Obstetric outcomes after treatment of periodontal disease during pregnancy: systematic review and meta-analysis

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ABSTRACT

Objective To examine whether treatment of periodontal disease with scaling and root planing during pregnancy is associated with a reduction in the preterm birth rate.

Design Systematic review and meta-analysis of randomised controlled trials.

Data sources Cochrane Central Trials Registry, ISI Web of Science, Medline, and reference lists of relevant studies to July 2010; hand searches in key journals.

Study selection Randomised controlled trials including pregnant women with documented periodontal disease randomised to either treatment with scaling and root planing or no treatment.

Data extraction Data were extracted by two independent investigators, and a consensus was reached with the involvement a third. Methodological quality of the studies was assessed with the Cochrane’s risk of bias tool, and trials were considered either high or low quality. The primary outcome was preterm births (<37 weeks).

Secondary outcomes were low birthweight infants (≤2500 g), spontaneous abortions/stillbirths, and overall adverse pregnancy outcome (preterm births <37 weeks and spontaneous abortions/stillbirths).

Results 11 trials (with 6558 women) were included. Five trials were considered to be of high methodological quality (low risk of bias), whereas the rest were of low quality (high or unclear risk of bias). Results among low and high quality trials provided clear evidence that no such effect exists. Among high quality studies, treatment had no significant effect on the overall rate of preterm birth (odds ratio 1.15, 95% confidence interval 0.95 to 1.40; P=0.15). Furthermore, treatment did not reduce the rate of low birthweight infants (odds ratio 1.07, 0.85 to 1.36; P=0.55), spontaneous abortions/stillbirths (0.79, 0.51 to 1.22; P=0.28), or overall adverse pregnancy outcome (preterm births <37 weeks and spontaneous abortions/stillbirths) (1.09, 0.91 to 1.30; P=0.34).

Conclusion Treatment of periodontal disease with scaling and root planing cannot be considered to be an efficient way of reducing the incidence of preterm birth. Women may be advised to have periodical dental examinations during pregnancy to test their dental status and may have treatment for periodontal disease. However, they should be told that such treatment during pregnancy is unlikely to reduce the risk of preterm birth or low birthweight infants.

INTRODUCTION

Periodontitis is a relatively common clinical condition, which occurs in more than 30% of people in some populations;1 it has a prevalence of between 5% and 20% in pregnant women.2 Treatment in pregnancy is safe and easily applicable and involves scaling and root planing.3 An association between periodontal disease and preterm birth has engendered much interest. Despite advances in obstetric care, preterm birth continues to be the leading cause of perinatal morbidity and mortality. This suggestion has led many investigators to seek evidence in this field. Since 1996, when a relation of periodontal disease with preterm birth was proposed,4 many observational studies have been carried out. Although the pathophysiological mechanism remains unclear, several studies support the hypothesis that periodontal disease is associated with preterm labour and other conditions complicating pregnancy, such as pre-eclampsia and fetal growth restriction.5 This association has also been reported by most of the 17 observational studies up to 2005 that were included in a meta-analysis published by Vergnes,6 which concluded that pregnant patients with periodontal disease have a 2.8-fold increased risk of preterm birth.

Researchers have suggested that periodontal disease causes the release of pathogens or inflammatory products such as cytokines, which then affect embryonic tissue or amniotic fluid through haematogenous transport.7 Taking into account the fact that treatment of periodontal disease is simple and safe during pregnancy,3 many studies have examined whether active management of periodontal disease has a potential beneficial effect on outcomes of pregnancy. In 2006 several of the US insurers offered their clients scaling and root planing during pregnancy at no extra cost,8 saying that they expected that spending more on
preventive dental care would yield big savings in the medical treatment of costly chronic illnesses. Although the assumption that treatment of periodontitis would have a positive result on reducing the incidence of preterm births seems logical, broad investigation is essential to reach valid conclusions. In the past, treatment of conditions correlated with preterm birth, such as vaginitis, failed to alter the incidence of preterm birth.9,11

Between 2003 and 2008 seven prospective randomised studies were published, and most of them showed a decrease in preterm birth and low birthweight infants among women with periodontal disease when the disease was treated, with scaling and root planning, compared with no treatment. In contrast, one of the largest studies showed no difference.12 Whereas pooled results of the early trials supported the hypothesis that applying treatment in patients with periodontal disease may reduce the risk of preterm birth and low birthweight infants,13 the low methodological quality of most of them prevented us from prompt implementation of the findings. Today, after the publication of more recent, well designed randomised trials, a more comprehensive meta-analysis is essential to provide solid guidelines for the treatment of periodontal disease during pregnancy.

**METHODS**

**Search strategy and eligibility criteria**

Two independent investigators (DM and AV) searched the Cochrane Central Trials Registry, ISI Web of Science, and Medline without language restriction up to July 2010 by using the search algorithm “(periodontal disease OR periodontitis OR gingivitis) AND (preterm labor OR preterm birth OR premature rupture of membranes OR low birthweight OR PTB OR PROM OR LBW).” They compared the results and reached a consensus on the eligibility of the trials with the involvement of a third investigator (IPP). In addition, we reviewed the references of all eligible trials, did cross searches in Medline by using the names of the investigators who were lead authors on at least one eligible trial, and hand searched the last two year volumes of two key dentistry journals (Journal of Periodontology and Journal of Clinical Periodontology).

All randomised controlled trials that allocated pregnant women to receive treatment with scaling and root planning were included, as well as studies that assessed the natural history of pregnancy in women with periodontal disease. Excluded were reviews, editorials, letters, not relevant studies, and trials that did not follow patients. Other exclusions were trials that did not randomise women to receive treatment with scaling and root planning, those that did not have a control group, and those that did not use a validated clinical or radiographic periodontal index. Eligible randomised controlled trials were included. The Cochrane Central Trials Registry, ISI Web of Science, and Medline without language restriction were searched for retrieved dates up to July 2010. We included trials that enrolled pregnant women, and excluded studies that did not enrol patients with threatened preterm birth receiving tocolytic agents. Eligible randomised controlled trials were included. Where possible, we attempted to contact authors for additional data and information.

**Fig 1| Flow chart of selection of trials**

**Table 1| Baseline characteristics of included trials**

| Author, year | Country | Total No of patients | No of patients followed | No of live births | Gestational age at enrolment (weeks) | Gestational age at completion of treatment (weeks) | Definition of periodontal disease | Reclassification of severity of disease inclusion criteria according to CDC AAP 2003* |
|--------------|---------|----------------------|------------------------|------------------|------------------------------------|-----------------------------------------------|----------------------------------|----------------------------------|
| Lopez, 200221 | Chile   | 400                  | 372                    | 358              | 9-21                               | 28                                            | ≥4 teeth with ≥1 sites with PD≥4 mm and CAL≥2 mm | Mild periodontitis               |
| Jeffcoat, 200325† | USA     | 246                  | 246                    | 246              | 21-25                              | NA                                           | ≥3 sites with CAL≥3 mm            | Mild periodontitis               |
| Lopez, 200522 | Chile   | 870                  | 856                    | 845              | ≤22                                | 28                                            | BOP>25% of sites and no sites with CAL≥2 mm | Gingivitis                      |
| Michalowicz, 200623† | USA       | 823                  | 812                    | 793              | 13-17                              | Until delivery when necessary                 | 4 or more teeth with PD≥4 mm and CAL≥2 mm and BOP≥35% of sites | Mild periodontitis               |
| Offenbacher, 200623† | USA     | 74                   | 67                     | 67               | ≤22                                | NA                                           | ≥2 sites with PD≥5 mm and CAL 1-2 mm at ≥1 sites with PD≥5 mm | Moderate periodontitis           |
| Sadramansuri, 200624 | Iran      | 30                   | 30                     | 30               | 13-20                              | 30                                            | ≥4 teeth with ≥1 sites with PD≥4 mm and CAL≥3 mm | Mild periodontitis               |
| Taranum, 200724 | India    | 220                  | 192                    | 188              | 9-21                               | 28                                            | CAL≥2 mm at ≥50% of examined sites | Mild periodontitis               |
| Offenbacher, 2009† | USA      | 1806                 | 1760                   | 1761             | ≤24                                | NA                                           | ≥3 periostal sites with CAL≥3 | Mild periodontitis               |
| Newham, 200923† | Australia | 1087                | 1080                   | 1073             | 12-20                              | 28                                            | ≥12 probing sites with PD≥4 | Mild periodontitis               |
| Macones, 2010† | USA      | 756                  | 713                    | 720              | 6-20                               | NA                                           | CAL≥3 mm on ≥3 teeth             | Mild periodontitis               |
| Oliveira, 201019 | Brazil   | 246                  | 239                    | 233              | 12-20                              | 30-32                                         | ≥1 sites with PD≥4 mm and CAL≥3 mm | Mild periodontitis               |

BOP—bleeding on probing site; CAL—clinical attachment loss; NA—no data available; PD—probing depth.

*Mild periodontitis refers to category neither moderate nor severe periodontitis according to Centres for Disease Control and Prevention and American Association of Periodontology 2003 criteria; gingivitis is not included in clinical case definitions by CDC AAP 2003 but is in accordance with term proposed by International Workshop for a Classification of Periodontal Diseases and Conditions in 1999.

†Trials considered to be of high methodological quality.
planning versus no treatment or prophylaxis were eligible for inclusion. We considered trials to be eligible if they included patients with documented periodontal disease (periodontitis or gingivitis), as defined by the International Workshop for a Classification of Periodontal Diseases and Conditions. All trials were eligible regardless of the depth and the severity of periodontal disease. We adopted a further classification of the severity of periodontal disease based on the conclusions of the working group by the Centers for Disease Control and Prevention and the American Association of Periodontology in 2003. We defined moderate and severe periodontitis, according to this classification, in terms of probing depth and clinical attachment loss to enhance case definitions and provide distinct categories.

For trials that, according to their protocol, had included arms in which patients received concomitant treatment (such as antibiotics), we focused on the subgroups of eligible patients. We excluded from the analysis randomised trials that included patients with threatened preterm delivery who received tocolytic agents, non-randomised trials, and pseudo-randomised trials.

Data extraction and assessment of methodological quality
Two independent investigators (AZ and IPP) were involved in the data extraction. A third investigator (NPP) examined the results, and a consensus was reached. We extracted the following data from each arm of the eligible trials: authors’ names, journal and year of publication, country of origin, enrolment years, gestational age at enrolment, gestational age at completion of treatment, number of patients randomised and eligible, number of live births, and patients’ inclusion criteria. In addition, we recorded the methodological quality of the trials by using Cochrane’s risk of bias tool. Two independent investigators (NPP and DM) assessed the methodological quality of the trials, and consensus was reached.

The primary outcome was the rate of preterm births, defined as the number of preterm births before 37 weeks of gestation (spontaneous or indicated) among all successful pregnancies (all randomised trials). The data for this outcome are presented in the table below.
We included all trials in the final analysis. Because of the increased heterogeneity seen after we pooled the data from all the trials, we did separate meta-analyses including only high quality or low quality trials. We considered high quality trials to be those that had a low risk of bias as was assessed by Cochrane’s risk of bias tool. We detected the possibility of publication bias visually by using contour funnel plots and tested for small study effect bias with Harbord’s modified test.

We used Review Manager (RevMan) version 5 statistical software to analyse data. All P values were two tailed with a level of significance of <0.05. We used Stata SE 10.0 to do contour enhanced funnel plots and Harbord’s test for small study effect bias.

RESULTS
Characteristics of eligible trials
We retrieved 613 reports through searches in the Cochrane Central Trials Registry, ISI Web of Science, and Medline. We did not identify any additional trials through additional searches. We retrieved 12 randomised controlled trials. However, we considered one randomised trial to be ineligible because randomised patients had been admitted to hospital for threatened preterm birth and received tocolytic agents. Another recent trial did not clarify whether all patients in the intervention arm received treatment with scaling and root planing; however, personal contact with the primary investigators of the trial clarified that all patients had received such treatment. Finally, we considered 11 trials to be eligible. These trials included 6558 patients —3438 allocated to periodontal disease treatment and 3120 allocated to control groups.

| Study or subgroup | Treatment Events/total | No treatment Events/total | Odds ratio (M-H, fixed) (95% CI) | Weight (%) | Odds ratio (M-H, fixed) (95% CI) |
|-------------------|-----------------------|--------------------------|---------------------------------|------------|---------------------------------|
| **Low quality trials** |                       |                          |                                 |            |                                 |
| Lopez 2002        | 1/168                  | 7/190                    | 3.1 (0.2 to 1.29)               |            |                                 |
| Lopez 2005        | 4/563                  | 3/282                    | 1.9 (0.15 to 2.99)              |            |                                 |
| Sadatmansuri 2006 | 0/15                   | 1/15                     | 0.7 (0.01 to 8.28)              |            |                                 |
| Taranum 2007      | 26/99                  | 48/89                    | 17.9 (0.17 to 0.56)             |            |                                 |
| Oliveira 2010     | 23/116                 | 31/118                   | 11.8 (0.38 to 1.28)             |            |                                 |
| **Subtotal (95% CI)** | 54/961                | 90/694                   | 35.4 (0.30 to 0.66)             |            |                                 |
| **High quality trials** |                   |                          |                                 |            |                                 |
| Michalowicz 2006  | 40/402                 | 43/391                   | 18.8 (0.57 to 1.41)             |            |                                 |
| Offenbacher 2009  | 72/881                 | 71/880                   | 31.3 (1.07 to 1.43)             |            |                                 |
| Macones 2010      | 48/359                 | 35/361                   | 14.5 (0.91 to 2.28)             |            |                                 |
| **Subtotal (95% CI)** | 160/1642              | 149/1632                 | 64.6 (0.85 to 1.36)             |            |                                 |
| **Total (95% CI)** | 214/2603               | 239/2326                 | 100.0 (0.70 to 1.04)            |            |                                 |

Fig 4 | Meta-analysis plot for low birthweight infants (<2500 g). M-H=Mantel-Haenszel model.

pregnancies except patients lost to follow-up and pregnancies that led to spontaneous abortions or stillbirths. Secondary outcomes included the rate of low birthweight infants, defined as the number of infants under 2500 g among all successful pregnancies; the rate of spontaneous abortions/stillbirths, defined as the number of spontaneous abortions or stillbirths among all patients randomised except those lost to follow-up; and the overall rate of adverse outcomes of pregnancy, defined as the number of preterm births (<37 weeks) and the number of spontaneous abortions and stillbirths, in an intention to treat analysis among all randomised patients. Finally, other secondary outcomes included the rate of spontaneous preterm births, rate of preterm birth before 35 weeks of gestation, and rate of very low birthweight infants (<1500 g).

Data analysis
We constructed two by two tables and calculated the odds ratio for each primary study to estimate the relative risk of preterm birth, low birthweight infants, and spontaneous abortion/stillbirth among the treatment group compared with the control group. To test the homogeneity of the estimates of odds ratios among eligible studies, we used the $\chi^2$ test with a level of significance of 0.1, and we further quantified the degree of heterogeneity by using the I^2 test. We synthesised data across studies by using the fixed effects (Mantel-Haenszel) model whenever no statistical heterogeneity was apparent, or by using the random effects model (DerSimonian and Laird). Whenever studies reported zero events in both arms (treatment and no treatment), we excluded these trials from the final analysis.

RESULTS
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We included all trials in the final analysis. Because of the increased heterogeneity seen after we pooled the data from all the trials, we did separate meta-analyses including only high quality or low quality trials. We considered high quality trials to be those that had a low risk of bias as was assessed by Cochrane’s risk of bias tool. We detected the possibility of publication bias visually by using contour funnel plots and tested for small study effect bias with Harbord’s modified test.

We used Review Manager (RevMan) version 5 statistical software to analyse data. All P values were two tailed with a level of significance of <0.05. We used Stata SE 10.0 to do contour enhanced funnel plots and Harbord’s test for small study effect bias.
3120 allocated to no treatment or placebo (fig 1). The table shows the baseline characteristics of eligible trials.

### Methodological quality of trials

We used Cochrane’s risk of bias tool to assess the methodological quality of the trials (fig 2). Among eligible trials, eight reported an adequate randomisation mode,1 5 12 20-24 four used an adequate mode of allocation concealment,1 5 12 20-23, and six used blinding.1 5 12 20-25 Overall, we considered five trials to be of high methodological quality,1 5 12 20-23 with a low risk of bias. One study had an unclear risk of bias, whereas the five remaining studies had a high risk of bias.

### Primary outcome—preterm birth (<37 weeks)

Preterm births were reported in all of the eligible trials. Overall, 364 preterm births were reported in women receiving treatment compared with 366 in patients who received no treatment. Meta-analysis of the overall preterm birth rate indicated no difference when we used either the random effects model (odds ratio 0.79, 95% confidence interval 0.58 to 1.06; P=0.12) or the fixed effects model (0.93, 0.79 to 1.10; P=0.39). Subgroup analysis according to the methodological quality of randomised studies showed diverse results. Whereas low quality trials supported a significant beneficial effect of treatment with scaling and root planing on the rate of preterm birth (odds ratio 0.52, 0.38 to 0.72; P=0.0001), no such effect was apparent among high quality studies (1.15, 0.95 to 1.40; P=0.15). Heterogeneity was present when we pooled all the trials together but not for the separate analyses of high quality or low quality trials (fig 3).

### Secondary outcomes

#### Low birthweight infants (<2500 g)

The number of low birthweight infants was reported in eight of the eligible trials. Meta-analysis found no significant difference between compared arms (odds ratios 0.85, 0.70 to 1.04 [P=0.11] with fixed effects model and 0.74, 0.49 to 1.12 [P=0.16] with random effects model). Low quality and high quality trials again differed in the effect of treatment (odds ratios 0.44, 0.30 to 0.66 [P<0.0001] for low quality trials and 1.07, 0.85 to 1.36 [P=0.55] for high quality trials), whereas heterogeneity was significant when we included all the trials (Q=20.23, P=0.005, I²=65%) but not for either high quality or low quality studies separately (fig 4).

#### Spontaneous abortion/stillbirth

All trials reported data for spontaneous abortions or stillbirths. Three trials reported no outcomes in both treatment and control arms,20 25 26 so we excluded these from the final analysis. The pooled odds ratio for the rate of spontaneous abortion/stillbirth was 0.84 (0.58 to 1.22; P=0.37), suggesting that no significant difference exists. Results remained non-significant
This meta-analysis shows that treatment of periodontitis with scaling and root planing did not significantly improve the rate of spontaneous preterm birth (<37 weeks) (odds ratio 0.66, 0.37 to 1.17; P=0.12), preterm birth <35 weeks (1.22, 0.88 to 1.68; P=0.23), or very low birthweight infants (<1500 g) (0.99, 0.61 to 1.60; P=0.97) (fig 7).

**Publication bias**

We assessed the presence of publication bias by using the contour enhanced funnel plots.

**DISCUSSION**

This meta-analysis shows that treatment of periodontitis with scaling and root planing in pregnant women has no significant effect on the incidence of preterm birth. Furthermore, treatment does not seem to have a significant effect on the incidence of low birthweight infants or spontaneous abortions/stillbirths or on the overall rate of adverse outcomes of pregnancy (preterm births and spontaneous abortions/stillbirths).

**Comparison with other studies**

The results of this meta-analysis are in contrast to those of a previous meta-analysis published in 2009. A
potential reason for this discrepancy may be the fact that the trials included in the earlier study had considerable methodological shortcomings. Today, after the publication of three new, well designed, large randomised trials,152 treatment of periodontal disease during pregnancy does not seem to offer any clear benefit for the reduction of preterm births or low birthweight infants and therefore should not be routinely recommended in pregnant women as a measure for prevention of preterm birth.

Another reason for the discrepancy between this meta-analysis and the previous one may be the potential threat of publication bias. Considerable evidence from many clinical domains indicates that trials with “negative” results, especially small ones, may have difficulty getting published or may be published with considerable delays compared with trials that find significant benefits for the tested interventions28 29; the odds of publication are approximately four times greater if the results are statistically significant.30 Thus, early trials that did not favour treatment of periodontal with scaling and root planing during pregnancy may have been left unpublished, resulting in the accumulation of small trials with significant results in favour of treatment. According to the funnel plot constructed, publication bias is highly likely to exist.

| Study or subgroup | Treatment Events/total | No treatment Events/total | Odds ratio (M-H, fixed) (95% CI) | Weight (%) | Odds ratio (M-H, fixed) (95% CI) |
|-------------------|-----------------------|---------------------------|---------------------------------|------------|---------------------------------|
| **Spontaneous preterm birth <37 weeks** | | | | | |
| Low quality trials | | | | | |
| Lopez 2002 | 2/168 | 12/190 | 9.8 | 0.18 | (0.04 to 0.81) |
| Lopez 2005 | 8/563 | 16/282 | 18.1 | 0.24 | (0.10 to 0.57) |
| Oliveira 2010 | 24/116 | 26/117 | 22.4 | 0.91 | (0.49 to 1.71) |
| Subtotal (95% CI) | 34/847 | 54/589 | 50.3 | 0.38 | (0.13 to 1.13) |
| Test for heterogeneity: $\tau^2=0.68$, $\chi^2=8.18$, df=2, $P=0.02$, $I^2=75\%$ | | | | | |
| Test for overall effect: $z=1.74$, $P=0.08$ | | | | | |
| High quality trials | | | | | |
| Michalowicz 2006 | 33/402 | 30/391 | 24.5 | 1.08 | (0.64 to 1.80) |
| Macones 2010 | 38/359 | 37/361 | 25.2 | 1.04 | (0.64 to 1.67) |
| Subtotal (95% CI) | 71/761 | 67/752 | 49.7 | 1.05 | (0.74 to 1.50) |
| Test for heterogeneity: $\tau^2=0.00$, $\chi^2=0.01$, df=1, $P=0.92$, $I^2=0\%$ | | | | | |
| Test for overall effect: $z=0.30$, $P=0.77$ | | | | | |
| **Total (95% CI)** | 105/1608 | 121/1341 | 100.0 | 0.66 | (0.37 to 1.17) |
| Test for heterogeneity: $\chi^2=28.2$, $\tau^2=13.90$, df=4, $P=0.008$, $I^2=71\%$ | | | | | |
| Test for overall effect: $z=1.43$, $P=0.15$ | | | | | |
| **Preterm birth <35 weeks** | | | | | |
| High quality trials | | | | | |
| Jeffcoat 2003 | 1/123 | 6/123 | 8.8 | 0.16 | (0.02 to 1.35) |
| Michalowicz 2006 | 18/402 | 12/391 | 17.2 | 1.48 | (0.70 to 3.12) |
| Offenbacher 2009 | 36/881 | 33/880 | 46.9 | 1.09 | (0.68 to 1.77) |
| Macones 2010 | 31/359 | 20/361 | 27.0 | 1.61 | (0.90 to 2.88) |
| **Total (95% CI)** | 86/1765 | 71/1755 | 100.0 | 1.22 | (0.88 to 1.68) |
| Test for heterogeneity: $\chi^2=4.83$, df=3, $P=0.18$, $I^2=38\%$ | | | | | |
| Test for overall effect: $z=1.20$, $P=0.23$ | | | | | |
| Test for subgroup differences: Not applicable | | | | | |
| **Low birth weight <1500 g** | | | | | |
| High quality trials | | | | | |
| Michalowicz 2006 | 8/402 | 15/391 | 44.5 | 0.51 | (0.21 to 1.21) |
| Offenbacher 2009 | 15/881 | 13/880 | 38.2 | 1.16 | (0.55 to 2.44) |
| Macones 2010 | 11/359 | 6/361 | 17.3 | 1.87 | (0.68 to 5.11) |
| **Total (95% CI)** | 34/1642 | 34/1632 | 100.0 | 0.99 | (0.61 to 1.60) |
| Test for heterogeneity: $\chi^2=5.95$, df=2, $P=0.14$, $I^2=49\%$ | | | | | |
| Test for overall effect: $z=0.04$, $P=0.97$ | | | | | |
Furthermore, the application of Harbord’s modified test for detecting small study effect bias showed that small studies were more likely to provide inflated outcomes in favour of the treatment arm. Despite the fact that recent evidence suggests that in most meta-analyses the application of funnel plot asymmetry tests to detect publication bias is inappropriate or not meaningful, especially when few trials are included in the meta-analysis, we applied the contour enhanced funnel plots, which seem to be more convincing in detecting publication bias.

**Strengths and limitations of study**

The most important strength of this study is the large sample size of patients included among high quality trials. Despite the fact that less than half of the trials were of high methodological quality, they cumulatively included more than 4500 patients, with a 65% weight in the overall estimate of the pooled study effect. This constitutes a large sample size, sufficiently powered to exclude at least a 2.5% reduction in the incidence of preterm birth after treatment with scaling and root planing, given that the preterm birth rate in the United States is around 12%.32

Another advantage of this meta-analysis is that we separately analysed our results according to the methodological quality of the studies. The most interesting observation is the completely different results obtained when we pooled the data separately from low quality and high quality trials for most of the primary and secondary outcomes. Strong statistical significance was achieved when we included only low quality trials, and this significance completely disappeared when we considered only high quality trials. One should be cautious when interpreting the data from low quality randomised trials and possibly reconsider the adoption of outcomes retrieved from such trials. Considering that only two of the early trials published up to 2008 reported key methodological parameters for randomised trials, the discrepancy between the results of our previous analysis and this one should be undeniably attributed to the suboptimal quality of the early trials. The subgroups analyses we did for all the outcomes clearly support this hypothesis, as low quality trials showed a strong significant effect of treatment whereas high quality trials reported no difference in any of the outcomes tested.

Poor reporting of the methodological characteristics of randomised trials has been previously described. Although a substantial improvement in key aspects of trials’ methods has been seen over the past years, the quality of reporting remains well below an acceptable level.24 The most important aspect related to the suboptimal methodological quality of randomised trials is the substantial threat to the validity of the results and conclusions obtained. Reporting of methodological quality parameters is directly correlated with the provision of outcomes that favour the experimental arm.35 36 Previous reports have shown an inverse relation between the methodological quality score and the efficacy of preventive strategies, whereas the quality of reports of randomised trials seems to affect estimates of the efficacy of interventions reported in meta-analyses.28 Thus, taking into account the fact that studies of low methodological quality in which the estimate of quality is incorporated into the meta-analyses can alter the interpretation of the benefit of the intervention, this meta-analysis has a major strength in that it separately considered the effect of treatment among high quality and low quality trials. This is in accordance with a large meta-epidemiological study, which clearly recommends that systematic reviewers should present meta-analyses restricted to trials at low risk of bias for each outcome, either as the primary analysis or in conjunction with less restrictive analyses.29 Furthermore, the adoption of Cochrane’s risk of bias tool to assess the quality of the randomised trials included in our review is another advantage, given that it is a validated tool developed to overcome some of the shortcomings of existing quality assessment instruments.

Our meta-analysis has certain limitations. Firstly, the definition of periodontal disease differed among the trials included. A recent position paper commissioned by the European Association of Dental Public Health showed a distinct lack of consensus and uniformity in the definition of periodontitis in epidemiological studies.41 Furthermore, a review showed that the significance of the association between periodontal disease and pregnancy outcomes may be determined by the definition or measurement of periodontal disease used.42 This discrepancy in the definition of periodontal disease is also present in the trials of treatment of periodontal disease during pregnancy. However, we re-evaluated the severity of periodontal disease among patients included within the trials, on the basis on the classification proposed by the Centres for Disease Control and Prevention and the American Association of Periodontology in 2003.15 On the basis of trials’ inclusion criteria, the five high quality trials had comparable inclusion criteria as regards the severity of disease (table). Consequently, differences in the definition of the disease are unlikely to have contributed to the lack of effect of treatment.

Secondly, one of the trials included in the main analysis enrolled only patients with gingivitis who were treated with scaling.22 These patients might have had...
WHAT IS ALREADY KNOWN ON THIS TOPIC

Periodontal disease is associated with an increased risk of preterm birth, and a causal relation may exist.
Existing reports on the effect of treatment with scaling and root planning on the incidence of preterm birth are conflicting.

WHAT THIS STUDY ADDS

Treatment of periodontal disease with scaling and root planning during pregnancy does not reduce the risk of preterm birth and should not be routinely recommended as a measure to prevent preterm birth.
Randomised trials of low methodological quality tend to overestimate the effect of treatment, whereas high quality trials provide strong evidence that no significant effect of treatment exists.

considerable improvement in their disease with treatment, as they were women with less severe periodontal disease and the effect of treatment on the rate of preterm birth may have been greater. However, this study had important methodological shortcomings and had a significantly higher percentage of patients with a previous history of preterm birth in the treatment arm, so we did not include it in the analysis of high quality studies. Furthermore, considering the positive results of the trial, exclusion from the main analysis would have not affected our results.

Finally, according to our tests, an increased likelihood of publication bias exists in our analysis. However, given that publication bias usually involves trials with “negative” results,49 any unpublished trial would be highly unlikely to have favoured the treatment arm. Furthermore, taking into account the quality of the five highest quality trials included here and their weight in the overall analysis, the effect of any unpublished or missed trial on the overall effect of treatment would probably be detrimental.

Conclusions and policy implications

In an attempt to explain the lack of a significant effect of treatment of periodontal disease during pregnancy on perinatal outcomes, we could speculate that treatment with scaling and root planning does not alter the sequence of events leading from regional inflammation due to periodontitis to systemic inflammation and onset of preterm birth. This lack of treatment effect may be related to the diverse systemic response after scaling and root planning on different stages of periodontal disease. For example, in non-pregnant patients with documented periodontal disease, non-surgical treatment results in different local and systemic inflammatory responses.43 Furthermore, in more severe cases, treatment may lead to higher levels of circulating inflammatory cytokines.7,44 A subgroup analysis in our previous report showed that treatment was significantly more effective when applied in patients with less severe disease. However, our updated meta-analysis now includes five high quality trials, among which the definition of periodontal disease does not differ widely. Thus, taking into account the fact that none of the outcomes among these trials favoured treatment with scaling and root planning during pregnancy, such treatment is unlikely to have any beneficial effect on perinatal outcomes.

Some investigators suggest that adjuvant antibiotic treatment could be used in patients with periodontal disease to prevent preterm birth. However, the beneficial effect of antibiotic treatment in reducing the incidence of preterm birth, either among patients with periodontal disease or among those with other infections, is controversial. The only randomised trial that allocated patients to a combination of scaling and root planning and administration of antibiotic failed to support this hypothesis,20 and previous meta-analyses failed to show a significant effect of antibiotics on the incidence of preterm birth when given for the treatment of bacterial vaginosis.8-11

Another consideration is that the success of periodontal disease treatment may indeed be a determinant of the effect of scaling and root planning on the overall incidence of preterm birth. Some randomised trials included in our meta-analysis examined periodontal status only before delivery and not after treatment. Consequently, a well designed prospective randomised trial to test whether pregnant patients who have successfully controlled their periodontal disease may eventually have a substantially reduced incidence of preterm birth would be of interest.

Finally, despite the fact that we failed to show any benefit of treatment during pregnancy, we cannot exclude the possibility that women who start treatment and have their periodontal disease controlled early in the first trimester of pregnancy or even before becoming pregnant, may have a substantial improvement in perinatal outcomes. Although we cannot be sure that a causal relation between periodontal disease and preterm birth exists, the likelihood of an association is very high.4 Therefore, pre-pregnancy treatment seems to be a reasonable alternative, and a randomised trial including patients allocated to scaling and root planning or control before pregnancy and examining its effect in a subsequent pregnancy may show significant differences in the future. However, given that many pregnancies are unplanned, such a trial would be very difficult to do and would need a large sample size.

In conclusion, updated evidence does not encourage the use of scaling and root planning as an efficient method of reducing the rate of preterm birth or improving perinatal outcomes. Women may be advised to evaluate their dental status during pregnancy and may have treatment for periodontal disease. However, they should be told that such a treatment during pregnancy is unlikely to reduce the risk of preterm birth or low birthweight infants and therefore should not be considered as routine antenatal care.

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