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Effects of acupuncture on rates of pregnancy and live birth among women undergoing in vitro fertilisation: systematic review and meta-analysis

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ABSTRACT
Objective To evaluate whether acupuncture improves rates of pregnancy and live birth when used as an adjuvant treatment to embryo transfer in women undergoing in vitro fertilisation.

Design Systematic review and meta-analysis.

Data sources Medline, Cochrane Central, Embase, Chinese Biomedical Database, hand searched abstracts, and reference lists.

Review methods Eligible studies were randomised controlled trials that compared needle acupuncture administered within one day of embryo transfer with sham acupuncture or no adjuvant treatment, with reported outcomes of at least one of clinical pregnancy, ongoing pregnancy, or live birth. Two reviewers independently agreed on eligibility; assessed methodological quality; and extracted outcome data. For all trials, investigators contributed additional data not included in the original publication (such as live births). Meta-analyses included all randomised patients.

Data synthesis Seven trials with 1366 women undergoing in vitro fertilisation were included in the meta-analyses. There was little clinical heterogeneity. Trials with sham acupuncture and no adjuvant treatment as controls were pooled for the primary analysis. Complementing the embryo transfer process with acupuncture was associated with significant and clinically relevant improvements in clinical pregnancy (odds ratio 1.65, 95% confidence interval 1.27 to 2.14; number needed to treat (NNT) 10 (7 to 17); seven trials), ongoing pregnancy (1.87, 1.40 to 2.49; NNT 6 (to 15); five trials), and live birth (1.91, 1.39 to 2.64; NNT 9 (6 to 17); four trials). Because we were unable to obtain outcome data on live births for three of the included trials, the pooled odds ratio for clinical pregnancy more accurately represents the true combined effect from these trials rather than the odds ratio for live birth. The results were robust to sensitivity analyses on study validity variables. A prespecified subgroup analysis restricted to the three trials with the higher rates of clinical pregnancy in the control group, however, suggested a smaller non-significant benefit of acupuncture (odds ratio 1.24, 0.86 to 1.77).

Conclusions Current preliminary evidence suggests that acupuncture given with embryo transfer improves rates of pregnancy and live birth among women undergoing in vitro fertilisation.

INTRODUCTION

Some 10-15% of couples have difficulty conceiving at some point in their reproductive lives and seek specialist fertility treatment.1 A commonly used option is in vitro fertilisation, which involves retrieving a woman’s egg, fertilising the egg in the laboratory, and then transferring the embryo back into the woman’s uterus through the cervix.2 This entire process is typically referred to as an in vitro fertilisation “cycle” because it involves several procedures, typically over the course of about two weeks, starting when a woman begins taking drugs to stimulate egg production. In 2003, over 120 000 treatment cycles were performed in clinics in the United States.2 In 2000, about 200 000 babies worldwide were conceived through in vitro fertilisation.3

Because each cycle is expensive, lengthy, and stressful, new drugs and technologies have been developed to improve success rates. Progress, however, has been limited. Although use of some procedures, initiated before the cycle, have been shown to improve pregnancy rates in women with a poorer prognosis because of specific conditions (such as surgical treatment for tubal disease,4 long term treatment with gonadotrophin releasing hormone agonists for women with endometriosis5), few adjuvant procedures have been shown to be effective for women in general. One exception is luteal phase support, which has been shown to increase pregnancy rates6 and is routinely used.

Acupuncture has been used in China for centuries to regulate the female reproductive system.7 Three potential mechanisms for its effects on fertility have been postulated.8 Firstly, acupuncture may mediate the release of neurotransmitters,9 which may in turn stimulate secretion of gonadotrophin releasing hormone, thereby influencing the menstrual cycle,
ovulation, and fertility. Instead, acupuncture may stimulate blood flow to the uterus by inhibiting uterine central sympathetic nerve activity. Thirdly, acupuncture may stimulate the production of endogenous opioids, which may inhibit the central nervous system outflow and the biological stress response.

We conducted a systematic review and meta-analysis of randomised controlled trials to determine whether acupuncture given with embryo transfer improves the rates of pregnancy and live birth among women undergoing in vitro fertilisation.

METHODS

Identification of studies

We searched the computerised databases Medline, Embase, Cochrane Central, and the Chinese Biomedical Database from inception to January 2007. We searched the following terms as free text terms and MeSH terms (shown in italics): (acupuncture, acupuncture therapy, auriculotherapy; electroacupuncture; Medicine, Oriental Traditional; Medicine, Chinese Traditional; moxibustion) and (reproductive techniques, assisted, fertilization in vitro, embryo transfer; oocytes; egg collection). We combined this search strategy with a methods filter for clinical trials.

We also searched the proceedings of three major annual conferences on assisted reproduction technology for 2001-6: the American Society for Reproductive Medicine, the European Society for Human Reproduction and Embryology, and the Pacific Coast Reproductive Society. We also scanned reference lists of relevant publications.

Selection criteria, data extraction, and quality assessment

We selected randomised controlled trials that compared acupuncture with sham acupuncture or no adjuvant treatment. Because we were evaluating acupuncture as a complement to embryo transfer, we considered only trials in which acupuncture was administered within one day of the procedure, with the objective of improving success rates. Trials that included intracytoplasmic injection of sperm as well as in vitro fertilisation were eligible. We excluded trials that evaluated acupuncture as an alternative to conventional analgesia for egg removal.

For trials to be eligible, we had to be able to extract data on at least one of the following outcomes, as recommended: (1) clinical pregnancy (that is, presence of at least one gestational sac or fetal heartbeat, confirmed by transvaginal ultrasound); (2) ongoing pregnancy (that is, pregnancy beyond 12 weeks of gestation, as confirmed by fetal heart activity on ultrasound), or live birth. We included cross over trials only if relevant outcome data from the first phase (that is, before the cross over occurred) were available. We ignored any data after cross over occurred.

We included only trials in which acupuncture involved the insertion of needles into traditional meridian points. The needles could be inserted into tender points in addition to the traditional meridian points, and the needles could also be electrically stimulated. We excluded trials of dry needling or trigger point therapy. We also excluded trials of laser acupuncture and electro-acupuncture without needle insertion because most authorities believe acupuncture involves needle insertion.

We imposed no restrictions on publication type (that is, either full article or abstract) or language of publication.

Two authors (EM and GZ) independently selected articles and extracted data, with disagreements resolved by discussion. We extracted data pertaining to quality of the methods, participants, interventions, and outcomes. Methodological quality of the trials was evaluated with the internal and external validity criteria from the checklist created by the Cochrane menstrual disorders and subfertility group. We contacted corresponding authors with specific questions related to the design and outcomes of their trials and asked them to review the information we extracted from their trials and clarify any ambiguities.

Data synthesis and analysis

The measure of treatment effect was the pooled odds ratio of achieving a clinical pregnancy, ongoing pregnancy, or live birth for women in the acupuncture group compared with women in the control group. The odds ratio is the measure of choice for outcomes in subfertility trials because this measure is more likely to show homogeneity when control rates differ between trials. To allow for a more clinically relevant interpretation, however, we also calculated pooled rate differences between the acupuncture and control groups and converted these rate differences to numbers needed to treat. For our meta-analyses, we used a random effects model because of the expected heterogeneity of the studies’ protocols and settings.

All meta-analyses were based on the number of women randomised (rather than on the number of treatment attempts—that is, cycles of in vitro fertilisation) with the intention to treat approach to analysis. We included randomised women who began the in vitro fertilisation process but did not complete the treatment (that is, had no embryo transfer) in the intention to treat meta-analyses. Although inclusion of women without an embryo transfer will tend to underestimate the effect of acupuncture, it is the more conservative and appropriate analytical approach because it preserves the groups created by the randomisation and reduces the chance of a type I error. All trials reported pregnancy outcomes resulting from a single cycle.

Subgroup analyses

To assess whether treatment effects varied with internal validity of studies we performed six subgroup analyses that evaluated whether any pooled results that were significant in analyses of all the trials remained significant when we restricted them to trials judged adequate on each of six internal validity components most commonly used for evaluating quality of randomised trials. These components, each evaluated with
a separate subgroup analysis (prespecified, except where indicated), were concealment of allocation of randomisation sequence, blinding of patients, blinding of physicians (post hoc analysis), loss to follow-up (withdrawal rate above or below 10% of the study population), intention to treat analysis, and generation of allocation sequence.

We evaluated heterogeneity using the I² test, which indicates the proportion of variability across trials not explained by chance alone. If the overall I² value for all trials was reduced when we separated the trials into subgroups according to control with sham acupuncture and no adjuvant treatment, then we would use the subgroup results as primary. Otherwise the pooled results from all trials would be used for our primary analysis, but with the results from the two subgroups also presented. A priori, we would not expect important heterogeneity of results based on whether or not a sham acupuncture control group was used because all outcomes are entirely objective (that is, pregnancy and births), and unlikely to be largely affected by expectations and placebo effects.23-25

We assessed whether effects of acupuncture varied with three clinical characteristics that might influence success (prespecified, except where indicated): use of extra acupuncture sessions in addition to the sessions before and after the embryo transfer [yes/no] (post hoc analysis); eligibility restricted to women with good quality embryos [yes/no]; and low versus high rates of clinical pregnancy in control groups [28% or more, the European average26]. For the subgroup analyses on each of the dichotomised methodological and clinical variables, we also used a significance test, as described by Deeks et al.19,27 to investigate whether differences in effects of acupuncture between the two subgroups were significant. Subgroup analyses on various clinical characteristics related to the patient (age, cause of diagnosis, duration of infertility, number of previous attempts) were not possible because the trials included typical heterogeneous populations. Other factors (such as electrical stimulation and intracytoplasmic sperm injection) also could not be examined in subgroup analyses because there was little or no heterogeneity across the trials on these factors.

As a sensitivity analysis, we evaluated whether the overall conclusions were affected if we excluded reports published only as abstracts. In addition, in one trial group assignment was unavailable for four randomised patients, none of whom completed the treatment [that is, no embryo transfer]. For our main analysis, we allocated two to the acupuncture group and two to the no adjuvant treatment group. In a post hoc sensitivity analyses, we assumed all four were randomised to the acupuncture group.

RESULTS

Figure 1 shows details of the selection process. Seven randomised controlled trials with a total of 1366 participants met inclusion criteria (table). All trials were published in English since 2002, and conducted in four different Western countries. Four were published as full reports2,6-7 and three as abstracts,8-10,14 in two of the leading reproductive medicine journals. We obtained unpublished methodological information for all seven and unpublished outcome data on live births for three. All seven trials used a pragmatic design,17 including typical clinical populations and using typical interventions before and after randomisation. All included a broad selection of women undergoing in vitro fertilisation, with a wide range of ages, diagnostic categories of infertility, durations of infertility, and numbers of previous treatment cycles. The only difference in the inclusion criteria was that two trials included only women with good quality embryos whereas the five others included women with embryos of varying quality. All trials reported use of intracytoplasmic sperm injection for some women.

The timing of the acupuncture sessions relative to embryo transfer differed somewhat among trials (table). In all trials, however, women received acupuncture immediately before or immediately after the embryo transfer.

![Fig 1 Flow of studies through selection process](image_url)
In all trials, the acupuncture protocol and selection of acupuncture points was designed for the sole purpose of improving rates of pregnancy. All trials used a fixed selection of acupuncture points for all patients for the sessions before and after embryo transfer. The fixed selection of points for these sessions was similar in all but one trial,\(^4\) and the points selected were largely based on the points selected in the first published trial that evaluated acupuncture as an adjuvant to embryo transfer.\(^4\) Three trials also included one extra acupuncture session, in addition to the sessions before and after embryo transfer. The fixed selection of points for all patients for the acupuncture group was designed for the sole purpose of improving rates of pregnancy. All trials used a fixed selection of acupuncture points for all patients for the sessions before and after embryo transfer. The fixed selection of points for these sessions was similar in all but one trial,\(^4\) and the points selected were largely based on the points selected in the first published trial that evaluated acupuncture as an adjuvant to embryo transfer.\(^4\) Three trials also included one extra acupuncture session, in addition to the sessions before and after embryo transfer.\(^4\) Six trials used ear acupuncture as a supplement to body acupuncture,\(^1-6\) and one of these stimulated the true and sham ear acupuncture points using a Chinese herb rather than a needle.\(^2\) In all the trials, the acupuncture sessions lasted 25-30 minutes. Five trials reported that the "de qi" needling sensation was sought,\(^2\) whereas the two others did not report on de qi.\(^1\) No trial used electro-acupuncture.

For all trials, there were no significant differences between the randomised groups in the mean numbers of embryos transferred.

Methodological quality of included studies

The trials generally had high internal validity, in terms of randomisation procedures (table) and follow-up of participants. For all trials but two,\(^2\) investigators confirmed no losses to follow-up (which is usual for in vitro fertilisation cycles\(^8\)). For one of the two trials with drop outs,\(^4\) these were limited to two women in the sham group with ongoing pregnancies, both of whom we assumed to have had live births.\(^1\) In two other trials,\(^6\) some randomised women began the in vitro fertilisation process but did not complete the treatment (that is, no embryo transfer); however, as noted, these women were still included in the meta-analyses.

Three of the trials used a sham acupuncture control,\(^2,6\) with one trial\(^2\) using needles that penetrated the skin at acupuncture points selected not to influence fertility\(^29,30\) and two\(^6\) using non-penetrating sham needles. For the four other trials,\(^1,4,6,7\) women in the control group received no adjuvant treatment (table).

Six trials\(^1,4,6,7\) reported their source of funding: three were funded by the in vitro fertilisation clinic,\(^1,4,7\) one by government support,\(^7\) and one by a company that manufactures fertility drugs.\(^6\) Four trials did not report on calculation of sample size.\(^1,4,6,7\)

Efficacy analysis

Our primary analysis is based on results from all included trials because dividing trials according to control group (sham acupuncture and no adjuvant care) increased, rather than reduced, heterogeneity. Embryo transfer with acupuncture was associated with a higher pooled odds for clinical pregnancy (1.65, 95% confidence interval 1.27 to 2.14), ongoing pregnancy

### Characteristics of included trials in meta-analysis of acupuncture on acupuncture and in vitro fertilisation

<table>
<thead>
<tr>
<th>Study</th>
<th>No randomised</th>
<th>Acupuncture sessions</th>
<th>Control*</th>
<th>Allocation sequence generation</th>
<th>Allocation concealment</th>
<th>Baseline comparability†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benson(^1)</td>
<td>103</td>
<td>Two sessions: one 25 minutes before transfer and one immediately after (n=53)(\dagger)</td>
<td>No adjuvant treatment (n=50)(\dagger)</td>
<td>Adequate</td>
<td>Inadequate</td>
<td>Age only</td>
</tr>
<tr>
<td>Dieterle(^2)</td>
<td>225</td>
<td>Two sessions: one immediately after transfer and one 3 days after (n=116)</td>
<td>Needle acupuncture, with needles inserted in real acupuncture points not expected to influence fertility (n=109)</td>
<td>Adequate</td>
<td>Adequate</td>
<td>Adequate</td>
</tr>
<tr>
<td>Domar(^3)</td>
<td>150</td>
<td>Two sessions: one 25 min before transfer and one immediately after (n=81)(\dagger)</td>
<td>No adjuvant treatment (women lay quietly for same time as those in that intervention group) (n=69)(\dagger)</td>
<td>Adequate</td>
<td>Adequate</td>
<td>Age only</td>
</tr>
<tr>
<td>Paulus(^4)</td>
<td>160</td>
<td>Two sessions: one 25 min before transfer and one immediately after (n=80)</td>
<td>No adjuvant treatment (n=80)</td>
<td>Adequate</td>
<td>Adequate</td>
<td>Adequate</td>
</tr>
<tr>
<td>Paulus 2003(^5)</td>
<td>200</td>
<td>Two sessions: one 25 min before transfer and one immediately after (n=100)</td>
<td>Sham needles on real acupuncture points did not penetrate skin (n=100)</td>
<td>Adequate</td>
<td>Adequate</td>
<td>Adequate</td>
</tr>
<tr>
<td>Smith 2006(^6)</td>
<td>228</td>
<td>Three sessions: day 9 of stimulating injections, 25 min before embryo transfer, and immediately after (n=110)</td>
<td>Sham needles placed close to, but not on, real acupuncture points did not penetrate skin, (n=118)</td>
<td>Adequate</td>
<td>Adequate</td>
<td>Adequate</td>
</tr>
<tr>
<td>Westergaard(^7)</td>
<td>300</td>
<td>Group 1: one session 25 min before transfer and one after (n=100); group 2: same as group one, but one additional acupuncture session two days after transfer (n=100)(\dagger)</td>
<td>No adjuvant treatment (n=100)</td>
<td>Adequate</td>
<td>Adequate††</td>
<td>Adequate</td>
</tr>
</tbody>
</table>

*For all sham controlled trials, sham acupuncture procedure was given on same schedule as that used in true acupuncture group.
††Trials judged to have adequate baseline comparability if groups created by randomisation were comparable at baseline on both age and prognostic factors related to conception. Two trials\(^1,3\) reported only that age of groups were comparable at baseline, as indicated; for one of these trials,\(^3\) this comparability was based on egg donor age.
\(\dagger\) In addition to needle acupuncture intervention group, this trial\(^4\) also included laser acupuncture intervention group (n=53); data from laser acupuncture group excluded because we considered only needle acupuncture as an eligible test intervention.
\(\dagger\) Two additional control groups (sham laser and relaxation) excluded from meta-analysis.
\(\dagger\) Of 150 patients randomised, group assignment was unavailable for four. For our main analysis, for which we assumed numbers randomised as shown in table, we allocated two to acupuncture group and two to no adjuvant treatment control.
\(\dagger\) Acupuncture groups one and two combined for meta-analysis.\(^5\)
††Randomised treatment assignments placed in sealed, opaque envelopes, which were shuffled and deposited in cardboard box, from which each patient selected only one. Procedure handled by independent nurse not responsible for obtaining information about patients and enrolling them. Although the envelopes were not sequentially numbered, we considered safeguards used in process to have provided adequate assurance of allocation concealment.
(1.87, 1.40 to 2.49), and live birth (1.91, 1.39 to 2.64) (fig 2). The pooled rate differences were 0.11 (0.06 to 0.16) for clinical pregnancy, 0.12 (0.07 to 0.17) for ongoing pregnancy, and 0.12 (0.06 to 0.18) for live birth. The numbers needed to treat (rounded up to the next whole number, as recommended31) were 10 (7 to 17) for clinical pregnancy, 9 (6 to 15) for ongoing pregnancy, and 9 (6 to 17) for live birth. For the clinical pregnancy outcome, I² values were 16% and 4% for the odds ratio and rate difference effect measures, respectively. All of the heterogeneity was caused by a single trial, w3 which reported only the clinical pregnancy outcome.

Of the nine subgroup analyses (seven prespecified) on clinical and methodological variables, only the subgroup analysis on the rates of clinical pregnancy in the control group showed a significant effect modification (P=0.04). Restriction to the three trials with the higher rates of clinical pregnancy in the control group suggested a smaller non-significant benefit of acupuncture (odds ratio 1.24, 0.86 to 1.77). No other subgroup restriction resulted in a change to a non-significant effect.

There were no significant adverse effects of acupuncture reported in the two trials that reported on this outcome, w2 w6

DISCUSSION

Although still somewhat preliminary, this review suggests that acupuncture given with embryo transfer improves rates of pregnancy and live birth among women undergoing in vitro fertilisation. The strengths of this review include the number of trials and their relatively large sample sizes; pooled odds ratios that are highly significant and clinically important; fairly consistent effect sizes across trials; homogeneity of the acupuncture protocols; use of objective and clinically relevant outcomes; adherence to the intention to treat approach for all meta-analyses; and overall high validity of the trials, as well as robustness of the results to sensitivity analyses on the effects of study validity variables. Because we obtained additional information for all of the trials, the review extends beyond the individual trial publications. For example, we included outcome data on live births from three trials, none of which had those data available when the studies were published.

Methodological strengths and limitations of included trials

The included trials generally had sound methods. Although some used suboptimal methods of concealment of allocation to treatment, blinding,29,30 and intention to treat analyses,28 the designs were adequate overall, and any minor design concerns would not be expected to result in a substantial risk of bias. In terms of randomisation, six out of the seven trials w2 w7 used an allocation procedure that would be considered as concealed.14,32 Four of these six trials w2 w3 w6 w7 however, concealed allocation by using sealed envelopes managed by clinical investigators. This is not ideal because of the greater potential for subversion and errors than use of off site treatment allocation.32

As for blinding, three trials w2 w6 w7 used a sham control and four w3 w4 w7 did not blind women to treatment assignment. The necessity to blind participants, however, is arguable when the outcomes are entirely objective (that is, pregnancy and birth), such as in these trials.24 It seems unlikely that a woman’s knowledge of whether or not she was receiving acupuncture would affect her ability to become pregnant. If the increases in pregnancies in this meta-analysis were due to expectation or placebo effects, then we would also have expected to see a smaller or no effect of acupuncture relative to sham treatment. The effects of acupuncture, however, were the same, regardless of whether acupuncture was compared with sham or no treatment. In addition, an indirect comparison of the proportions of pregnancies pooled from each of the two control arms did not indicate that women allocated to sham acupuncture were more likely to get pregnant compared with women in the no treatment control groups. This indirect comparison is of limited value, however, because it does not maintain the randomisation and therefore has all the limitations of observational data.

A recent methodological study compared the placebo effects of sham acupuncture with the placebo effects of oral pills and showed that patients receiving sham acupuncture can report significantly greater decreases in pain and severity of symptoms than patients receiving oral placebo pills.23 While this study found that acupuncture can have an enhanced placebo effect for subjective outcomes, it also found that the placebo effects (of both the sham acupuncture and oral placebo pills) were irrelevant for entirely objective outcomes. Irrelevance of shams for objective outcomes is also supported by recent meta-epidemiological data, which suggest that failure to double blind is associated with exaggerated treatment estimates, but only for trials with subjective outcomes.23 Finally, a systematic review has also shown that a placebo can result in a benefit over a “no treatment” control in studies with subjective outcomes such as pain, but that the placebo has no significant effect in studies with objective or binary outcomes.24

Blinding of physicians performing the embryo transfer is another potential source of bias (performance bias), and three of the seven included trials did not blind the physicians. w1 w4 w7 Considering the cost of embryo transfer and the importance of successful transfers to maintaining high pregnancy rates at clinics, we think that physicians would be motivated primarily to perform a successful procedure for all patients, rather than to show that acupuncture, a non-proprietary treatment, is an effective adjuvant procedure. In subfertility trials in general, where outcomes are entirely objective, blinding of either patients or physicians is “infrequently attempted.”25 Such blinding components are not always considered as critical elements related to the evaluation of risk of bias.25
For all trials but one, the data were reported in sufficient detail to allow us to conduct full intention to treat analyses for clinical pregnancies. For one other trial, intention to treat analyses were not used in the analysis for publication, but we could extract the relevant data for our meta-analysis. In this trial, the patients were randomised too early [at the point of oocyte retrieval], resulting in some women who did not undergo embryo transfer because they had a failure of fertilisation or poor embryo development. These women were excluded from the investigators’ analyses, but we included them in our meta-analysis.

**Limitations of the systematic review and meta-analysis**

Limitations of the meta-analysis include heterogeneity of baseline rates across trials, as well as the potential for publication and orientation biases. Substantial heterogeneity of baseline rates is indicated by the fact that the rates of pregnancy in the control group differed across trials by a factor of three. This heterogeneity is probably not caused by differential selection of patients across trials because each trial was pragmatic, including typical clinic patients with minimal inclusion or exclusion criteria applied. Also, the mean age of the women, an important predictive prognostic variable, was similar across trials, although one abstract did not report mean age. While distribution of other factors—such as diagnostic category of infertility, fertilisation procedure (in vitro fertilisation or intracytoplasmic sperm injection), and number of previous cycles—varied somewhat across trials, as expected by chance, such factors have not been shown to be strong predictors of pregnancy rates and none of these factors seemed to correlate well with the variability in baseline rate. The heterogeneity in baseline rate might more likely be explained by geographical differences in the nature of the typical clinic patients with minimal inclusion or exclusion criteria applied. As with any systematic review, we cannot exclude the possibility that publication bias might have affected our results. Although we conducted extensive searches to identify relevant studies and funnel plots did not suggest that there were small studies with negative results that were unpublished or not identified, we cannot rule out publication bias and this must be acknowledged as a potential limitation.

Another limitation is the possibility of “hypothesis” or “orientation” bias, which has been described as a type of interpretive bias in which the orientation or enthusiasm of the investigator for a treatment can be an unintentional determinant of a study’s results. The nature of this research might have increased the investigators’ enthusiasm or conviction of the benefits of the treatments investigated. On the other hand, a primary reason for orientation bias is widely acknowledged to be competing financial interests and as acupuncture is a non-proprietary treatment these might be reduced. We cannot exclude an effect, however, because if acupuncture were included as an adjuvant to in vitro fertilisation, clinics would be able to charge extra for acupuncture and thereby increase profits. While orientation bias cannot be excluded, the potential for such bias is also likely to be minimised in these trials because of the objective outcome measures and the methodologically sound study designs.

A final potential limitation is the inclusion of three studies presented only as abstracts, which may reflect premature analyses that were not peer reviewed. The corresponding authors, however, provided extensive additional information about their methods and results, beyond what was presented in the three abstracts, which increases our confidence in these studies.

**Clinical implications**

The odds ratio of 1.65 suggests that acupuncture increased the odds of clinical pregnancy by 65% compared with the control groups. It is important to note, however, that the odds ratio significantly overestimates the rate ratio in this context, in which the event (pregnancy) is relatively frequent. In absolute terms, the number needed to treat was 10, suggesting that 10 patients would need to be treated with acupuncture to bring about one additional clinical pregnancy. These are clinically relevant benefits. The subgroup analysis restricted to three trials with the higher pregnancy rates at baseline suggested a smaller non-significant benefit of acupuncture. A possible explanation for this non-significant finding is that in in vitro fertilisation settings, where the baseline pregnancy rates are already high, the relative added value of additional co-interventions, such as acupuncture, may be reduced. The dependency of the magnitude of the effect of acupuncture on the baseline pregnancy rate warrants further study.

Safety and costs are other considerations. Two large prospective surveys of practitioners show that serious adverse events after acupuncture are rare. Among women in labour and women at various stages of pregnancy, systematic reviews and randomised trials have shown acupuncture to be safe, although limited sample sizes preclude definitive conclusions. The effects of acupuncture in early pregnancy on complications later in pregnancy and on perinatal and infant outcomes have also been investigated in one trial, and no safety concerns were detected. In vitro fertilisation is an expensive procedure, costing an
WHAT IS ALREADY KNOWN ON THIS TOPIC

In vitro fertilisation is lengthy, expensive, and stressful. Safe, low cost, adjuvant treatments to improve success rates would benefit patients and reduce costs.

WHAT THIS STUDY ADDS

Current evidence from methodologically sound trials showed an odds ratio of more than 1.6 for clinical pregnancy after in vitro fertilisation with adjuvant acupuncture.

On average, 10 women would need to be treated with acupuncture to bring about one additional clinical pregnancy. The magnitude of this effect depended on the baseline pregnancy rate.

average of $12 400 (£6300, €8480) per cycle in the United States. If acupuncture increased the likelihood of success of an individual cycle, then the need for a subsequent cycle would be reduced, and overall costs would be decreased. Even if such increases were small, and, for example, 17 patients needed to be treated with acupuncture to bring about one additional pregnancy (that is, the lowest range of our 95% confidence interval), an acupuncture contervention may still be cost effective, considering the negligible costs of two to four sessions of acupuncture, relative to the high costs of in vitro fertilisation.

Conclusion and future research

Although current estimates of the effects of adjuvant acupuncture on in vitro fertilisation are significant and clinically relevant, they are still somewhat preliminary. Additional randomised trials are needed to quantify findings further and investigate the relation between baseline rate of pregnancy and the efficacy of adjuvant acupuncture. Future trials should adhere to CONSORT guidelines and guidelines developed specifically for subfertility trials and should evaluate an acupuncture protocol similar to that first reported by Paulus and colleagues, and suggested to have some beneficial effects, according to the trials included in this review.

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Contributors: EM and GZ determined inclusion eligibility of trials, extracted data from the trials, and assessed the methodological quality of the trials. AH provided a third assessment of the methodological quality of all the trials. EM entered and organised the data, conducted the analyses, and drafted the manuscript. EM, GZ, LU, AH, PL, BMB, and LMB all contributed to the conception and design. EM, GZ, LU, AH, PL, and LB contributed to the analysis and interpretation of data. EM, GZ, LU, AH, PL, BMB, and LMB critically revised the article for important intellectual content and approved the final version. EM is guarantor.

Fig 2  Effects of acupuncture on clinical pregnancy, ongoing pregnancy, and live birth outcomes
Competing interests: None declared.

Ethical approval: Not required.

Provenance and peer review: Not commissioned; externally peer reviewed.

29 Domar AD. Acupuncture and infertility: we need to stick to good science. Fertil Steril 2006;85:1359-61.