

75. Moore LE, Clokey D, Rappaport VJ, et al. Metformin compared with glyburide in gestational diabetes: a randomized controlled trial. *Obstet Gynecol* 2010;115:55–6.

76. George A, Mathews JE, Sam D, et al. Comparison of neonatal outcomes in women with gestational diabetes with moderate hyperglycaemia on metformin or glibenclamide - a randomised controlled trial. *BMJ* 2010;340:e6.

77. Silva JC, Fachin DR, Coral ML, et al. Perinatal impact of the use of metformin and glyburide for the treatment of gestational diabetes mellitus. *J Perinat Med* 2012;40:225–8.

78. Lawlor DA. The Society for Social Medicine John Pemberton Lecture 2011. Developmental overnutrition - an old hypothesis with new importance? *Int J Epidemiol* 2013;42:7–29.

79. O’Sullivan JB, Mahan CM. *Diabetes* 1967;13:278–85.

80. Metzger BE, Lowe LP, Dyer AR, et al. *HAPO Study Cooperative Research Group*. Hyperglycemia and adverse pregnancy outcomes. *N Engl J Med* 2008;358:1991–2002.

81. Farrar D, Fairley L, Santorelli G, et al. Association between hyperglycaemia and adverse perinatal outcomes in South Asian and white British women: analysis of data from the born in Bradford cohort. *Lancet Diabetes Endocrinol* 2015;3:785–804.

82. World Health Organization. Diagnostic criteria and classification of hyperglycaemia first detected in pregnancy. 2013 http://apps.who.int/iris/bitstream/10665/85975/1/WHO_NMH_MND_13.2_eng.pdf?ua=1%2520%2520