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Acupuncture for treatment of irritable bowel syndrome (Review)

Manheimer E, Cheng K, Wieland LS, Min LS, Shen X, Berman BM, Lao L

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[Intervention Review]

Acupuncture for treatment of irritable bowel syndrome

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ABSTRACT

Background

Irritable bowel syndrome (IBS) is a common, costly, and difficult to treat disorder that impairs health-related quality of life and work productivity. Evidence-based treatment guidelines have been unable to provide guidance on the effects of acupuncture for IBS because the only previous systematic review included only small, heterogeneous and methodologically unsound trials.

Objectives

The primary objectives were to assess the efficacy and safety of acupuncture for treating IBS.

Search methods

MEDLINE, the Cochrane Central Register of Controlled Trials, EMBASE, the Cumulative Index to Nursing and Allied Health, and the Chinese databases Sino-Med, CNKI, and VIP were searched through November 2011.

Selection criteria

Randomized controlled trials (RCTs) that compared acupuncture with sham acupuncture, other active treatments, or no (specific) treatment, and RCTs that evaluated acupuncture as an adjuvant to another treatment, in adults with IBS were included.

Data collection and analysis

Two authors independently assessed the risk of bias and extracted data. We extracted data for the outcomes overall IBS symptom severity and health-related quality of life. For dichotomous data (e.g. the IBS Adequate Relief Question), we calculated a pooled relative risk (RR) and 95% confidence interval (CI) for substantial improvement in symptom severity after treatment. For continuous data (e.g. the IBS Severity Scoring System), we calculated the standardized mean difference (SMD) and 95% CI in post-treatment scores between groups.

Main results

Seventeen RCTs (1806 participants) were included. Five RCTs compared acupuncture versus sham acupuncture. The risk of bias in these studies was low. We found no evidence of an improvement with acupuncture relative to sham (placebo) acupuncture for symptom severity (SMD -0.11, 95% CI -0.35 to 0.13; 4 RCTs; 281 patients) or quality of life (SMD = -0.03, 95% CI -0.27 to 0.22; 3 RCTs; 253 patients). Sensitivity analyses based on study quality did not change the results. A GRADE analysis indicated that the overall quality of the evidence for the primary outcomes in the sham controlled trials was moderate due to sparse data. The risk of bias in the four Chinese language comparative effectiveness trials that compared acupuncture with drug treatment was high due to lack of blinding. The risk of bias in the other studies that did not use a sham control was high due to lack of blinding or inadequate methods used for randomization and allocation concealment



or both. Acupuncture was significantly more effective than pharmacological therapy and no specific treatment. Eighty-four per cent of patients in the acupuncture group had improvement in symptom severity compared to 63% of patients in the pharmacological treatment group (RR 1.28, 95% CI 1.12 to 1.45; 5 studies, 449 patients). A GRADE analysis indicated that the overall quality of the evidence for this outcome was low due to a high risk of bias (no blinding) and sparse data. Sixty-three per cent of patients in the acupuncture group had improvement in symptom severity compared to 34% of patients in the no specific therapy group (RR 2.11, 95% CI 1.18 to 3.79; 2 studies, 181 patients). There was no statistically significant difference between acupuncture and Bifidobacterium (RR 1.07, 95% CI 0.90 to 1.27; 2 studies; 181 patients) or between acupuncture and psychotherapy (RR 1.05, 95% CI 0.87 to 1.26; 1 study; 100 patients). Acupuncture as an adjuvant to another Chinese medicine treatment was significantly better than the other treatment alone. Ninety-three per cent of patients in the adjuvant acupuncture group improved compared to 79% of patients who received Chinese medicine alone (RR 1.17, 95% CI 1.02 to 1.33; 4 studies; 466 patients). There was one adverse event (i.e. acupuncture syncope) associated with acupuncture in the 9 trials that reported this outcome, although relatively small sample sizes limit the usefulness of these safety data.

Authors' conclusions

Sham-controlled RCTs have found no benefits of acupuncture relative to a credible sham acupuncture control for IBS symptom severity or IBS-related quality of life. In comparative effectiveness Chinese trials, patients reported greater benefits from acupuncture than from two antispasmodic drugs (pinaverium bromide and trimebutine maleate), both of which have been shown to provide a modest benefit for IBS. Future trials may help clarify whether or not these reportedly greater benefits of acupuncture relative to pharmacological therapies are due entirely to patients' preferences for acupuncture or greater expectations of improvement on acupuncture relative to drug therapy.

PLAIN LANGUAGE SUMMARY

Acupuncture for irritable bowel syndrome

Irritable bowel syndrome (IBS) is a chronic gastrointestinal condition characterized by altered bowel habits and abdominal pain and discomfort. It is a common, costly, and difficult to treat disorder that also impairs health-related quality of life and work productivity. Some pharmacological (i.e. drug) therapies for treating IBS have modest benefits and a risk for side effects, and therefore, it is important to evaluate the effectiveness and safety of non-drug therapies, including acupuncture. One problem with trials in IBS is that placebo effects are often seen in IBS treatment. Placebo effects are improvements in symptoms that are due to patient beliefs in a particular treatment rather than the specific biological effects of the treatment.

This review included 17 randomized controlled trials (RCTs) including a total of 1806 participants. Five RCTs (411 participants) compared acupuncture to sham acupuncture for the treatment of IBS. Sham acupuncture is a procedure in which the patient believes he or she is receiving true acupuncture. However, in sham acupuncture the needles either do not penetrate the skin or are not placed at the correct places on the body, or both. Sham acupuncture is intended to be a placebo for true acupuncture. The sham-controlled studies were well designed and of high methodological quality. These studies tested the effects of acupuncture on IBS symptom severity or health-related quality of life. None of these RCTs found acupuncture to be better than sham acupuncture. Evidence from four Chinese language comparative effectiveness trials showed acupuncture to be superior to two antispasmodic drugs (pinaverium bromide and trimebutine maleate), both of which provide a modest benefit for the treatment of IBS, although neither is approved for treatment of IBS in the United States. It is unclear whether or not the greater benefits of acupuncture reported by patients in these unblinded studies are due entirely to patients' greater expectations of improvement from acupuncture than drugs or preference for acupuncture over drug therapy. There was one side effect (i.e. fainting in one patient) associated with acupuncture in the nine trials that reported side effects, although relatively small sample sizes limit the usefulness of this safety data.

SUMMARY OF FINDINGS

Summary of findings for the main comparison. Acupuncture versus sham acupuncture for irritable bowel syndrome

Acupuncture versus sham acupuncture for irritable bowel syndrome

Patient or population: patients with irritable bowel syndrome **Settings:** Canada (1), Germany (1), UK (1), US (2) **Intervention:** True acupuncture **Comparison:** Sham acupuncture

Outcomes	mes Illustrative comparative risks* (95% CI)			No of Partici- pants	Quality of the evidence	Comments	
	Assumed risk	Corresponding risk	- (95% CI)	(studies)	(GRADE)		
	Sham acupuncture	True acupuncture					
symptom severity (con- tinuous outcome) IBS Severity Scoring Sys- tem (IBS-SSS) ¹ . Scale from: 0 to 500. Follow-up: 3-13 weeks ²	The mean symptom severity (continuous outcome) in the con- trol groups was 193 points ³	The mean symptom severity (contin- uous outcome) in the intervention groups was 9.2 lower (29.2 lower to 10.8 higher) ⁴ (Bet- ter values are indicated by lower scores.)		281 (4 studies)	⊕⊕⊕⊝ moderate ⁵	SMD -0.11 (-0.35 to 0.13)	
quality of life (continu- ous outcome) IBS Quality of Life (IBS- QOL) Scale ⁶ . Scale from: 0 to 100. Follow-up: 3-5 weeks ²	The mean quality of life (continuous out- come) in the control groups was 73.8 points ³	The mean quality of life (continuous outcome) in the intervention groups was .53 lower (4.8 lower to 3.9 higher) ⁷ (Better val- ues are indicated by higher scores.)		253 (3 studies)	⊕⊕⊕⊝ moderate ⁵	SMD -0.03 (-0.27 to 0.22)	

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** Confidence interval;

GRADE Working Group grades of evidence

High quality: Further research is very unlikely to change our confidence in the estimate of effect.

Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate.

Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. Very low quality: We are very uncertain about the estimate.

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¹ The IBS Severity Scoring System (IBS-SSS) possesses responsiveness, face and construct validity (Francis 1997), and is one of the most widely used IBS symptom severity outcome measures. Better values are indicated by lower scores.

² This outcome is for the short-term follow-up time point, defined as the time point closest to 8 weeks and less than or equal to 3 months after randomization. Outcomes were measured at the end of the treatment period in all studies, and because in all studies the end of treatment time point coincided with the time point closest to 8 weeks and less than or equal to 3 months, our short term outcome time points were the end of treatment for all trials.

³ We used the Lembo 2009 trial as the representative trial for the final value scores of symptom severity and quality of life in the control group because this trial was sufficiently large; the patient characteristics and the baseline means and SDs of symptom severity and quality of life in the control group of this trial were similar to and representative of the other trials; and this trial used the familiar IBS-SSS scale for symptom severity and the well-validated IBS-QoL for quality of life.

⁴ The standardized mean difference (SMD) was re-expressed into a mean difference by applying the calculated SMD back into the Lembo 2009 study and depicted on the IBS-SSS scale used in that study. This calculation was made by multiplying the post-treatment standard deviation of the IBS-SSS score of the sham group in the Lembo trial by the pooled SMD.

⁵ Imprecision due to sparse data (less than 400 events) and confidence intervals include possibility of benefit.

⁶ The IBS Quality of Life measure (IBS-QoL) (Patrick 1998) is an extensively validated IBS quality of life scale (Bijkerk 2003; Irvine 2006). Better values are indicated by higher scores. ⁷ The standardized mean difference (SMD) was re-expressed into a mean difference by applying the calculated SMD back into the Lembo 2009 study and depicted on the IBS-QoL scale used in that study. This calculation was made by multiplying the post-treatment standard deviation of the IBS-QoL score of the sham group in the Lembo trial by the pooled SMD.

Summary of findings 2. Acupuncture versus pharmaceutical medications for irritable bowel syndrome

Acupuncture versus pharmaceutical medications for irritable bowel syndrome

Patient or population: patients with irritable bowel syndrome

Settings: China (5)

Intervention: Acupuncture

Comparison: Pharmaceutical medications

Outcomes	Illustrative comparative risks* (Relative effect (95% CI)	No of Partici- pants	Quality of the evidence	Comments	
	Assumed risk	Corresponding risk	- (55% CI)	(studies)	(GRADE)	
	Pharmaceutical medications	Acupuncture				
symptom severity (di- chotomous outcome) Dichotomous measure of overall symptom severi- ty ¹ Follow-up: 3-7 weeks ²	633 per 1000 ³ had improved overall symptom severity in the pharmaceutical medication group	810 per 1000 (709 to 918) had improved overall symptom severity in the acupuncture group	RR 1.28 (1.12 to 1.45)	449 (5 studies)	⊕⊕⊙⊝ low ^{4,5}	

*The basis for the **assumed risk** (e.g. the median control group risk across studies) is provided in footnotes. The **corresponding risk** (and its 95% confidence interval) is based on the assumed risk in the comparison group and the **relative effect** of the intervention (and its 95% CI). **CI:** Confidence interval; **RR:** Risk ratio;

GRADE Working Group grades of evidence

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High quality: Further research is very unlikely to change our confidence in the estimate of effect. Moderate quality: Further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate. Low quality: Further research is very likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate. Very low quality: We are very uncertain about the estimate.

¹ In cases where dichotomous outcomes such as improvement in IBS symptoms were presented in the form of multiple strata, such that we had the option of choosing cutpoints for the dichotomous outcome, we created a dichotomous measure in which all positive outcomes were combined into a single positive category (i.e., improvement) and the remaining strata constituted the negative category (i.e., no improvement). When investigators selected a cutpoint on a continuous scale to dichotomize between improvement and no improvement, we used the same cutpoint to define the dichotomous outcome.

² This outcome is for the short-term follow-up time point, defined as the time point closest to 8 weeks and less than or equal to 3 months after randomization.

³ The assumed risk in the control group is based on the percentage of all participants in the control group who experienced improvement in symptom severity. There were a total of 229 control group participants, and 145 of these participants experienced improvement in symptom severity.

⁴ The primary limitation is that treatment in these studies was unblinded, and it is unclear whether or not the greater benefits of acupuncture reported by patients in these unblinded trials are due entirely to patients' greater expectations of improvement from acupuncture than drugs. Additionally, one study author (Xue Y 2009) explained that allocation to treatment was by means of a random number table and did not give further details.

⁵ Imprecision due to sparse data (less than 400 events).

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BACKGROUND

Irritable bowel syndrome (IBS) is a chronic, relapsing gastrointestinal disorder characterized by altered bowel habits and abdominal pain and discomfort (Brandt 2009). A systematic review (Saito 2002) has estimated that 10 to 15% of adults in North America have IBS, as diagnosed by either the Rome (Vanner 1999) or Manning (Saito 2000) objective diagnostic criteria. IBS is associated with significant reductions in both health-related quality of life (El-Serag 2002) and work productivity (Maxion-Bergemann 2006; Brandt 2009) and increased consumption of medical resources. Indeed, people with IBS consume over 50% more health care resources than age-matched controls without IBS (Longstreth 2003; Talley 1995). The combined direct and indirect costs associated with IBS patients in the United States in 2004 were estimated at over \$1 billion (Everhart 2008).

Effective treatments for IBS are needed to relieve symptoms, improve quality of life, and to reduce healthcare utilization. In 2009, the American College of Gastroenterology Task Force conducted a series of systematic reviews to evaluate the efficacy of both pharmacological and non-pharmacological therapies for treating IBS (Brandt 2009). In terms of pharmacological treatments, the Task Force found "poor quality of evidence" for certain antispasmodics and "moderate guality of evidence" for tricyclic antidepressants, selective serotonin reuptake inhibitors, non-absorbable antibiotics (for diarrhea-predominant IBS), and C-2 chloride channel activators (for constipation-predominant IBS). The Task Force found "good quality of evidence" for 5HT₃ antagonists and 5HT₄ agonists, but noted that these agents carry a possible risk of ischemic colitis and cardiovascular events, respectively, which may limit their utility. A subsequent systematic review showed that the benefits of these 5HT₃ antagonists and 5HT₄ agonists relative to placebo are "modest" (Ford 2009a). In terms of non-pharmacological therapies, the Task Force found "poor quality of evidence" for psyllium fiber and peppermint oil. The Task Force also noted that preliminary evidence suggested that some probiotics may be effective in reducing IBS symptoms (Brandt 2009). A subsequent systematic review (Brenner 2009) concluded that the specific probiotic B. infantis 35624 has shown repeated efficacy in well-designed randomized controlled trials (RCTs), and can be considered an effective treatment for IBS.

The Task Force was unable to make any recommendations either for or against acupuncture for treating IBS because the only systematic review available at the time was a Cochrane review (Lim 2006) which was inconclusive because it included only small, heterogeneous, and methodologically unsound trials. Given the safety of acupuncture (MacPherson 2001; White 2001; Melchart 2004) and the limited availability of other safe and effective treatments for IBS, the question of whether acupuncture is effective for treating IBS is highly relevant. Recently, RCTs have been published which provide greater evidence to estimate the effects of acupuncture for treating IBS. We have therefore updated our previous Cochrane systematic review and meta-analysis of acupuncture for IBS (Lim 2006) to assess whether the pooled effects of currently available trials show any benefit of acupuncture for improving symptoms or health-related quality of life in patients with IBS.

OBJECTIVES

The primary objectives were to assess the efficacy and safety of acupuncture for treating IBS.

METHODS

Criteria for considering studies for this review

Types of studies

We included randomized controlled trials (RCTs) in any language, published as either full articles or abstracts. Recent research indicates that a large proportion of Chinese-language RCT reports are of studies that are not truly randomized (Wu 2009). An author (KC) therefore contacted the investigators of Chineselanguage RCTs by telephone to determine whether they had used randomization. The interviews were conducted using questions adapted from the survey developed by Wu et al to verify the authenticity of "claimed" randomized trials (Wu 2009). Responses were recorded and forwarded to a second review author (EM) for confirmation of RCT authenticity. The same questions were asked of authors of English-language RCTs that did not include details about randomization methods in their published reports. Trials that were found to assign patients by alternation, rotation, or hospital record number were excluded. Trials that used a random method of assignment, but with flaws or suspected flaws in the random assignment process were included, but with their limitations described.

Types of participants

We included trials involving adult participants diagnosed with irritable bowel syndrome (IBS). Examples of diagnostic criteria included but were not limited to the Manning criteria (Manning 1978), and Rome I (Drossman 1994), Rome II (Thompson 2000), or Rome III criteria (Longstreth 2006). We did not exclude trials in which patients were stated to be diagnosed with IBS but no diagnostic criteria were described.

Types of interventions

We included trials evaluating Traditional Chinese Medicine (TCM) acupuncture. TCM acupuncture involves inserting needles into traditional meridian points, usually with the intention of influencing energy flow in the meridian. Needles may also be inserted at additional tender points and electrical stimulation of the needles may be used. Since TCM acupuncture is often accompanied by moxibustion, we included trials using moxibustion as a co-intervention with acupuncture. We excluded trials of dry needling or trigger point therapy, a therapy which is based on principles of Western anatomy and physiology and rejects TCM concepts of energy and meridians. We also excluded RCTs of laser acupuncture, non-invasive electrostimulation (i.e. using electrodes on the skin rather than needles to stimulate acupuncture points (Ezzo 2005)), and acupressure, to restrict our focus to the effects of traditional needle acupuncture. Finally, we excluded trials of micropuncture, a non-traditional acupuncture practice which is based on the principle that the ear (or nose, eye, etc.) is a microsystem of the entire body, and in which needles are only inserted on that microsystem. Identified trials using types of interventions that are not eligible are described in the 'Characteristics of excluded studies' table.

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We included trials comparing acupuncture to any of the following control interventions: sham (placebo) acupuncture, no (specific) treatment, an active non-TCM treatment, or evaluated acupuncture as an adjuvant to another treatment. Because our objective was to evaluate the effects of TCM acupuncture compared to a sham treatment, no treatment or a Western medicine control, we excluded RCTs in which one form of acupuncture was compared with another form of acupuncture or a different type of TCM (e.g. Chinese herbal medicine). Adjunctive treatments, either Western or TCM, were allowed as long as they had been given to both the acupuncture and control groups.

Types of outcome measures

Our primary outcomes were overall IBS symptom severity and IBS health-related quality of life. Studies that did not report at least one of these outcomes were excluded. Although there is no consensus on how to define and measure clinically meaningful improvement in IBS (Irvine 2006), two recent evaluations of symptom and quality of life measures in IBS concluded that the IBS Adequate Relief question (IBS-AR) (Mangel 1998) and the IBS Symptom Severity Scale (IBS-SSS) (Francis 1997) possessed responsiveness, face and construct validity and were two of the most appropriate IBS symptom outcome measures, while the IBS Quality of Life measure (IBS-QoL) (Patrick 1998) was the most extensively validated quality of life scale (Bijkerk 2003; Irvine 2006). When an individual study reported more than one overall IBS symptom severity measure, and either the IBS-AR question or the IBS-SSS scale was present, we gave preference to the IBS-AR question for dichotomous outcomes and to the IBS-SSS scale for continuous outcomes. When an individual study reported more than one quality of life measure, we gave preference to the IBS-QoL if present. In cases where dichotomous outcomes such as improvement in IBS symptoms were presented in the form of multiple strata, such that we had the option of choosing cutpoints for the dichotomous outcome, we followed the model of Ford et al. and created a dichotomous measure in which positive outcomes were combined into a positive category (i.e. improvement) and the remaining strata constituted the negative category (i.e. no improvement) (Ford 2009a; Ford 2009b). In cases where investigators selected a cutpoint on a continuous scale to dichotomise between improvement and no improvement, we used the same cutpoint to define the dichotomous outcome (Ford 2009a).

Timing of outcome assessment

We extracted outcome data for both short and long-term followup points. Short-term follow-up was defined as three months or less after randomization, and long-term follow-up was defined as closest to six months but more than three months after randomization. When we observed multiple short-term follow-up points, we chose to extract the data closest to eight weeks after randomization, which coincided with end of treatment.

In randomized crossover trials, only outcomes from the first period were eligible for inclusion, due to the risk of carry-over effects.

Adverse outcomes

We extracted data on adverse events when such data were present in the trial report.

Search methods for identification of studies

Electronic searches

To identify RCTs, we searched the Cochrane Central Register of Controlled Trials (*The Cochrane Library*, Issue 11, 2011), the Cochrane Inflammatory Bowel Disease and Functional Bowel Disorders Review Group Specialized Register, the Cochrane Complementary Medicine Field Specialized Register, MEDLINE/ PubMed, EMBASE, the Cumulative Index to Nursing and Allied Health, and the Chinese databases Sino-Med, CNKI and VIP, during November 15 to 28, 2011. We developed search strategies for each database, based upon the original Medline search strategy constructed for the previous version of this review (Lim 2006). The strategies and results for each database are presented in Appendix 1.

Searching other resources

We considered all RCTs included in the previous version of this review for inclusion in this update. We scanned bibliographies of included articles and systematic reviews for further references. Finally, we searched ClinicalTrials.gov to identify trials that may be relevant for future updates of this review.

Data collection and analysis

Selection of studies

All records identified by searching were independently screened by at least one author. The full text of potentially relevant reports was obtained and independently reviewed by two authors for eligibility. Disagreements between reviewers were resolved by discussion.

Data extraction and management

All data were independently extracted by two authors. In addition to the outcomes of overall IBS symptom severity and IBS healthrelated quality of life for all time points reported, we extracted data pertaining to the methods of the trial, characteristics of the participants, details of the acupuncture and control interventions, and treatment outcomes. Any disagreements were resolved by discussion. When reported data were incomplete or ambiguous, we requested additional information or clarification from the corresponding authors.

The method of selecting acupuncture points was categorized as fixed, flexible or individualized. For the fixed method, the same points are used for all participants. For the flexible method, a fixed formula is used and some additional points are chosen for the individual participant, based upon individual diagnosis or symptoms. In individualized point selection, the practitioner is free to choose any points.

Assessment of risk of bias in included studies

We assessed risk of bias using the Cochrane Collaboration's risk of bias tool (Higgins 2011a). We used the following six criteria:

- Adequate sequence generation: Was the allocation sequence adequately generated?
- Allocation concealment: Was the allocation adequately concealed?
- Blinding: Was knowledge of the allocated intervention adequately prevented during the study?

Library

- Incomplete outcome data: Were incomplete outcome data adequately addressed?
- Selective reporting: Were reports of the study free of suggestion of selective outcome reporting?
- Other risk of bias: Was the study apparently free of other problems that could put it at a high risk of bias?

For each risk of bias question, an answer of 'Yes' indicates low risk of bias, 'No' indicates high risk of bias and 'Unclear' indicates unclear or unknown risk of bias. We used the criteria described in the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011a) for judgments of 'Yes', 'No' and 'Unclear'. For "Blinding", we assigned sham-controlled trials a judgment of "Unclear" unless we felt certain that the sham control was sufficiently credible in fully blinding participants to the treatment being evaluated (Manheimer 2007; Manheimer 2010). We assigned the "Yes" score to shamcontrolled trials that either 1) evaluated the credibility of the sham and found the sham to be indistinguishable from true acupuncture or 2) used a penetrating needle or a previously validated sham needle (i.e. Streitberger needle). There is no universally agreedupon instrument to measure outcomes in trials of IBS treatment, however overall IBS symptom severity is a key outcome in the majority of IBS trials and health-related quality of life is an important secondary outcome that may legitimately be considered as a primary outcome measure (Irvine 2006). For the selective reporting item, we assigned the "Yes" score to trials that both 1) reported outcomes for overall IBS symptom severity or healthrelated quality of life, or both, and 2) reported, at the end of treatment (and follow-up, if done), the results of each outcome measured according to the Methods section. For "Other risk of bias", we evaluated 2 other risk of bias-related criteria: baseline comparability and use of an intention to treat analysis. For the baseline comparability criterion to be considered adequate, a comparison of the symptom scores between the treatment and control group(s) at baseline needed to be reported

Although we did not explicitly consider this to be a risk of bias measure, we also extracted data on the sources of funding for each trial.

As a first step in evaluating risk of bias, we copied information relevant for making a judgment on a criterion from the original publication into a table. If available, we also entered any additional information from the RCT authors into this table. Two review authors (EM and LSW or KC for English-language trials, and LL and KC for Chinese-language trials) independently judged whether the risk of bias for each criterion should be considered low, high or unclear. Any disagreement was resolved by discussion.

We used the GRADE approach to rate the overall quality of evidence for the primary outcomes. RCTs start as high quality evidence, but may be downgraded due to: (1) limitations in design and implementation (risk of bias), (2) indirectness of evidence, (3) inconsistency (unexplained heterogeneity), (4) imprecision (sparse data), and (5) reporting bias (publication bias). The overall quality of evidence for each outcome was determined after considering each of these elements, and categorized as high quality (i.e. further research is very unlikely to change our confidence in the estimate of effect); moderate quality (i.e. further research is likely to have an important impact on our confidence in the estimate of effect and may change the estimate); low quality (i.e. further research is very likely to have an important impact on our confidence in the

estimate of effect and is likely to change the estimate); or very low quality (i.e. we are very uncertain about the estimate) (Guyatt 2008; Schünemann 2011).

Assessment of acupuncture adequacy

Two acupuncturists (LL, XS) who have a combined clinical experience of nearly fifty years in treating IBS with acupuncture, and who have previously worked on RCTs of acupuncture, independently assessed the adequacy of the acupuncture administered in the trials. Six aspects of the acupuncture intervention were assessed for adequacy: choice of acupuncture points; total number of sessions; treatment duration; treatment frequency; needling technique; and acupuncturist's experience (Furlan 2005). The likelihood of the sham intervention to have physiological activity was also assessed, using an open-ended question. The acupuncturist assessors were provided with only the part of the publications that described the acupuncture and sham procedures, so that their assessments could not be influenced by the results of the trials. To test the success of blinding the assessors to the study publication and results, we asked the assessors to guess the identity of each study being assessed. The acupuncturists assessed adequacy independently and achieved consensus by discussion.

Measures of treatment effect

Dichotomous data

Dichotomous outcomes (e.g. adequate symptom relief) were expressed as relative risks (RR) with 95% confidence intervals (CI).

Continuous data

For continuous outcomes on the same scale (e.g. symptom severity as measured by the IBS-SSS), the mean difference (MD) and 95% confidence intervals were calculated. For continuous outcomes on different scales that assess the same underlying construct (e.g. different measures of symptom severity), the standardized mean difference (SMD) and 95% confidence intervals were calculated.

Dealing with missing data

In cases where participants were lost to follow-up, and intentionto-treat (ITT) analyses were conducted using baseline observations carried forward, multiple imputation, or other methods to impute the missing values, we used the ITT data for our primary analysis, if group means and standard deviations were present or could be estimated, and if the method for imputing data was described and did not bias the effect size calculation. If only the ITT data were reported, and not the available case data, we used the ITT data. When statistics such as standard deviations were not present in the study report, we used the methods suggested in Chapter 16 of the Cochrane Handbook for Systematic Reviews of Interventions (Higgins 2011b) to calculate or estimate the values of the missing statistics.

Assessment of heterogeneity

We evaluated heterogeneity using the I² statistic (Higgins 2003), which indicates the proportion of variability across trials not explained by chance alone. Roughly, I² values of 50% to 90% may represent substantial heterogeneity, and I² values of 75% to 100% represent considerable heterogeneity (Higgins 2003; Deeks 2011). We also checked for heterogeneity by visual examination of forest plots. When heterogeneity was observed, we attempted to determine potential reasons for it by examining individual study characteristics, such as study population, type and duration of treatment, and type of control intervention.

Data synthesis

Data from individual trials were combined when the trials were sufficiently similar in terms of control interventions (sham (placebo) acupuncture, no treatment, another active treatment, or acupuncture as adjuvant to another treatment), outcome measures (overall IBS symptom severity, IBS-related quality of life), and timing of outcome assessment (short-term, long-term). For pooled data, summary test statistics were calculated using a random-effects effect model to account for expected heterogeneity. If the I² statistic was greater than or equal to 50%, the summary measures of effect were interpreted with caution, and heterogeneity between trials was investigated.

For the acupuncture versus sham comparison, data for the symptom severity outcome were presented in some studies as dichotomous data (e.g. adequate symptom relief) and in other studies as continuous data (e.g. symptom severity as measured by the IBS-SSS). We re-expressed odds ratios as standardized mean

differences (SMDs), thereby allowing dichotomous and continuous data to be pooled together for this comparison/outcome (Deeks 2011), using the generic inverse variance method in RevMan.

Sensitivity analysis

We conducted sensitivity analyses using fixed-effect and randomeffects meta-analytic estimates, and for available case and imputed data, when such data were available.

To assess whether treatment effects varied with internal validity of studies or treatment adequacy, we attempted sensitivity analyses to evaluate 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 the risk of bias and treatment-adequacy criteria.

RESULTS

Description of studies

Results of the search

Figure 1 shows details of the search and selection process.



Figure 1. Flow of studies through selection process.

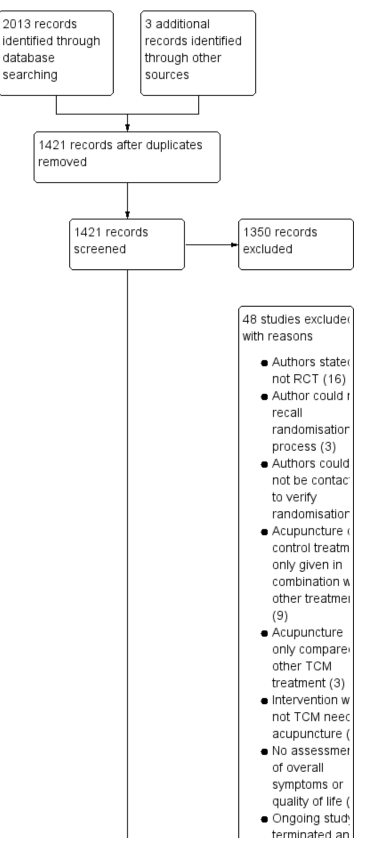
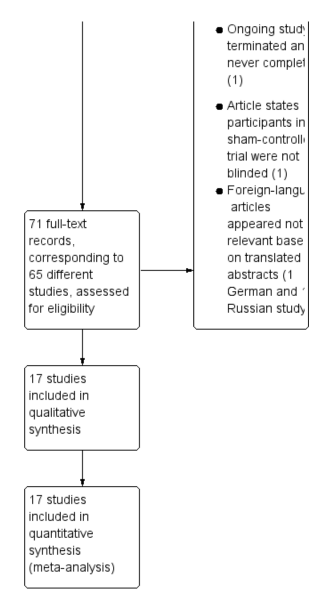




Figure 1. (Continued)



Included studies

Seventeen studies with a total of 1806 participants were included (Liu 1997; Lowe 2000; Forbes 2005; Schneider 2006; Reynolds 2008; Xiong 2008a; Anastasi 2009; Chen 2009; Lembo 2009; Li 2009; Xue 2009; An 2010; Liu 2010; Shi 2010; Zeng 2010; Chen 2011; Sun 2011; See 'Characteristics of included studies' table). Five of the studies were small, including between 29 and 59 participants (Lowe 2000; Forbes 2005; Schneider 2006; Reynolds 2008; Anastasi 2009). The largest study included 230 participants (Lembo 2009). The studies were conducted in the US (Anastasi 2009; Lembo 2009), the UK (Forbes 2005; Reynolds 2008), China (Liu 1997; Xiong 2008a; Chen 2009; Li 2009; Xue 2009; An 2010; Liu 2010; Shi 2010; Zeng 2010; Chen 2011; Sun 2011), Canada (Lowe 2000) and Germany (Schneider 2006). The studies conducted in China were published in Chinese, and the remaining studies were published in English. All were published as full articles, with the exception of the Lowe 2000 trial, which was published as an abstract. Three of the studies included in our 2006 Cochrane review (Liu 1995; Liao 2000; Fireman 2001) were excluded from this update because either an adequate

randomization process was not used (Liao 2000; Fireman 2001) or the randomization procedure could not be recalled by the author (Liu 1995). Appendix 2 includes an overview description of trial characteristics and acupuncture and control interventions.

Excluded studies

The 'Characteristics of excluded studies' table lists the studies that we excluded as well as the reasons for exclusion.

Risk of bias in included studies

All sham-controlled trials reported adequate methods for sequence generation and allocation concealment, and all trials except the Lowe 2000 trial adequately described and addressed losses to follow-up. The Lowe 2000 trial was reported only as an abstract, and the completeness of outcome data ascertainment could not be assessed.

In 4 out of 5 of the sham-controlled trials (Forbes 2005; Schneider 2006; Lembo 2009; Anastasi 2009), we judged that the shams



were likely to be indistinguishable from true acupuncture. In two of these sham-controlled trials (Schneider 2006; Lembo 2009), the Streitberger placebo needle was used, which has been previously validated as a sufficiently credible sham (Streitberger 1998). The Streitberger needles were placed close to the genuine acupuncture points in both trials. In both trials, the participants were acupuncture naïve. In the Anastasi 2009 trial, the investigators superficially inserted needles at non points which were 2 to 3 cm away from the true points, and placebo moxibustion was performed above the same sham points without generating a heat sensation. In the Forbes 2005 trial, acupuncture needles were inserted at areas on the body that do not correspond to acupuncture points and are deemed to have no therapeutic value. The points and needling technique were varied somewhat each week, as was also done in the true acupuncture group, who received individualized point selection. In each of these four trials, the sham was likely to be indistinguishable from true acupuncture. For the Lowe 2000 trial, the sham was judged to have been potentially detectable as a fake treatment by the trial participants. The sham procedure involved tapping a blunt needle on the skin and then taping the needle in place. Although this procedure was described as "validated" in the Lowe 2000 abstract, we are unaware of a validation study for this procedure. Also, in the trial report there was no description of whether or not the patients were required to have never previously used acupuncture, and there were no reported tests for checking the success of the blinding.

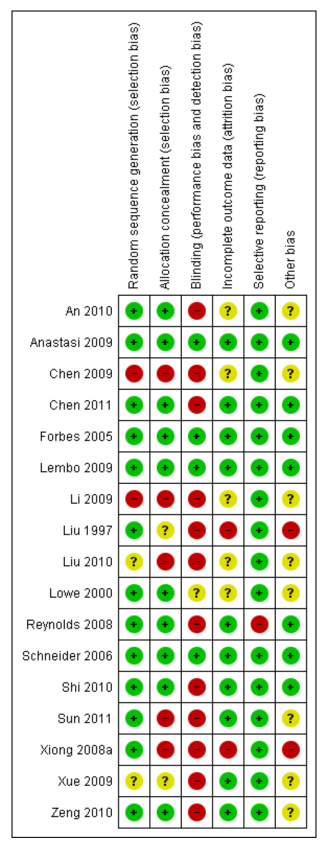
In the trials comparing acupuncture with another active treatment (Liu 1997; Li 2005; Xue 2009; An 2010; Liu 2010; Shi 2010; Zeng 2010; Chen 2011; Sun 2011), no (specific) treatment (Reynolds 2008; Lembo 2009), or evaluating acupuncture as an adjuvant to

another treatment received by all trial participants (Liu 1997; Xiong 2008a; Chen 2009; Li 2009; Liu 2010), blinding of participants was not possible, and this likely represents the major risk of bias in these trials. In these pragmatic trials, there were also risks of bias associated with the randomization procedure and the follow-up of patients. Although all of these trials reported to use a random sequence generation for the treatment assignment, 5 of these trials (Liu 1997; Chen 2009; Li 2009; Xue 2009; Liu 2010) reported an equal number of participants in each group, which would be unlikely to occur by chance with the simple randomization methods used in these trials. The authors of the Chen 2009 and Li 2009 trials confirmed in telephone interviews that a few patients were nonrandomly assigned to achieve identical sized treatment groups. The authors of the other 3 trials (i.e. Liu 1997; Xue 2009; Liu 2010), were unable to explain how equal sample sizes were achieved. In 6 of the Chinese language trials (Liu 1997; Xiong 2008a; Chen 2009; Li 2009; An 2010; Liu 2010), incomplete outcome data were not adequately addressed. In four trials (Liu 1997; Xiong 2008a; Li 2009; An 2010), the authors did not report the numbers of drop-outs in the publication, and did not have records of the numbers of dropouts to provide during the telephone interviews. For the Liu 2010 trial, the authors reported no drop-outs, which would be unusual in a 4 week trial of 300 participants. For the Chen 2009 trial, the authors endeavoured to maintain equal group sizes, by eliminating participants who withdrew during the trial and replacing them with new patients, and the number of such replacements was not recorded by the author.

A 'risk of bias graph' displays the judgments for each risk of bias item for each included study (Figure 2).



Figure 2. Risk of bias summary: review authors' judgements about each risk of bias item for each included study.



Acupuncture adequacy



All trials included in this review were judged adequate on "Choice of acupoints" and "Needling technique", except for the Lowe 2000 trial, for which the acupuncture points and needling technique were not reported. The acupuncture frequency was judged adequate in all trials except for the Forbes 2005 and Reynolds 2008 trials. All trials that reported on the acupuncturist's experience were judged adequate, except for the Lowe 2000 trial, for which a physiotherapist with Level 1 accreditation in acupuncture was used, which was judged inadequate. Also, for two trials (Forbes 2005; Anastasi 2009), the acupuncture adequacy assessors noted that the sham needling may have had physiologic activity. The results of the acupuncture adequacy assessments are reported in Appendix 3.

Effects of interventions

See: Summary of findings for the main comparison Acupuncture versus sham acupuncture for irritable bowel syndrome; Summary of findings 2 Acupuncture versus pharmaceutical medications for irritable bowel syndrome

The GRADE analyses for the main comparisons are presented in Summary of findings for the main comparison and Summary of findings 2.

Acupuncture versus sham

Five trials (Lowe 2000; Forbes 2005; Schneider 2006; Anastasi 2009; Lembo 2009) compared the effects of acupuncture and sham acupuncture. The Schneider 2006 trial did not measure

the outcome of symptom severity and the Anastasi 2009 and Lowe 2000 trials did not report quality of life. The 5 individual sham-controlled RCTs, and also the pooled analyses, found no statistically significant differences between acupuncture and sham acupuncture for the outcomes symptom severity or quality of life.

Data for the symptom severity outcome were presented as dichotomous data (e.g. adequate symptom relief) in some studies and as continuous data (e.g. symptom severity as measured by the IBS-SSS) in other studies. We re-expressed odds ratios as standardized mean differences (SMDs), thereby allowing dichotomous and continuous data to be pooled together for this comparison/outcome (Deeks 2011), using the generic inverse variance method in RevMan. This pooled analysis for symptom severity included 4 studies and 281 patients. There was no statistically significant difference between acupuncture and sham acupuncture (SMD - 0.11, 95% CI -0.35 to 0.13; See Figure 3). The GRADE analysis indicated that the overall quality of the evidence for this outcome was moderate due to sparse data (less than 400 events; See Summary of findings for the main comparison). Three studies (231 patients) reported symptom severity as a continuous outcome. There was no statistically significant difference between acupuncture and sham acupuncture (SMD - 0.09, 95% CI -0.35 to 0.17; See Analysis 1.1). Three studies (262 patients) reported on adequate symptom relief. Fifty-five per cent of patients in the acupuncture group had adequate symptom relief compared to 49% of patients in the sham acupuncture group. There was no statistically significant difference between acupuncture and sham acupuncture (RR 1.09, 95% CI is 0.87 to 1.37; See Analysis 1.2).

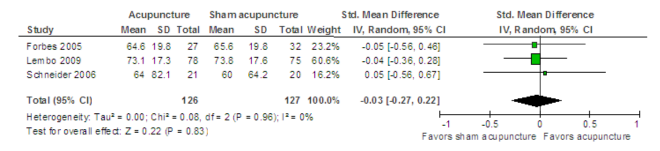
Figure 3. Acupuncture versus sham acupuncture: Symptom severity

Study	Acupuncture Total	Sham ac upunct Total	ture Weight	Std. Mean Difference IV, Random, 95% (
An astasi 2009	12	15	9.9%	0.25 [-0.51, 1.01]	
Forbes 2005	22	29	18.6%	0.11 [-0.45, 0.66]	-
Lem bo 2009	78	75	56.7%	-0.21 [-0.53, 0.11]	-
Lowe 2000	28	22	14.9%	-0.26 [-0.88, 0.36	j <u> </u>
Total (95% CI)	140	141	100.0%	-0.11 [-0.35, 0.13]	
Heterogeneity: Tau ²	= 0.00; Chi ² = 2.05, df = 3	3 (P = 0.56); I ² = 0	%		
Test for overall effec	t: Z = 0.91 (P = 0.36)				-1 -0.5 0 0.5 1 Favors acupuncture Favors sham acupunctu

For the acupuncture versus sham comparison, all 3 studies (253 patients) that included the quality of life outcome reported continuous data. There was no statistically significant difference in quality of life scores between acupuncture and sham acupuncture groups (SMD -0.03, 95% CI -0.27 to 0.22; See Figure 4). A GRADE analysis indicated that the overall quality of the evidence for this outcome was moderate due to sparse data (less than 400 events; See Summary of findings for the main comparison). One study reported quality of life as a dichotomous outcome (153 patients).

Seventeen per cent of acupuncture patients had an improvement in quality of life compared to 14% of sham acupuncture patients. There was no statistically significant difference (RR 1.25, 95% CI 0.58 to 2.68; See Analysis 1.5). For both outcomes, the results of all shamcontrolled trials were homogeneous ($I^2=0\%$; See Figure 3; Figure 4). One trial (Schneider 2006) assessed quality of life at long-term follow-up, and this trial did not find a difference in effect between acupuncture and sham acupuncture at six months (SMD 0.07, 95% CI -0.54 to 0.69; 41 patients; See Analysis 1.6).

Figure 4. Acupuncture versus sham acupuncture: Quality of life



Acupuncture versus other active treatments, as an adjuvant to other active treatments, and versus no specific treatment

All trials reported dichotomous outcome data for all non-sham comparisons, We pooled these trials as relative risks (see Figure 5). In cases where continuous outcome data were also reported

in the trials, we pooled the available continuous data as well, and present these as additional forest plots (See Analysis 2.2; Analysis 7.2; Analysis 7.3).There were no important differences between continuous and dichotomous results for any comparison or outcome. Librarv

Figure 5. Acupuncture versus another active treatment, as adjuvant to another active treatment, or compared to no specific treatment: Symptom severity

Study			Control Events			Risk Ratio M-H, Random, 95% Cl	Risk Ratio M-H, Random, 95% Cl
Acupuncture ver	sus phar	macol	ogical me	dicatio	ons		
Chen 2011	25	29	22	30	17.7%	1.18 [0.91, 1.53]	
Shi 2010	26	32	20		11.4%	1.54 [1.09, 2.18]	
Sun 2011	27	30	24	30	22.9%	1.13 [0.91, 1.39]	
Xue 2009	81	100	55	100	24.9%	1.47 [1.20, 1.80]	— —
Zeng 2010 Subtotal (95% Cl	27	29 220	24		23.0% 100.0%	1.20 [0.97, 1.49] 1.28 [1.12, 1.45]	•
Total events	186		145				
Heterogen eity: Ta Test for overall eff					0.21); 2=	31%	
Acupuncture ver	sus Bifid	obacte	erium				
An 2010	41	47	26	34	61.8%	1.14 [0.92, 1.42]	
Liu 2010 Subtotal (95% Cl	33)	50 97	34		38.2% 100.0%	0.97 [0.74, 1.28] 1.07 [0.90, 1.27]	
Total events	74		60				
Heterogen eity: Ta Test for overall eff				1 (P =	0.35); 1² =	0%	
Acupuncture ver							
Liu 1997 Subtotal (95% Cl	42	50 50	40		100.0% 100.0%	1.05 [0.87, 1.26] 1.05 [0.87, 1.26]	
Total events Heterogeneity: No Test for overall eff			40 = 0.60)				
Acupuncture plu	us psych	otheraj	py versus	psych	otherapy	alone	
Liu 1997 Subtotal (95% Cl	48	50 50	40		100.0% 100.0%	1.20 [1.03, 1.39] 1.20 [1.03, 1.39]	
Totalevents	48		40				
Heterogeneity: No Test for overall eff			= 0.02)				
Acupuncture plu	s anothe	r tradit	tional Chi	nese n	nedicine (TCM) treatment versus the	other TCM treatment alone
Chen 2009	29	30	28	30	28.6%	1.04 [0.92, 1.16]	- a
Li 2009	60	62	44	62	23.4%	1.36 [1.16, 1.61]	│ — ∎ —
Liu 2010	135	150	37	50	22.6%	1.22 [1.02, 1.45]	
Xiong 2008 Subtotal (95% Cl)	40)	42 284	34		25.4% 100.0%	1.12 [0.97, 1.30] 1.17 [1.02, 1.33]	
Totalevents Heterogeneity: Ta Test for overalleff				3 (P =	0.02); 1² =	69%	
Acupuncture ver	sus no si	pecific	treatmen	ıt			
Lembo 2009	46	78	27		63.0%	1.68 [1.18, 2.40]	
Reynolds 2008 Subtotal (95% Cl	9	9 87	5	17	37.0% 100.0%	3.11 [1.53, 6.33] 2.11 [1.18, 3.79]	
Total events	55		32				
Heterogen eity: Ta Test for overall eff	u ² = 0.11;		2.31, df=	1 (P =	0.13); l² =	57%	
							0.5 0.7 1 1.5 Favors control condition Favors acupunct

Acupuncture versus other active treatments

The five trials (Xue 2009; Shi 2010; Zeng 2010; Chen 2011; Sun 2011) that compared acupuncture to pharmacological therapies for IBS found that participants receiving acupuncture reported a significantly greater improvement in symptom severity than participants receiving pharmacological therapies. Eighty-four per

cent of acupuncture patients reported improvement in symptom severity compared to 63% of patients in the pharmacological treatment group. (RR 1.28, 95% CI 1.12 to 1.45, 449 patients, See Figure 5; Analysis 2.1). The GRADE analysis indicated that the overall quality of the evidence for this outcome was low due to risk of bias (i.e. all studies were not blinded) and sparse data (less than 400

events; See Summary of findings 2). Three studies (190 patients) reported symptom severity as a continuous outcome. Acupuncture patients had significantly lower mean symptom severity scores than patients in the pharmacological treatment group (WMD -0.72, 95% CI -1.02 to -0.43; See Analysis 2.2).

Two studies (181 patients) compared acupuncture with probiotics (An 2010; Liu 2010). There was no statistically significant difference in improvement between patients treated with acupuncture and probiotics (*Bifidobacterium*). Seventy-six per cent of acupuncture patients improved compared to 71% of probiotic patients (RR 1.07, 95% CI 0.90 to 1.27; See Figure 5; Analysis 3.1). Participants receiving acupuncture were not more likely to have responded to treatment than those treated with psychotherapy (Liu 1997). Eight-four per cent of patients in the acupuncture group improved symptomatically compared to 80% of patients in the psychotherapy group (RR 1.05, 95% CI 0.87 to 1.26, 100 patients (See Figure 5; Analysis 4.1).

Acupuncture as an adjuvant to other active treatments

Five trials (Liu 1997; Xiong 2008a; Chen 2009; Li 2009; Liu 2010) compared the combination of adjuvant acupuncture plus another IBS treatment received by all trial participants to the other IBS treatment alone. Patients who received acupuncture and psychotherapy were significantly more likely to have improvements in symptom severity than patients who received psychotherapy alone. Ninety-six per cent of patients in the combined acupuncture and psychotherapy group improved compared to 82% of patients in the psychotherapy group (RR 1.16, 95% CI 1.04 to 1.29, 2 studies, 182 patients; See Analysis 5.1). Pooled results (4 studies, 466 patients) showed that participants receiving adjuvant acupuncture were significantly more likely to have reported improvement than those treated with another Chinese medicine treatment alone (although there was substantial heterogeneity of results and high risks of bias in these trials). Ninety-three per cent of patients in the adjuvant acupuncture group reported improvement in symptom severity compared to 79% of patients who received traditional Chinese medicine alone (RR 1.17, 95% CI 1.02 to 1.33; See Figure 5; Analysis 6.1).

Acupuncture versus no specific treatment

Two trials (Lembo 2009; Reynolds 2008) compared the effects of acupuncture to no specific treatment. In both trials, all participants were allowed to continue receiving standard medical care for IBS, including any prescribed medications, but control group participants were not assigned to any additional IBS treatment. Both of these trials showed a statistically significant benefit of acupuncture for improving IBS symptom severity, although there was substantial heterogeneity of results between the 2 trials (181 patients). Sixty-three per cent of patients in the acupuncture group reported improvement in IBS symptom severity compared to 34% of patients in the no treatment group (RR 2.11, 95% CI 1.18 to 3.79; See Figure 5; Analysis 7.1). At short-term follow-up in a single trial (Lembo 2009), post-treatment quality of life scores showed acupuncture to be associated with significant improvement for the continuous quality of life measure (Analysis 7.3), but not the dichotomous quality of life measure (Analysis 7.4).

Sensitivity analyses

For the sham-controlled trials, sensitivity analyses based on risk of bias or treatment adequacy-related variables would be uninformative because all sham-controlled trials had similar results and no combination of these trials resulted in a pooled statistically significant benefit, for either the symptom severity or quality of life outcome. For the five trials comparing acupuncture versus pharmacological therapies, restriction to the four trials that compared acupuncture versus evidence-based (Ruepert 2011) antispasmodic pharmacological therapies (Shi 2010; Zeng 2010; Chen 2011; Sun 2011) had similar results (RR 1.21, 95% CI 1.07 to 1.37, 249 participants, I²=0). For the other comparisons, there were too few trials to attempt sensitivity analyses (Deeks 2011).

For the Forbes 2005 trial, which reported both intent-to-treat (ITT) and available case data for the symptom severity outcome, a sensitivity analysis using ITT values instead of the available case values did not result in important differences in the SMDs for this trial. The statistical significance of the pooled results did not change depending on whether the random-effects or fixed-effect analyses were used.

Safety of acupuncture

Nine trials included descriptions of adverse events associated with acupuncture (Forbes 2005; Reynolds 2008; Anastasi 2009; Lembo 2009; An 2010; Liu 2010; Shi 2010; Chen 2011; Sun 2011). For 8 of these 9 trials, no serious adverse events were reported, while the Shi 2010 trial reported that 1 participant in the electro-acupuncture group withdrew because of syncope (see Appendix 4).

DISCUSSION

Summary of main results

Five sham-controlled RCTs have tested the effects of acupuncture for treating IBS, and four of these trials used adequate methods for randomization and blinding, and had few withdrawals or dropouts. None of these sham-controlled RCTs found a statistically significant benefit of acupuncture relative to sham acupuncture for the outcomes symptom severity or quality of life. Similarly, pooling the data from these sham-controlled trials did not result in statistically significant benefits of acupuncture for either outcome. Five unblinded Chinese-language comparative effectiveness trials found that patients receiving acupuncture reported greater improvements in IBS symptoms compared to patients receiving pharmacological therapies for IBS.

How should physicians, researchers, and policy-makers interpret these seemingly contradictory trial findings, i.e., that acupuncture had no greater effects than a sham placebo, but acupuncture did show greater effects compared with two pharmacological treatments (pinaverium bromide and trimebutine maleate) that have both previously been shown to be superior to a placebo (Jailwala 2000; Ruepert 2011)? First, both the comparative effectiveness trials and the sham-controlled trials have important limitations that complicate their interpretations. An important limitation of the trials comparing acupuncture to pharmacological therapy is that the patients in these trials are not "blinded" to whether they received acupuncture or drug therapy, and expectation effects (i.e. defined as "the impact of expectations on subjective outcomes" (Flum 2006)), may differ between acupuncture and drug treatment (Kaptchuk 2006; Manheimer 2007; O'Connell 2009). That is, if patients randomized to acupuncture



expect greater improvements than patients randomized to drugs, the greater expectations of benefits from acupuncture may contribute to a larger placebo effect (i.e. a larger improvement in symptoms due to an inert treatment, or an inert component of a treatment) in the acupuncture group than in the drug treatment group. Because of the possibility of differential expectations of a benefit from acupuncture versus drugs in these trials (Linde 2007; Manheimer 2007; O'Connell 2009), it cannot be determined whether any of the reported benefits of acupuncture are due to a larger biological effect of acupuncture needling relative to drugs, or rather due entirely to the impact of the trial participants' greater expectations of a benefit of acupuncture, on the subjective outcomes that they reported.

A limitation of the sham-controlled trial design is that the high placebo effects of sham acupuncture may preclude the detection of any small, true biological benefits of true acupuncture relative to a credible sham acupuncture control, when subjective patient self reports are the outcome measures used. Two "methodological" trials have evaluated the placebo effects of sham acupuncture, on both subjective and objective outcome measures (Kaptchuk 2006; Wechsler 2011). One such methodological trial (Kaptchuk 2006), designed to compare placebo effects of placebo pills and sham acupuncture, found that, relative to placebo pills, sham acupuncture was more credible as an authentic treatment and resulted in higher subjective patient self reports of improvement. This trial also found that the placebo effect was confined to selfreported, subjective outcomes (e.g. pain) and that there was no placebo effect (i.e. no improvement from baseline) for either the placebo acupuncture or placebo pill on the objective outcome that they measured (i.e. grip strength). Another recent methodological trial (Wechsler 2011) compared albuterol (i.e. a proven asthma drug) versus sham acupuncture for asthma patients, and found that while only the albuterol had a biological effect on the objective outcome of airway flow, both the sham acupuncture and albuterol groups had dramatic and comparable improvements from baseline on the subjective outcome of patient self-reports of improvement, such that the albuterol showed no benefit relative to the sham acupuncture on self-reported improvement.

These methodological studies suggest that relying exclusively on subjective patient reports, such as those used as outcomes in IBS trials, may result in a failure to detect small biological effects of an active treatment (i.e. true acupuncture) relative to a highly credible, but physiologically inert, sham acupuncture control. Thus, while the high placebo effects among IBS patients (Spiller 1999) make it difficult to show that any pharmacological treatment is superior to an inert placebo pill, demonstrating such an effect may be even more difficult when the placebo control is sham acupuncture.

Overall completeness and applicability of evidence

How externally valid are the results of this review? Namely, do the types of interventions investigated in these studies represent current best practice of acupuncture treatment for IBS? Assessing the adequacy of the acupuncture treatment procedure is important because, for instance, basing conclusions about acupuncture efficacy on a suboptimal procedure is "analogous to a pharmaceutical trial formulating conclusions about the efficacy of a drug based on an inadequate dose" (Ezzo 2001).

For the sham-controlled trials, a possible reason for the lack of benefit might be explained by the fact that only the Forbes 2005 trial

used individualized acupuncture, in which the acupuncturist tailors the point selection individually to each patient, and which is the typical approach used in everyday acupuncture clinical practice. Among the other four sham-controlled trials that did not use an individualized approach, two (Anastasi 2009; Lembo 2009) used a "flexible formula" approach, and two (Lowe 2000; Schneider 2006) used a fixed formula approach. With the fixed formula approach, the same acupoints are used for all trial participants. With the flexible formula approach, some acupoints are required for all patients but other acupoints could be selected based on the individual patient's specific constellation of symptoms. In terms of RCT design, an advantage of the flexible formula approach over the entirely individualized approach is that using a flexible formula allows for the results of the RCT to be relatively reproducible and externally valid (Lembo 2009) while at the same time allowing for some discretion of the acupuncturist to individualize point selection. In an RCT setting, this flexible formula approach is also easier to blind than an individualized approach, because individualization requires increased contact between the patient and the acupuncturist, which increases the risk of unblinding. Thus, although four trials did not use individualized acupuncture, the two acupuncturist systematic reviewers who assessed the adequacy of the acupuncture judged that, when reported, the point selection, the needling technique, and the experience of the acupuncturists to be adequate in all four of these trials. For the reasons described above, it seems unlikely that the use of a flexible or fixed formula instead of individualized acupuncture in four of the five shamcontrolled trials is the reason for the lack of a benefit.

Alternative explanations for the negative results in these trials might include an inadequate number of treatment sessions, an insufficient duration of treatment, or an inadequate treatment frequency. All sham-controlled trials were judged by our acupuncture adequacy assessors to have used an adequate number of treatment sessions and a sufficient duration of treatment. Only the Forbes 2005 sham-controlled trial was judged to use an inadequate treatment frequency because this trial involved only one acupuncture session per week (for 13 weeks), which even though judged inadequate, probably still well reflects clinical practice in Western countries. The other sham-controlled trials all used two sessions per week, which was judged by the acupuncture adequacy assessors as an adequate treatment frequency, so it seems unlikely that an inadequate frequency of treatments explains the lack of benefit. While one trial (Lembo 2009) did not meet the Rome criteria recommendations for a minimum treatment duration of four weeks (Irvine 2006), the acupuncturist adequacy assessors judged that this trial's three week treatment duration, with twice weekly treatments, was adequate. Although the acupuncture assessors judged the treatment frequency of the sham-controlled trials to be largely adequate, the Chinese language comparative effectiveness trials used a much greater treatment frequency, with daily acupuncture treatments used in 9 out of 11 of these comparative effectiveness trials, and in all five of these trials that compared acupuncture to drug treatment. The higher acupuncture treatment frequency in the Chinese comparative effectiveness trials, relative to the shamcontrolled trials, might also help explain the different benefits of acupuncture relative to the controls in these two subsets of trials.

Quality of the evidence

For the sham-controlled trials, the continuous outcomes symptom severity and quality of life were rated as 'moderate' quality using the GRADE criteria because of sparse data (i.e. less than 400 events). This indicates that further research could have an impact on our confidence in the estimates of effect and may change the estimates.

Four out of the five sham-controlled trials in this review (Forbes 2005; Schneider 2006; Anastasi 2009; Lembo 2009) did not have limitations related to a risk of bias criterion. Each of these four trials used adequate randomization (which was not reported in the publications but ascertained by contacting the corresponding authors), each adequately addressed incomplete outcome data, and each used a sham control that was likely to adequately blind participants to the treatment received. Only the Lowe 2000 trial used a sham intervention that might not have been sufficiently believable as true acupuncture, such that this sham could adequately blind study participants as to whether they were receiving true acupuncture or a sham. However, if the sham acupuncture group participants realized they were getting a sham treatment, then this unblinding to the treatment received would likely have resulted in an overestimate of the effects of true acupuncture in this trial (Higgins 2011a).

A potential methodological limitation is that two of the five sham-controlled RCTs (Forbes 2005; Anastasi 2009) used a sham control that involved skin penetrating needles inserted at nonacupuncture points, that the acupuncture assessors in our review judged to have potential weak physiological activity that might influence the outcome, and which might therefore have biased these two RCTs to the null. However, we would not expect this to explain the lack of benefit of acupuncture relative to sham, both because these two shams were judged to have potential for only weak physiological activity and also because the other three shamcontrolled RCTs (Lowe 2000; Schneider 2006; Lembo 2009) used non-penetrating shams that were judged to be unlikely to have physiological effects. These three RCTs also found no benefit of acupuncture relative to sham.

The quality of the evidence is also limited by the fact that all shamcontrolled trials except the Lembo 2009 study had small sample sizes and were each underpowered to detect a small benefit of the acupuncture protocol evaluated. Although these trials may have been adequately powered to detect a moderate to large benefit of acupuncture relative to sham, an effect size of this magnitude may have been unreasonable to expect, considering that specific 5HT₄ agonists (i.e. tegaserod) and 5HT₃ antagonists (i.e. alosetron and cilansetron), which are the only treatments with "good quality of evidence" for treating IBS (Brandt 2009) provide only a modest benefit. Although a meta-analysis of the five sham-controlled trials increases the statistical power to detect an effect, a limitation of pooling trials with different acupuncture protocols is that we cannot rule out the possibility that larger trials or meta-analyses focusing on one of these protocols might show a benefit of treatment. In addition, although the meta-analysis point estimates suggest no effect, the 95% confidence intervals include the possibility that there could be small benefits which could be important to patients. A final limitation of the shamcontrolled trial evidence base, related to the small sample sizes, and also the heterogeneity of participants, is that these trials did not restrict eligibility to specific subtypes of IBS patients, and the proportions of patients with different IBS subtypes differed across

trials. An individual patient data meta-analysis would be necessary to address whether acupuncture has different effects on different subtypes of IBS patients, although the relatively small numbers of patients would be unlikely to provide a confident answer to this question.

For the Chinese language comparative effectiveness trials the dichotomous outcome symptom severity was rated as 'low' quality using the GRADE criteria because of high risk of bias (e.g. none of the studies were blinded) and sparse data (i.e. less than 400 events). This indicates that further research is likely to have an important impact on our confidence in the estimate of effect and is likely to change the estimate.

In the Chinese language comparative effectiveness trials, in addition to the primary risk of bias associated with the absence of patient blinding, there were also risks of bias associated with the randomization procedure and the follow-up of patients. Notably, for five of these trials (Liu 1997; Chen 2009; Li 2009; Xue 2009; Liu 2010), there were equal sized treatment groups, and during our telephone surveys the trial investigators either stated that some participants were non-randomly assigned to treatment or could not adequately explain how the equal group sizes were achieved. This raises the possibility that the randomization might not have been adequately generated or concealed (Schulz 2002). The notion that randomized trials should have equal numbers in each treatment group has been shown to commonly lead clinical trial investigators to force equality by unscientific means (Schulz 2002). Indeed, previous methodological reviews of this issue have found that over one-half of trials using simple, unrestricted randomization schemes report equal numbers in each group (Adetugbo 2000; Schulz 2002), and 88% of reported randomized trials have been shown to exclude some randomized participants from their analysis (Adetugbo 2000). The Chinese trials with high risks of bias associated with randomization or the accounting of randomized patients in the outcomes assessments evaluated acupuncture as an adjuvant to either another Chinese medicine treatment or psychotherapy; or compared acupuncture to psychotherapy, probiotics, or a drug not indicated or commonly used for IBS (i.e. sulfasalazine (Xue 2009)). Therefore, the findings from these studies should be considered to be hypothesis generating, and are not included in our overall conclusions. In contrast, there was a lower overall risk of bias in the four comparative effectiveness trials that found acupuncture to be more effective than two antispasmodic pharmacological therapies shown to be effective for IBS (Jailwala 2000; Ruepert 2011) (i.e. pinaverium bromide (Chen 2011; Shi 2010; Sun 2011) and Trimebutine maleate (Zeng 2010)). These four studies were rated as high risk of bias due to lack of blinding but were otherwise methodologically sound studies.

Potential biases in the review process

Potential biases in the review process were avoided by conducting comprehensive searches to identify all relevant studies, conducting dual and independent data extraction, and blinding the acupuncture treatment adequacy assessors to the results of the trials they assessed. In addition, we contacted the corresponding authors of all potentially eligible, claimed "randomized" trials published in Chinese language journals, as well as English language RCTs that did not include details about randomization methods in their published reports, to confirm their authenticity.

Agreements and disagreements with other studies or reviews

The conclusions of this review have changed from the conclusions of our 2006 Cochrane review (Lim 2006) which is the only other systematic review focused on acupuncture for IBS. While our 2006 Cochrane review found that it is "inconclusive whether acupuncture is more effective than sham acupuncture", the current review concludes that currently available sham-controlled trials suggest that acupuncture is not superior to sham acupuncture for reducing symptom severity or improving quality of life in patients with IBS. Comparative effectiveness Chinese trials have found that patients report greater benefits from acupuncture than from pharmacological therapies. However, the results of the comparative effectiveness studies should be interpreted with caution due to risk of bias (lack of blinding) and expectation effects.

AUTHORS' CONCLUSIONS

Implications for practice

People with IBS have few treatment options available. Pharmacological therapies provide modest benefits (Ford 2009a), can have high costs, and some of the newer drugs have been withdrawn from the market because of adverse events (Thompson 2001; Pasricha 2007). Safe, non-pharmacological therapies that may allow patients to feel more empowered and more in control of their symptoms should be evaluated for effectiveness. However, evaluating complex non-pharmacological therapies for IBS (e.g. mindfulness meditation (Gaylord 2011) or hypnotherapy (Lindfors 2012)) poses challenges, particularly in regards to selecting a placebo control or a credible alternative treatment control.

While acupuncture can theoretically be compared with a sham acupuncture "placebo" control, a fundamental challenge has been developing a sham acupuncture control that is sufficiently believable to patients so as to be indistinguishable from true acupuncture, and yet at the same time not so similar to true acupuncture that the sham has a therapeutic effect of its own and is therefore not an inert placebo. The sham acupuncture controls used in four of the five sham-controlled trials in this review appeared to be believable as authentic treatments (Forbes 2005; Schneider 2006; Lembo 2009; Anastasi 2009), but two of the five sham-controlled trials used sham controls that might have had weak physiological activity (Forbes 2005; Anastasi 2009), and therefore these shams may not have been completely inert placebos. While none of the sham-controlled trials showed a benefit of acupuncture relative to sham acupuncture, it is still not clear whether these findings are because acupuncture has no true biological effect above and beyond a placebo; or whether instead acupuncture has small biological effects, but the small sample sizes and heterogeneity of participants and interventions in these trials precluded detecting a statistically significant pooled benefit of acupuncture over sham; or whether any biological effects of true acupuncture cannot be detected because they are overridden and obscured by the large placebo effects of the sham control (Kaptchuk 2006; Wechsler 2011). Evidence from four Chinese language comparative effectiveness trials (Shi 2010; Zeng 2010; Chen 2011; Sun 2011) showed acupuncture to be superior to two antispasmodic drugs (pinaverium bromide and trimebutine maleate), both of which have consistently been shown to provide a modest benefit in high quality trials (Jailwala 2000; Ruepert 2011), although neither is approved for treatment of

IBS in the United States (Jailwala 2000). Patient preferences and expectations may partly explain the positive findings of these trials comparing acupuncture to drug treatment. That is, if the trial participants had pretreatment preferences for acupuncture over drugs, these preferences may have influenced the participants' later assessments of their subjective states, as reported on the patient-reported outcome measures used (Kalauokalani 2001; Linde 2007; Manheimer 2007; O'Connell 2009).

In addition to efficacy, safety and costs are other considerations. Safety is best determined with large prospective surveys of practitioners and three such surveys (MacPherson 2001; White 2001; Melchart 2004) show that serious adverse events after acupuncture are rare. There was one adverse event associated with acupuncture in the nine trials that reported this outcome (Forbes 2005; Reynolds 2008; Anastasi 2009; Chen 2009; Lembo 2009; An 2010; Liu 2010; Shi 2010; Sun 2011), although relatively small sample sizes limit the usefulness of these safety data. Finally, patients would also need to consider costs because acupuncture treatment often needs to be paid for out of pocket.

Implications for research

Considering that our meta-analysis found no differences between acupuncture and sham, and also considering that there are limited resources available to conduct trials of acupuncture, a non-proprietary therapy, additional sham-controlled trials of acupuncture among IBS patients should not be a high priority in acupuncture research, at least until the large, ongoing shamcontrolled trial, which is expected to complete data collection in March 2013, is published (Anastasi). This trial (n = 171) (Anastasi) compares a sham control with two different acupuncture test treatment groups, one test group using a fixed formula and the other test group using an individualized treatment approach, for patients with diarrhea-predominant IBS (See Characteristics of ongoing studies). If this trial shows no benefit of acupuncture relative to the sham, then the need for additional sham-controlled trials would seem questionable. However, if this ongoing shamcontrolled trial shows a benefit, then it would certainly be warranted to conduct future sham-controlled trials building upon the results of this trial (e.g. restriction to diarrhea predominant IBS patients; using the same acupoints as used in this trial). Such future sham-controlled trials should use non-penetrating, but demonstrably credible, shams to control for placebo effects, and ideally these sham needles should be placed far away from the true acupuncture points.

Because of the difficulties of controlling for placebo effects in acupuncture for IBS trials, which typically evaluate strictly subjective, patient-reported outcomes (e.g. symptom severity, quality of life), another approach forward for research is the evaluation of objective or semi-objective outcomes in IBS patients, using pragmatic and cost-effectiveness trials. Indeed, a recently completed trial (n = 220) (Principal Investigator: MacPherson) compared the effectiveness and cost-effectiveness of acupuncture plus usual general practitioner (GP) care versus usual GP care alone, on the semi-objective outcomes of medication use, health service use, and days lost from work (MacPherson 2010). Although this trial does not include a placebo control, because the outcome measures being assessed in this trial are semi-objective, its results will be less influenced by expectation effects (Wood 2008; Hrobjartsson 2010; Manheimer 2011), than trials that assess only strictly subjective outcomes (i.e., patient reports of symptom improvement). Indeed,



the Rome criteria for design of IBS treatment trials note that placebo effects "are especially a problem where end points are subjective" (Irvine 2006). If this recently completed costeffectiveness trial shows that acupuncture reduces healthcare utilization, then whether the resulting cost-savings are due to a specific effect of acupuncture needling or non-specific effects (e.g. greater autonomy and empowerment of patients, positive patient-practitioner relationship) seems of secondary importance. However, it must be borne in mind that the patient population who elected to participate in this acupuncture trial may have stronger a priori beliefs about the benefits of acupuncture, than does the average population of IBS patients, and therefore, the non-specific effects experienced by the patients in this unblinded trial may not be generalizable to the results that would be obtained among an average population of IBS patients. However, the results of this trial may be generalizable to the subset of IBS patients in general practice who would elect to receive acupuncture because such patients may also have a priori expectations for acupuncture to be beneficial. To produce results generalizable to the average population of IBS patients, investigators of future pragmatic trials might minimize the recruitment of participants with an a priori preference for acupuncture by not specifying, in the recruitment of patients, that acupuncture is one of the treatment options being investigated.

Future comparative effectiveness trials would also be helpful to validate and extend the preliminary evidence in this review, which suggests that acupuncture is associated with greater improvements in subjective patient self-assessments than pharmacological therapies. As previously mentioned, a limitation of the acupuncture versus pharmacological therapy trials in this review is that they did not use a design that controlled for the effects of patients' expectations for improvement, patient preferences, and non-specific therapeutic factors. Indeed, in the Chinese trials included in this review, the patients may well have had pre-treatment preferences for acupuncture, considering that these trials were conducted at hospitals of traditional Chinese medicine. Because acupuncture may elicit a greater expectation effect than pharmacological therapies or other active treatments (Kaptchuk 2006; Manheimer 2007; O'Connell 2009), particularly among participants who have a preference for acupuncture, investigators conducting future trials that compare acupuncture with other active therapies should consider asking participants about their preferences and expectations (before and after the intervention), and studying the potential effects of pre-treatment preferences on study outcomes. Such trials should also include a credibility questionnaire to establish that the treatments being compared are perceived by the patients as equally credible treatments for IBS symptoms (Gaylord 2011). Future comparative effectiveness trials in the West should also consider using a daily frequency of acupuncture, as was used in the Chinese trials in this review. However, even with additional well-designed trials, the truth about the effects of acupuncture for IBS will likely always be difficult to assess because the complexities and potential biases inherent to both the comparative effectiveness and sham acupuncture control designs makes it difficult to evaluate the subjective, patient-reported outcomes typically used in IBS trials.

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CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

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Wu T, Li Y, Bian Z, Liu G, Moher D. Randomized trials published in some Chinese journals: how many are randomized?. *Trials* 2009;**10**:46.

* Indicates the major publication for the study

Methods	Design: Parallel Blinding: No Attempt to confirm patient blinding?: No				
	Dropouts/withdrawals: Not reported in the article				
	Total duration: 27 days treatment immediately followed by an evaluation, no follow up reported Type of analysis reported: NA				
Participants	Setting: Outpatients of Shanghai Xuhui District Fenglin Community Health Center and Shanghai Re- search Institute of Acupuncture and Meridian, Shanghai, China				
	Mean age (+/- Standard Deviation or Range): 43 (13) for acupuncture group, 42 (12) for bifidobacteria group				
	Men/Women (number/number): 31/50. Acu/Moxa: 18/29, Medication: 13/21				
	Recruitment method: Not reported				
	Minimum duration of IBS symptoms: Rome II criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis				
	Actual duration of IBS symptoms before enrollment: 9.12 (5.05) years for acupuncture group, 9.15 (5.22 years for bifidobacteria group				
	Diagnosis of IBS required for eligibility?: Rome II criteria				
	Evaluation to rule out organic gastrointestinal disease: Not reported Predominant symptoms or stool patterns of participants: Not reported				
	Hospital inpatients?: No Were people with a history of acupuncture treatment excluded?: Not reported				
Interventions	ACUPUNCTURE plus moxibustion Number allocated to acupuncture: 47 [The participant numbers in both groups were not the number randomized, but the number remaining at the end of treatment] Style of acupuncture: Chinese Points selection: Fixed formula Points stimulated: Bilateral ST 25 and ST 37 Description of acupuncturist qualifications: Not reported				
	Total length of treatment period (weeks): ~4 (27 days) Number of sessions target (mean): 24				
	Times per week: ~6 Number of points used (mean): 2 (bilateral 4) Insertion depth: Not reported Was De Qi [1] reportedly sought?: [Yes] Duration (minutes): [20 min] Method of stimulation: Manipulation				
	[1] De Qi means 'arrival of energy' and is a sensation of numbness or distension sometimes generated by stimulating acupuncture needles by hand or with an electrical current. According to acupuncture theory, activation of De Qi may be one indication that acupuncture is exerting its beneficial effects				
	Moxibustion: Mild moxibustion on bilateral ST 25 only for 20 min and acupuncture on ST 37 only				
	CONTROL GROUP A INTERVENTION: Bifidobacterium (Product name: Pei Fei Kang). The English name for "Pei Fei Kang" is Live Combined <i>Bifidobacterium Lactobacillus</i> and <i>Enterococcus faecium</i> capsules				
	Number allocated to control group A: 34 Total length of treatment period: 4 weeks Number of sessions target (mean): 28 Days per week: 7				

An 2010 (Continued)							
	Times per day: 3 Dosage: 2 pills						
		all groups?: "Patients in both groups were given suggestions for regulating ther medications were stopped during treatment"					
Outcomes	Symptom severity: The	Symptom severity: The overall effect					
	All the symptoms were observed, including abnormal stools (abnormal frequency, number of defe- cation and aspect, three sub-items), abdominal pain (frequency, intensity, duration), abdominal dis- tension or abdominal discomfort (frequency, intensity, duration), borborygmus, defecation incom- plete sensation, bowel urgency, mucus stools, poor appetite, epigastric discomfort or pain, fatigue, hot flashes with night sweats, psychological disorders – 12 items used to calculate an overall IBS symptom score.						
	Overall IBS symptom se	core (3-point scoring system): 1) improved 2) effective 3) no effect					
	ment * 100%. 1) markedly effective: t 2) effective: the freque	fore treatment - symptom score after treatment) / symptom score before treat- the defecation became normal and other symptoms were relieved. $R > = 70\%$. ncy of defecation decreased and other symptoms improved, $30\% = < R = < 70\%$ ncy of defecation did not decrease obviously and other symptoms did not im-					
	Quality of life: Not assessed						
	Time points of symptom measurement: The responder rates of overall IBS symptom score were report- ed after 4 weeks treatment						
	Author's conclusion: "Acupuncture and moxibustion are more effective than medication for treating IBS. Acupuncture and moxibustion are a safe and effective way to treat IBS"						
	Additional outcomes reported in the trial but not extracted: The improvement of twelve symptoms items including defection, abdomen pain, abdominal distension or discomfort, stomach rumbling, feeling of unfinished defection, feeling of urgent need to defect, mucus stool, indigestion and loss of appetite, stomach discomfort or distending pain, asthenia, hot flushes and night sweats and mental aberration were reported after 4 weeks treatment Adverse effects: "No adverse events were observed"						
Notes	Comments:						
	Funding sources: Project of Xuhui District Health Bureau of Shanghai (SHXH20070402)						
	randomization: Referring to a random number table						
	Allocation: [Sequentially numbered, opaque, sealed envelopes]						
	Additional information in square brackets was provided by the first author An GQ, in a telephone survey on September 1, 2011						
	We dichotomized improved, effective and no effect into the 2 categories of improved or not improved according to the methods section of the review						
Risk of bias							
Bias	Authors' judgement	Support for judgement					
Random sequence genera- tion (selection bias)	Low risk	"Referring to a random number table, we assigned 47 participants into acupuncture-moxibustion group and 34 participants into bifidobacteria group"					

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Trusted evidence. Informed decisions. Better health.

An 2010 (Continued)

Allocation concealment (selection bias)	Low risk	[Sequentially numbered, opaque, sealed envelopes]
Blinding (performance bias and detection bias) All outcomes	High risk	Single blind: The evaluators were blind to the participant assignment
Incomplete outcome data (attrition bias) Short term	Unclear risk	Dropouts were not reported in the article. [There were dropouts during the 4 weeks of treatment, but the author could not recall the number of the dropouts in both groups]
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall IBS symptom score at the completion of the 4 week treatment period, together with results for each outcome measured
Other bias	Unclear risk	ITT analysis was not reported and the author also could not recall the number of dropouts, so we could not estimate whether the "missingness" would affect the treatment effect in this trial
		"The two groups were comparable at baseline in terms of sex (P = 0.084), age (P = 0.389) and disease duration (P=0.708)." There was no description or P value reported for the comparison of symptom score between two groups at baseline

Anastasi 2009 Methods **Design: Parallel** Blinding: Yes. "The SF monitored and recorded Acu/Moxa point sequencing and duration and maintained blinding. The PI, data manager, statistician, DA, and study coordinator were all blinded to study assignment until the final statistical analysis." Also, "all subjects were blindfolded during treatment..."Attempt to confirm patient blinding?: No Dropouts/withdrawals: During the 4 weeks, 2/14 drop-outs in acupuncture/moxibustion group and 0/15 drop-outs in the sham group. An additional 1/14 participants in the acupuncture/moxibustion group and 3/15 in sham acupuncture/moxibustion group did not complete the symptom diary at end of follow-up, however the symptom diary was not an outcome for this review Total duration: 4 weeks of treatment, evaluation at 3 weeks and 4 weeks after baseline (symptom diary) Type of analysis reported: Not specified Participants Setting: outpatient research lab, USA Mean age (+/-SD or Range): 40.4 (34.6 to 46.2) Men/Women (n/n): 10/19 Recruitment method: "Subjects were recruited by the study team by informing GI practitioners, internists, and nurse practitioners in the New York City area about the study. Advertisements were also posted in select New York newspapers and community boards. Potential subjects responding to advertisements or practitioner referrals were screened by the primary investigator (PI) and research staff for eligibility" Minimum duration of IBS symptoms: 6 months Actual duration of IBS symptoms before enrollment: Not stated Diagnosis of IBS required to be eligible? (if yes, describe how patients were verified to have IBS): Rome III criteria Evaluation to rule out organic gastrointestinal disease: "Subjects had to have verification from their medical provider confirming their IBS diagnosis and absence of pathology confirmed by diagnostic workup' Predominant symptoms or stool patterns of participants: Not reported Hospital inpatients? (Y/N; if Y list number inpatients): No

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Anastasi 2009 (Continued)	Were people with a history of acupuncture treatment excluded?: Yes, if they had received acupuncture during the past 6 months
Interventions	ACUPUNCTURE/MOXIBUSTION N allocated to acupuncture: 14 Style of acupuncture: Chinese Points selection: Flexible formula for specific symptoms Points stimulated: The active acupuncture and moxibustion (Acu/Moxa) protocol consisted of eight core points (CV-12, CV-6; bilateral ST-25, BL-23, BL-25) and additional points (two-six additional needles selected from a table of additional points for the TCM diagnoses listed above)
	Description of acupuncturist qualifications: "All TAs [treating acupuncturists] were formally trained in TCM and were nationally certified and state licensed.
	Total length of treatment period (weeks): 4 Number of sessions target (mean): 8 Times per week: 2 Number of points used (mean): 10 to 14 points Insertion depth: "needle insertion at the appropriate depth" Was De Qi reportedly sought?: Yes. "The TA performed the reinforcing/tonification technique according to classic text, by needle insertion at the appropriate depth and needle rotation to elicit qi" Duration (mins): 60 min "All subjects attended 4 weeks of 1-hour, twice-weekly treatments" Method of stimulation: Manipulation
	Moxibustion procedure: "All moxibustion procedures were performed according to standard indi- rect techniques as described in the classic text (BCTCM, 1987). Point stimulation involved a burning, cigar-shaped moxa pole held approximately 1 inch over the acupuncture points. The moxa pole was moved in a clockwise circular motion directly over each point, stimulating each for a period of 2 min- utes or until the skin around the area of the point became pink (BCTCM, 1987)"
	CONTROL GROUP A INTERVENTION (sham, if used): Sham acupuncture/moxibustion with penetrating needles N allocated to control group A: 15 Points selection: Flexible
	Points stretction: receiver Points stimulated: "superficial needling at nonpoints according to previously established controls (Hammerschlag, 1997; Vincent, 1989)." "Sham points were 2 to 3 cm away from the traditional true point location and not on a meridian, with an insertion depth of 1 to 2 mm, sufficient to make the nee- dle stand vertically with very light stimulation without elicitation of qi"
	Total length of treatment period: 4 weeks Number of sessions target (mean): 8 Times per week: 2 Number of points used: 10 to 14 Insertion depth: 1 to 2 mm Was De Qi sought?: No Duration (mins): 60 Method of stimulation: Manipulation - "Very light stimulation"
	Sham moxibustion procedure: "Placebo moxibustion was performed by holding the burning moxa pole approximately 8 in. above and 2 to 3 cm away from traditional location for a period of 2 minutes. Special and meticulous attention was made on the part of the acupuncturist to not generate a heat sensation"
	Any co-interventions in all groups?: None stated
Outcomes	Symptom severity: "The Clinical Global Impression Scale (CGIS) measures the patient-perceived sever- ity of symptoms with a 7-point Likert-type scale ranging from 0 (no discomfort) to 6 (very severe dis- comfort).
	Quality of life: Not assessed

Anastasi 2009 (Continued)	
(continued)	Time points of symptom measurement: "The CGIS was administered at baseline before the interven- tion and again at completion of the 4-week treatment period"
	Author's conclusion: "The findings indicate that Acu/Moxa treatment shows promise in the area of symptom management for IBS"
	Additional outcomes reported in the trial but not extracted: Symptoms of 1) Abdominal pain/discom- fort, and 2) intestinal gas, bloating, and stool consistency
	Adverse effects: "Among the 29 subjects in this eight-session study, no adverse events were reported"
Notes	Comments: Carefully worked, especially for the blinding of subjects
	Weak points: Small sample size
	Standard deviations for CGIS at pretreatment and post-treatment were not reported. We used the con- fidence interval for the mean difference to calculate post-treatment standard deviations
	The first author Joyce Anastasi was contacted by e-mail to request information about randomization, allocation concealment and CGIS. She replied with detailed information on February 10, 2010
	Funding sources: Not stated
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Description from text: "Once deemed eligible, subjects were randomly as- signed to either the experimental group or the control group"
		[The statistician used SAS Proc Plan to develop a scheme of randomly permut- ed blocks: codes labelled # 1 or # 2 were balanced within blocks of 2, 4 or 6 and blocks randomly intermingled]
Allocation concealment (selection bias)	Low risk	[We assured that blinded and masked allocation of intervention was preserved during the study. These codes were delivered in a masked log sheet, consecu- tively numbered, again with only codes #1 or #2 (not the meaning of the code). The log sheet was kept in a binder (in a locked cabinet) for a blinded study co- ordinator with no involvement with the facilitation of the treatments. Once it was determined that the subject was eligible, the study facilitator contacted a blinded study coordinator to receive the code. The study facilitator did not have access to the randomization sequence log sheet. Upon receiving the code (#1 or #2), the study facilitator would translate the code into the actual treat- ment condition for the treating acupuncturist. The principal investigator, da- ta manager, statistician, diagnostic acupuncturist and the study coordinator were blinded to the treatment condition code. It was only after, the comple- tion of the last study subject and all data were entered and analyzed, the treat- ment condition was revealed (that is the corresponding treatment associated with codes #1 and #2)]
Blinding (performance bias and detection bias) All outcomes	Low risk	"The SF monitored and recorded Acu/Moxa point sequencing and duration and maintained blinding. The PI, data manager, statistician, DA, and study co- ordinator were all blinded to study assignment until the final statistical analy- sis"
		"All subjects were blindfolded during treatment"
		"Subjects received sham acupuncture by superficial needling at nonpoints according to previously established controls (Hammerschlag, 1997; Vincent, 1989). The sham points were as anatomically precise as an active treatment protocol so that they were reproducible and consistent. Sham points were 2 to 3 cm away from the traditional true point location and not on a meridian,

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Anastasi 2009 (Continued)		
		with an insertion depth of 1 to 2 mm, sufficient to make the needle stand verti- cally with very light stimulation <i>without</i> elicitation of <i>qi</i> . This procedure "min- imizes the specific effects of the needling while maintaining its psychological impact" (Vincent & Lewith, 1995) and has been assessed as a credible and ef- fective control procedure (Birch, Hammerschlag, Trinh, & Zaslawski, 2002; Stux & Hammerschlag, 2001; Vincent 1989; Vincent & Lewith 1995)."
		"Placebo moxibustion was performed by holding the burning moxa pole ap- proximately 8 inches above and 2 to 3 cm away from traditional location for a period of 2 minutes. Special and meticulous attention was made on the part of the acupuncturist to not generate a heat sensation. Both true treatment and sham/placebo sessions were identical in duration"
		Although one of the exclusion criteria in this trial was no use of acupunc- ture in the past 6 months, the patients in this trial were not required to have <i>never</i> used acupuncture (i.e. to be acupuncture naive) (i.e. "Anyone receiv- ing acupuncture over the past 6 months with a history of receiving moxibus- tionwas not enrolled.") Also, there were no tests for checking the success of the blinding. However, because a needle insertion sham was used and the needles were inserted close to the genuine acupuncture points, and because this procedure has reportedly been assessed as a credible and effective con- trol procedure, we judged that this sham was likely to be indistinguishable from the true acupuncture
Incomplete outcome data (attrition bias)	Low risk	"One withdrew, one lost to follow-up, and four did not submit final symptom diary"
Short term		At 4 weeks, 2/14 (14%) participants in the true acupuncture/moxibustion group did not complete the Clinical Global Impression Scale. All (15/15) partic- ipants in the sham control group completed the 4 week Clinical Global Impres- sion Scale
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall IBS symptom severity at the comple- tion of the 4 week treatment period, together with results for each outcome measured
Other bias	Low risk	Although ITT analysis was not reported, there were few drop-outs, so the de- gree of "missingness" would be unlikely to affect the estimate of the treatment effect in this trial. The two treatment groups were comparable at baseline in terms of the variables "abdominal pain/discomfort", "intestinal gas, bloating, and stool consistency" and "Clinical Global Impression Scale." The two treat- ment groups also appeared comparable at baseline in terms of demographic characteristics. However, there were no statistical tests or P values reported for these comparisons

Chan	2000
Chen	2009

Methods	Design: Parallel
	Blinding: No
	Attempt to confirm patient blinding?: No
	Dropouts/withdrawals: Not reported in the article [In order to ensure the number of 30 participants for both groups, the participants who withdrew during the 4 weeks treatment were eliminated from the tri- al and new participants were recruited. The author could not recall the number of the dropouts]
	Total duration: 4 weeks treatment followed by an evaluation, no follow up reported Type of analysis reported: Not reported



Chen 2009 (Continued)	
Participants	Setting: Outpatient and inpatient of the First Affiliated Hospital of Guangzhou University of Traditional Chinese Medicine from Jan 2005 to May 2006, Guangzhou, China
	Mean age (+/- Standard Deviation or Range): Not reported. The age range for acupuncture plus patent- ed Chinese medicine group was 18 to 63 years; the age range for patented Chinese medicine alone group was 18 to 65 years
	Men/Women (number/number): 27/33
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Rome II criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis
	Actual duration of IBS symptoms before enrollment: Not reported. The duration range for acupuncture plus patented Chinese medicine group was 14 months to 12 years; the duration range for patented Chinese medicine alone group was 12 months to 15 years
	Diagnosis of IBS required for eligibility?: Rome II criteria and diagnostic criteria for liver-qi stagnation with spleen deficiency according to the <i>Guiding Principles for Clinical Study of New Chinese Medicine</i>
	Evaluation to rule out organic gastrointestinal disease: Yes Predominant symptoms or stool patterns of participants: Not reported
	Hospital inpatients?: Yes, the number inpatients was not reported
	Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	ACUPUNCTURE (Warming needle moxibustion) plus patented Chinese medicine Number allocated to acupuncture: 30 Style of acupuncture: Chinese warming needle Points selection: Flexible formula Points stimulated: 2 main fixed acupoints and 2 to 5 optional points. The fixed points were bilateral ST 25 and BL 25. The optional points were HT 7, PC 6, LR 3 and SP6 for restlessness, insomnia, anxiety and suspiciousness; ST 37, GV 20, GV 1 and BL 57 for a sense of incomplete emptying and down-bearing sen sation in the anus; CV 12, ST 36, SP 6, LR 3, SP 9 for abdominal pain or distension, bowel sounds and se vere diarrhea; ST 37 and SJ 6 for constipation Description of acupuncturist qualifications: Not reported
	Total length of treatment period (weeks): 4 Number of sessions target (mean): 12 Times per week: 3 Number of points used (mean): 2 fixed acupoints plus 2 to 5 optional points Insertion depth: Not reported Was De Qi reportedly sought?: Yes Duration (minutes): 30 min Method of stimulation: Manipulation, even reinforcing - reducing. Warming needle moxibustion only on bilateral ST 25 and BL 25, acupuncture on other optional acupoints
	Warming needle moxibustion means using needles to puncture in the acupoint first, then put a moxa cone on the needle handle and burn it. Co-intervention: Patented Chinese medicine: <i>Kang Tai</i> Capsule (previously known as <i>Shun Ji</i> Decoction), 3 capsules, 3 times a day
	CONTROL GROUP A INTERVENTION: Patented Chinese medicine alone (Product name: <i>Kang Tai</i> Cap- sule (previously known as <i>Shun Ji</i> Decoction)) Number allocated to control group A: 30 Total length of treatment period: 4 weeks Number of sessions target (mean): 28 Days per week: 7 Times per day: 3 Dosage: 3 capsules / time



Chen 2009 (Continued)

Any co-interventions in all groups?: Both groups were administered patented Chinese medicine

Outcomes	Symptom severity:		
	 IBS symptom score for TCM symptom (4-point scoring system): 1) cured 2) improved 4) effective 4) no effect N (Efficacy index) = (symptom score before treatment - symptom score after treatment) / symptom score before treatment * 100% 1) Recovery: N > = 90% 2) Markedly effective: 70% = < N = < 90% 3) Improvement: 30% = < N = < 70% 4) Failure: N = < 30% Quality of life: Functional gastrointestinal disease and quality of life scale created by Prof. Zhou FS, et al included assessment of field of disease, field of physiology, field of psychology, field of society and total score. There are 44 items in the QOL questionnaire. This does not appear to be a validated scale Time points of symptom measurement: The responder rates of IBS symptom score for TCM symptom were reported after 4 weeks treatment. The quality of life score in total and for each field were reported before and after treatment Author's conclusion: "Both treatment methods could improve the clinical symptoms and increase the patient's quality of life. The total effective rate in the treatment group was slightly higher than the control group" Additional outcomes reported in the trial but not extracted: Symptom scales for 9 items including abdominal paindegree, frequency of abdominal pain, abdominal distension, upset and insomnia, anxiety and suspicion, loose stool, feeling of unfinished defection, asthenia and loss of appetite before treatment, 2 weeks and 4 weeks Adverse effects: "No adverse effects appeared in all participants" 		
Notes	Comments:		
	Funding sources: [No funding]		
	randomization: [Drawing of lots]		
	Allocation: [No specific method was used to conceal the allocation]		
	Reason for identical numbers assigned to the two groups: [The author said if the patient number of one group started to exceed 30, they assigned the additional patients into the other group. The number of patients assigned in this non-random manner was not available from the author]		
	Information in square brackets was provided by the first author Chen YQ, in a telephone survey on July 23, 2011		
	We dichotomized cured, improved, effective and no effect into the 2 categories of improved or not im- proved according to the methods section of the review		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	High risk	"Sixty IBS cases of liver-qi stagnation with spleen deficiency were randomized into a treatment and control group by single-blind method"	
		[We used drawing of lots. If the number of patient of one group started to exceed 30, they assigned the additional patients into the other group]	
Allocation concealment (selection bias)	High risk	[No specific method was used to conceal the allocation]	

Acupuncture for treatment of irritable bowel syndrome (Review)

Chen 2009 (Continued)	nen 2009 (Cor	ntinued)
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Blinding (performance bias and detection bias) All outcomes	High risk	[Quality of life scales were evaluated by participants themselves. The IBS symptom score for TCM symptom were evaluated by the first author who knew the group assignment of the participants]
Incomplete outcome data (attrition bias) Short term	Unclear risk	The dropouts were not reported in the article [In order to ensure the number of 30 participants for both groups, the partici- pants who withdrew during the 4 weeks treatment were eliminated from the trial and new participants were recruited. The first author could not recall the number of the dropouts
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for IBS symptom score for TCM symptom at the completion of the 4 week treatment period and quality of life scale before and after treatment
Other bias	Unclear risk	ITT analysis was not reported and the author also could not recall the number of dropouts, so we could not estimate whether the "missingness" would affect the treatment effect in this trial. "There was no significant difference in gender, age and duration between the
		two groups." The P values were 0.417 for gender, 1.0 for age and 0.919 for dura- tion between the two groups
		There were no P values reported for the comparison of quality of life scale and each symptom score between two groups at baseline, but according to the mean(SD) of quality of life scale in Table 8 and symptom score for each item in Table 4 in Chen YQ 2006 Master's thesis, the 2 groups appeared comparable at baseline.

Chen 2011

Methods	Design: Parallel Blinding: No Attempt to confirm patient blinding?: No Dropouts/withdrawals: Not reported in the article [one withdrew from acupuncture group] Total duration: 3 weeks treatment followed by an evaluation, no follow up reported Type of analysis reported: NA
Participants	Setting: Acupuncture outpatients of the First Affiliated Hospital of Guangzhou University of Traditional Chinese Medicine and digestion medicine outpatient of the Second Hospital of TCM, Guangzhou, China
	Mean age (+/- Standard Deviation or Range): Not reported. 14 patients in the acupuncture group and 14 patients in the Western medication control group were aged 18 to 35 years; 10 patients in the acupunc- ture group and 12 patients in the Western medication group were aged 36 to 50 years; and 5 patients in the acupuncture group and 4 patients in the Western medication control group were aged 56 to 70 years.
	Men/Women (number/number): 27/32 14/15, Medication: 13/17
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Rome III criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis
	Actual duration of IBS symptoms before enrollment: The duration for acupuncture group was 13.44 (7.63) months; the duration for western medication group was 13.36 (7.79) months



hen 2011 (Continued)	
	Diagnosis of IBS required for eligibility?: Rome III criteria and diagnostic criteria for 4 different TCM syn drome differentiations according to the <i>Traditional Chinese Internal Medicine (Version 7)</i> and <i>Integrated</i> <i>Traditional Chinese and Western Internal Medicine</i>
	Evaluation to rule out organic gastrointestinal disease: Yes Predominant symptoms or stool patterns of participants: Diarrhea-predominant IBS
	Hospital inpatients?: No Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	ACUPUNCTURE (plus mild moxibustion) Number allocated to acupuncture: 29 [30 participants were assigned to each treatment group, one pa- tient withdrew from the acupuncture group during treatment] Style of acupuncture: Chinese Points selection: Flexible formula Points stimulated: 4 main fixed acupoints and 1 to 2 optional points. The fixed points were bilateral ST 25 and ST 37, single GV 20 and CV 4. The optional points were ST 36 for deficiency of the spleen and stomach; BL 23 and GV 4 for Yang deficiency of spleen and kidney; LR 4 and ST 36 for stagnation of the Liver-Qi attacking the spleen; SP 9 for wetness-heat stagnated in the middle energizer Description of acupuncturist qualifications: Not reported
	Total length of treatment period (weeks): 3 Number of sessions target (mean): 15 Times per week: 5 Number of points used (mean): 4 fixed acupoints plus 1 to 2 optional points Insertion depth: Not reported Was De Qi reportedly sought? Yes
	Duration (minutes): 30 min Method of stimulation: Manipulation, reinforcing and reducing. Mild moxibustion on GV 20, CV 4 and S ⁻ 25. Mild moxibustion on BL 23 and GV 4 for Yang deficiency of spleen and kidney
	CONTROL GROUP A INTERVENTION: Montmorillonite was given to all patients and the loperamide and pinaverium bromide was added if the diarrhea did not stop
	Number allocated to control group A: 30 Total length of treatment period: [3 weeks] Number of sessions target (mean): 21 Days per week: 7 Times per day: smecta: 2 times/day, loperamide and pinaverium bromide: 3 times/day Dosage: 1 bag/time for smecta, 4 mg/time for loperamide and 50 mg/time for pinaverium bromide
	Any co-interventions in all groups? Not reported
Outcomes	Symptom severity:
	 Overall IBS symptom score (4-point scoring system) referring to <i>the guide for clinical trials of new drugs of Chinese medicine</i> at end of treatment: 1) cured 2) improved 3) effective 4) no effect. 1) cured: the symptom disappeared. Effect index=100%. 2) improved: the symptom obviously improved. 100%>Effect index >=75%. 3) effective: the symptom improved. 75%>Effect index >=30%. 4) no effect: the symptom did not improved. Effect index =<30%. Computation formula: Effect index = (symptom score before treatment - symptom score after treatment) / symptom score before treatment * 100%
	Quality of life: Not assessed
	Time points of symptom measurement: The responders for overall IBS symptom were reported after 3 weeks treatment
	Author's conclusion: Although acupuncture is slightly better than conventional western medication fo overall IBS symptom, there was no statistically significant difference (P > 0.05)

Chen 2011 (Continued)	Additional outcomes reported in the trial but not extracted: Symptom score for abdominal pain and abdominal distension, score for defecation frequency, score for stool characteristics before and after treatment were reported respectively in Table 5, 6 and 7; responder rates based on TCM syndrome differentiation, responder rates based on age, responder rates based on disease duration Adverse effects: Not reported
Notes	Funding sources: [The research was supported partly by my advisor, myself and the university.]
	randomization: "We used simple randomization method. The sample size was entered into the JMTJFX statistical software then the random number table was obtained."
	Reason for identical numbers assigned to the two groups: [The participants number assigned to both group were 30 respectively. I entered the sample size into the randomization program in JMTJFX, it automatically generated the randomized group assignment]
	Evaluation blinding: "All outcome data came from the daily record card filled by participants"
	Information in square brackets was provided by the author Chen QM, in a telephone survey on December 4, 2011
	We dichotomized cured, improved, effective and no effect into the 2 categories of improved or not im- proved according to the methods section of the review

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"We used simple randomization method. The sample size was entered into the JMTJFX statistical software then the random number table was obtained."
		[The participants number assigned to both group were 30 respectively origi- nally. I entered the sample size into the randomization program in JMTJFX, it automatically generated the randomized group assignment]
Allocation concealment (selection bias)	Low risk	[We used envelope method. The envelopes were sequentially numbered, opaque, sealed]
Blinding (performance bias and detection bias) All outcomes	High risk	"All data came from the daily record card filled by participants"
Incomplete outcome data (attrition bias) Short term	Low risk	[One withdrew from the acupuncture group during the treatment]
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall IBS symptom responders at the com- pletion of the 3 week treatment period, together with results for each outcome measured as described in the methods section
Other bias	Low risk	Although ITT analysis was not reported, there were few drop-outs, so the de- gree of "missingness" would be unlikely to affect the estimate of the treatment effect in this trial. The two groups were comparable at baseline in terms of sex, age and disease duration (P > 0.05). The two treatment groups were not signifi- cantly different at baseline in terms of abdominal pain, abdominal distention, defecation frequency and stool characteristics

Methods	Design: Parallel Blinding: Yes Attempt to confirm patient blinding?: Not reported Dropouts/withdrawals: 1/60 before randomization, 8/59 after randomization. Total of 59 participants randomized and followed Total duration: 10 weeks of treatment with a weekly evaluation (symptom diaries), follow-up evalua- tion at 13 weeks after baseline (or prior exit) Type of analysis reported: Intention to treat analysis
Participants	Setting: Hospital outpatient, UK Mean age (+/-SD or Range): 43.0 (range:19 to 67) for acupuncture group, 44.4 (range:17 to 79) for con- trol group Men/Women (n/n): 20/39 Recruitment method: Personal approach in the hospital clinics Minimum duration of IBS symptoms: 3 months
	Actual duration of IBS symptoms before enrollment: Not stated Diagnosis of IBS required to be eligible? (if yes, describe how patients were verified to have IBS): Man- ning and Rome II criteria Evaluation to rule out organic gastrointestinal disease: Reported Predominant symptoms or stool patterns of participants: Pain was the predominant symptom (21/59) diarrhea (16/59) was more common than constipation (4/59)
	Hospital inpatients? (Y/N; if Y list number inpatients): No Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	ACUPUNCTURE N allocated to acupuncture: 27 Style of acupuncture: Chinese Points selection: Flexible formula for specific symptoms Points stimulated: Names of points were not stated Description of acupuncturist qualifications: "fully trained in traditional Chinese acupuncture and ac- credited to the British Acupuncture Council"
	Total length of treatment period (weeks): 10 Number of sessions target (mean): 10 Times per week: 1 Number of points used (mean): 4 to 8 points (8 to 16 needles for both sides) Insertion depth: Not stated Was De Qi reportedly sought?: Yes Duration (mins): ~25 min Method of stimulation: Manipulation
	CONTROL GROUP A INTERVENTION (sham, if used): Sham acupuncture with penetrating needles N allocated to control group A: 32 Points selection: formula Points stimulated: Nonacupuncture points in three different areas on the body Total length of treatment period: 13 weeks
	Number of sessions target (mean): 10 Times per week: 1 Number of points used: Not reported Insertion depth: Not reported Was De Qi sought?: No Duration (mins): ~25 Method of stimulation: Manipulation
	Any co-interventions in all groups?: Specific dietary and lifestyle advice
Outcomes	Symptom severity: "Patients completed weekly symptom diariesThe symptom diary is based on the Bristol scale[15] and has been previously validated[16]. It employs eight Likert scales relating to ab- dominal symptomatology and their interference with activities (Appendix), and permits the compi-

Acupuncture for treatment of irritable bowel syndrome (Review)



Forbes 2005 (Continued)	lation of a numerical (non-linear) global "symptom score" between 0 and 30The principal outcome measure was a change in the symptom score. A reduction of four is considered indicative of a clinically useful response"
	Quality of life: EuroQol instrument, measured as change in score from baseline for each group
	Time points of symptom measurement:
	Global symptom score at baseline (Table 1) and 13 weeks after baseline (Results section); Number of patients with symptoms score reduction at 13 weeks after baseline (Results section); Number of patients with symptoms score reduction at least 4 points (number of responders) at 13 weeks after baseline (Results section); EuroQol at baseline (Table1) and 13 weeks after baseline (Results section);
	Author's conclusion: Acupuncture is relatively ineffective for IBS Additional outcomes reported in the trial but not extracted: Blinded clinician assessment of global sta- tus as "improved", "unchanged" or "worse" Adverse effects: "No direct adverse events were elicited. One patient was commenced on antidepres- sants after psychiatric consultation, and another on beta-blockers for anxiety. Six other patients failed to complete the study period. Of these eight withdrawals, five belonged to the acupuncture group and three to sham"
Notes	Comments: Carefully worked, especially for the blinding of evaluator
	Weak points: Small sample size
	SDs for quality of life outcome at end point were not reported, we estimated the standard deviation as 25% of the range (1-100), i.e. 25
	For responder rate of Symptom severity, the denominator for the response rate are the number avail- able for follow-up at that time-point, not the number randomized
	The first author Forbes was contacted by email to request information about allocation concealment. He replied with detailed information on January 27, 2011
	Funding resources: "Supported by the a donation gratefully received from the T-R Golden Charitable Trust"
Risk of bias	

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"Recruited patients were allocated according to computer-generated random numbers concealed in sealed envelopes"
Allocation concealment (selection bias)	Low risk	"Recruited patients were allocated according to computer-generated random numbers concealed in sealed envelopes"
		[The envelopes were opaque and they were numbered (according to the recruitment number for the study) so that we knew which one to open next]
Blinding (performance bias and detection bias) All outcomes	Low risk	"'Sham' acupuncture points were chosen from three different areas on the body (the anterior thigh distally, the posterior thigh, and the lateral aspect of the lower back), which do not correspond to recognized acupuncture points and are deemed to have no therapeutic value. These were varied to some de- gree each week as in genuine treatment, including variation in needle tech- nique and length of time of retention. No deqi needling sensation was sought or obtained at sham points" "The second "treating acupuncturist" (TA) for the duration of the study re- mained the only individual aware of treatment allocation"

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orbes 2005 (Continued)		There was no mention in the eligibility criteria of whether participants were re quired to be acupuncture-naïve, so we assume this was not a criterion for par- ticipation. Also, there was no attempt to confirm patient blinding. However, because a needle insertion sham was used, and the points and needling tech- nique were varied somewhat each week, as was also done in the true acupunc ture group, who received individualized point selection, we judged that this sham was likely to be indistinguishable from the true acupuncture
Incomplete outcome data (attrition bias)	Low risk	"Sixty patients (40 female) were recruited; one female withdrew after recruit- ment but before treatment"
Short term		At 13 week follow-up, "One patient was commenced on antidepressants after psychiatric consultation, and another on beta-blockers for anxiety. Six other patients failed to complete the study period. Of these eight withdrawals, five belonged to the acupuncture group and three to sham"
		During the 13 weeks, 5(27) drop-outs in acupuncture group and 3(32) in sham acupuncture group. Although this trial reported withdrawals or loss to fol- low-up of approximately 13%, we did not consider this an important source of bias because the drop-outs were approximately evenly distributed across treatment groups, the withdrawals were unlikely to be related to knowledge of treatment assignment or effects of the treatment, and the degree of 'miss- ingness' would be unlikely to affect the estimate of the treatment effect in this individual trial or in the meta-analytic estimates. The authors reported having conducted an ITT analysis but did not specify whether outcomes from partic- ipants who had withdrawn from treatment were available or unavailable for inclusion, or how the outcome values, if unavailable, had been imputed for inclusion in the ITT analysis. The authors reported, however, that when the analysis was restricted to those completing the intended therapy, the results were unchanged from the ITT analysis reported in the publication
Selective reporting (re- porting bias)	Low risk	"Patients completed weekly symptom diaries, supplemented by psychologica and quality of life scales at entry and at 12th wk (or prior exit) Patients also completed the Hospital Anxiety and Depression (HAD) scale[17], and the Euro- Qol[18] quality of life questionnaire." "The principal outcome measure was a change in the symptom score" "Secondary outcome measures included assessment by one of the blinded in- vestigators at the end of the study period, which made a simple categorization of patient status as "improved", "unchanged" or "worse"
		Symptom score and EuroQol LAS score were reported at baseline and 13 weeks The median score of pain, bloating, diarrhea and constipation for those with the respective four symptom predominance at enrollment were reported at baseline and 13 weeks HAD score was reported at baseline Number with reduction in symptom score and number with reduction in symptom score by at least four points, assessment by the blinded clinician, comparison of HAD scores between groups were reported at 13 weeks
		This trial reported outcomes for overall IBS symptom severity and health-re- lated quality of life at the completion of the 13 week treatment period, togeth- er with results for each outcome measured according to the Methods section
Other bias	Low risk	"Results were analyzed by intention to treat"
		"There was a good balancing of the two groups: for no parameter was there a statistically significant difference at study entry"

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Methods	Design: Parallel Blinding: Yes	
	Attempt to confirm patient blinding?: Not stated but the author stated that Streitberger needle, which was used in this trial, has previously been validated in other studies Dropouts/withdrawals: During the second 3 weeks, 8/78 drop-outs in acupuncture group and 5/75 in sham acupuncture group Total duration: 3 weeks treatment immediately followed by an evaluation, no follow up reported Type of analysis reported: Intention to treat analysis	
Participants	Setting: in a single outpatient research centre, USA	
	Mean age (+/-SD or Range): 37.5 (14.6) years for acupuncture group, 38.9 (14.1) years for sham acupunc ture group, 39.0 (14.0) for wait list group	
	Men/Women (n/n): 54/176	
	Recruitment method: Participants were recruited from advertisements in media, fliers, and through re- ferrals from health professionals	
	Minimum duration of IBS symptoms: Rome II criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis	
	Actual duration of IBS symptoms before enrollment: IBS was present for > 1 year in 96% of participants in acupuncture group, 91% of participants in sham acupuncture group and 95% of participants in wait list group	
	Diagnosis of IBS required to be eligible? (if yes, describe how patients were verified to have IBS): Rome II criteria with a score of ≥ 150 on the symptom severity scale	
	Evaluation to rule out organic gastrointestinal disease: Reported Predominant symptoms or stool patterns of participants: Alternating symptoms were most common (53%); diarrhea (25%) was slightly more common than constipation (21%). Diarrhea was most com- mon, and constipation was least common, in the sham acupuncture group	
	Hospital inpatients? (Y/N; if Y list number inpatients): No Were people with a history of acupuncture treatment excluded?: Yes	
Interventions	ACUPUNCTURE N allocated to acupuncture: 78 Style of acupuncture: Chinese Points selection: Flexible formula for specific symptoms Points stimulated: Six main fixed acupoints and 11 optional points. The fixed points are very common ly used in IBS patients and included Conception Vessel 10, Stomach 25, Liver 3, Spleen 4, Pericardiu 6, and Stomach 37. The optional points were Stomach 36 and Conception Vessel 4 (for the Chinese of agnosis of deficiency), Large Intestine 4 Liver 14 (for stagnant qi or energy), Stomach 40, Large Intest 11 (for dampness), Stomach 27 (for cold), Conception Vessel 12 (for "retention of food"), Gall Bladdee (for "damp heat"), and Spleen 10 and Spleen 6 (for "blood stasis") Description of acupuncturist qualifications: "Four licensed acupuncturists, each with more than 2,00 hours of professional training and more than 4 years of post-graduate experience, performed the acupuncture in this study"	
	Total length of treatment period (weeks): 3 Number of sessions target (mean): 6 Times per week: 2 Number of points used (mean): 6 fixed acupoints plus optional points. If the acupuncturist felt strong- ly that a fixed main point would be less desirable than an optimal point, they were allowed to replace a single fixed point with an optional point Insertion depth: Not reported Was De Qi reportedly sought?: Yes Duration (mins): 20 min	



Lembo 2009 (Continued)

Lembo 2009 (Continued)	Method of stimulation: Manipulation
	CONTROL GROUP A INTERVENTION (sham, if used): Sham acupuncture with Streitberger needles N allocated to control group A: 75 Points selection: Predetermined "non-acupuncture" points in the relative vicinity of the genuine points. At least 5 and a maximum of 11 "non-acupuncture" points in the relative vicinity of the genuine points. At least 5 and a maximum of 11 "non-acupuncture" points in the relative vicinity of the genuine points. At least 5 and a maximum of 11 "non-acupuncture" points. Acupuncturists were also encour- aged to slightly switch or adjust their point selection within the parameters of the study protocol from treatment to treatment Total length of treatment period: 3 weeks Number of sessions target (mean): 6 Times per week: 2 Number of points used: At least 5 and a maximum of 11 "non-acupuncture" points Insertion depth: Non-insertion Was De Qi sought?: No Duration (mins): ~20 Method of stimulation: No
	CONTROL GROUP B: Waitlist N allocated to control group B: 77 Total length of treatment period: 3 weeks Number of sessions target (mean): NA Times per week: NA
	Duration (mins): NA
	Any co-interventions in all groups?: Participants were allowed to continue their IBS medications (e.g. fiber, anti-spasmodics, and loperamide) as long as they had been on stable doses for at least 30 days before entering the study and agreed not to change medications or dosages during the trial
Outcomes	Symptom severity:
	1. IBS Adequate Relief (IBS-AR): "IBS-AR is a dichotomous single item that asks participants, 'Over the past week have you had adequate relief of your IBS symptoms?' A responder was defined as a patient who answered this question affirmatively"
	2. IBS Symptom Severity Scale: "The IBS-SSS contains five questions that are rated on a 100-point visu- al analog scale, namely, the severity of abdominal pain, the frequency of abdominal pain, the severity of abdominal distention, dissatisfaction with bowel habits, and interference with the quality of life. All five components contribute to the score equally, yielding a theoretical range of 0 to 500, with a higher score indicating a worse condition. Earlier studies have established that scores < 175 represent mild IBS symptoms, 175 to 300 represent moderate severity, and scores > 300 represent severe IBS. A decrease of 50 points on the IBS-SSS has been shown to correlate with improvement in clinical symptoms. We therefore defined patients with a decrease of 50 points on the IBS-SSS as responders in this study." Measured as change in score from baseline for each group
	Quality of life: "The IBS-QOL is a 34-item measure assessing the degree to which IBS interferes with a patient's quality of life. Each item is rated on a 5-point Likert scale and all items are summed. The total score is then converted linearly to a 100- point scale, with higher scores indicating an improved quality of life. We defined responders as patients whose IBS-QOL scores improved by 10 points from baseline." Measured as change in score from baseline for each group"
	Time points of symptom measurement:
	The responder rates for the IBS-AR, IBS-SSS and IBS-QOL were reported at 3 weeks. The scores (mean and SD) of IBS-SSS and IBS-QOL were reported at baseline and 3 weeks
	Author's conclusion: This study did not find evidence to support the superiority of acupuncture com- pared with sham acupuncture in the treatment of IBS Additional outcomes reported in the trial but not extracted: The responder rates and score (mean and SD) of IBS Global Improvement Scale (IBS-GIS) were reported at 3 weeks. All study personnel, except the practitioners, were blinded to participant assignment



Lembo 2009 (Continued)	Adverse effects: "Three adverse events were reported during the acupuncture vs. sham acupuncture phase of the study: (i) painful foot cramp after treatment (sham acupuncture), (ii) nausea or hip pain (true acupuncture), and (iii) rib pain after a fall (sham acupuncture). All of these events were considered to be unrelated to the study procedure"
Notes	Comments: Carefully worked, especially for the blinding of patients and evaluators For the quality of life outcome, symptom severity outcome and global improvement outcome, we com- bined the two acupuncture groups (i.e. augmented and limited) and the two sham groups (i.e. aug- mented and limited) shown in Table 1 by calculating a weighted mean for combined acupuncture and combined sham. For combining the two SDs for each group, we used the formula in Table 7.7.a in the <i>Cochrane Handbook for Systematic Reviews of Interventions</i> for combining SDs of global improvement outcome. We used the SDs of the baseline measure as an estimate of the SDs of the endpoint measure for symptom severity outcome and quality of life outcome
	For responder rates, the denominator for the response rate are the number available for follow-up at that time-point, not the number randomized Funding resources: "This research was made possible by National Institutes of Health (NIH) Grant 1R01 AT001414-01 from the National Center for Complementary and Alternative Medicine (NCCAM) and the National Institutes of Digestive, Diabetes and Kidney Disease, Grant 1R21 AT002860-01 from NCCAM and the OQ ce of Behavioral and Social Science Research (OBSSR), and Grants 1 R21 AT002564 and 1K24 AT004095 from NCCAM. 2 is research was also supported in part by Grant RR 01032 to the Beth Israel Deaconess Medical Center (BIDMC) General Clinical Research Center from the NIH and the Samueli In- stitute for Informational Biology."

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"at the end of week 3, all patients who had received sham acupuncture were re-randomized blindly to either acupuncture or continuation of sham acupuncture. This was stratified by the group assignment for the run-in period (augmented vs. limited) and by the post run in pain score (< 30 vs. 30 on a 100- point visual analog scale)."
		"We randomly assigned participants to the three study arms using permuted block randomization with variable block sizes and assignments provided in sequentially numbered opaque sealed envelopes At three weeks, we used similar methods to randomise patients in the sham acupuncture groups to continue sham acupuncture or to switch to genuine acupuncture. This ran- domization was stratified by the level of abdominal pain at the three week visit (<30 v ≥30 on a 100 point visual analogue scale)"
Allocation concealment (selection bias)	Low risk	"We randomly assigned participants to the three study arms using permut- ed block randomization with variable block sizes and assignments provided in sequentially numbered opaque sealed envelopes. An administrative assis- tant, not otherwise involved in the study, opened the assignment envelopes and recorded the assignment of each participant in a confidential log. At three weeks, we used similar methods to randomise patients in the sham acupunc- ture groups to continue sham acupuncture or to switch to genuine acupunc- ture"
Blinding (performance bias and detection bias) All outcomes	Low risk	"In this study, we used Streitberger needles (22), a validated sham acupunc- ture device. This device has been shown to be indistinguishable from an ac- tual acupuncture device; the "needle" does not pierce the skin but creates an illusion of doing so as it retracts into a hollow handle. Streitberger and true acupuncture needles were applied for an identical period of time. To avoid acupuncture pressure effect, sham needles were placed over predetermined "non-acupuncture" points in the relative vicinity of the genuine points. Our team's precise method of using the sham needles is described elsewhere"

Acupuncture for treatment of irritable bowel syndrome (Review)



Lembo 2009 (Continued)

		"Although the trial was prospectively designed to investigate non-specific effects in irritable bowel syndrome, its design included a nested acupuncture substudy that allowed potential participants in the "treatment" arms to be told, truthfully, that they had a 50% chance of receiving genuine acupuncture during the trial. When the study ended, a letter was sent to all participants explaining the exact purpose of the study and offering them the opportunity to withdraw their original consent to use their data. All study personnel, except the practitioners, were blinded to participant assignment. Blinded registered nurses who were otherwise unconnected to the study conducted assessments" No attempt to confirm patient blinding, although this may not be necessary, as the Streitberger needle has been previously validated
Incomplete outcome data (attrition bias) Short term	Low risk	There were 8(78) drop-outs in acupuncture group and 5(75) in sham acupunc- ture group
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall IBS symptom severity (IBS-SSS) and health-related quality of life (IBS-QOL) at the completion of the 3 week treat- ment period, together with results for each outcome measured according to the methods section
Other bias	Low risk	"All analyses were intent-to-treat" "Table 2 displays demographics and baseline symptoms by treatment group." Although no statistical comparisons are made between the two groups at baseline, the groups seem comparable on most variables. Most importantly, the true acupuncture and sham acupuncture groups had very similar baseline scores for the IBS-SSS and the IBS-QOL

Li 2009	
Methods	Design: Parallel Blinding: No Attempt to confirm patient blinding?: No
	Dropouts/withdrawals: Not reported in the article [There were dropouts during the 70 days treatment, but the author did not record the number of the dropouts for the three groups]
	Total duration: 70 days treatment followed by an evaluation, no follow up reported Type of analysis reported: not reported
Participants	Setting: Outpatient ward of digestion medicine and TCM in Changchun Central hospital from Aug 2007 to Apr 2009, Changchun, China
	Mean age (+/- Standard Deviation or Range): Not reported
	Men/Women (number/number): Not reported
	In each group, the difference of patients´gender, age, disease duration, clinical classification, was not statistically significant (P > 0.05)
	Recruitment method: Not stated
	Minimum duration of IBS symptoms: Rome III criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis)
	Actual duration of IBS symptoms before enrollment: Not reported.

Li 2009 (Continued)	
	Diagnosis of IBS required for eligibility? Yes, Rome III criteria for irritable bowel syndrome and Consen- sus on the diagnosis and treatment of irritable bowel syndrome. Chinese Medical Association, Chinese Society of Gastroenterology (Guangzhou, 2003).
	Evaluation to rule out organic gastrointestinal disease: Not reported Predominant symptoms or stool patterns of participants: Not reported
	Hospital inpatients?: No
	Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	 Were people with a history of acupuncture treatment excluded?: Not reported ACUPUNCTURE (Warming needle moxibustion) plus spinal manipulation (spinal Tui na massage) Number allocated to acupuncture: 62 Style of acupuncture: Chinese (warming needle) Points selection: Flexible formula Points stimulated: 7 main fixed acupoints and 2 to 3 optional points. The fixed points were ST 25, BL 25 CV 12, ST 36, SP 6, SP 4 and LR 2. The optional points were BL 20, ST 37 and SP 9 for diarrhea; SJ 6 and ST 40 for constipation. Description of acupuncturist qualifications: Not reported Total length of treatment period (weeks): 10 (70 days) Number of sessions target (mean): 60 Times per week: ~ 6 Number of points used (mean): 7 main fixed acupoints and 2 to 3 optional points Insertion depth: Not reported Was De Qi reportedly sought? Yes, before moxibustion on needle Duration (minutes): 40 min Wethod of stimulation: After De Qi was sought, the moxa rolls were put on the needle end and were ignited to make the participants feel comfortably warm, numb, sore and swelling. It was inferred from the text that the moxibustion was used at all points Spinal manipulation: Total length of treatment period (weeks): 10 (70 days) Number of sessions target (mean): 60 Times per week: ~ 6 Duration (minutes):12 min CONTROL GROUP A INTERVENTION: Spinal manipulation alone Number of sessions target (mean): 60 Times per week: ~ 6 Duration (second group A: 62 Total length of treatment period (weeks): 10 (70 days) Number of sessions target (mean): 60 Times per week: ~ 6 Duration (minutes):12 min CONTROL GROUP A INTERVENTION: Spinal manipulation alone Number of sessions target (mean): 60 Times per week: ~ 6
	Duration (minutes): 12 min CONTROL GROUP B INTERVENTION: Acupuncture (Warming needle moxibustion) alone Number allocated to control group B: 62 Style of acupuncture: Chinese Points selection: Flexible formula
	Points selection: Flexible formula Points stimulated: 7 main fixed acupoints and 2 to 3 optional points. The fixed points were ST 25, BL 25, CV 12, ST 36, SP 6, SP 4 and LR 2. The optional points were BL 20, ST 37 and SP 9 for diarrhea; SJ 6 and ST 40 for constipation Description of acupuncturist qualifications: Not reported Total length of treatment period (weeks): 10 (70 days)
	Number of sessions target (mean): 60 Times per week: ~6
	Number of points used (mean): 7 main fixed acupoints and 2 to 3 optional points Insertion depth: Not reported Was De Qi reportedly sought? Yes



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i 2009 (Continued)			
	Duration (minutes): 40 Method of stimulation:		
		oustion: After De qi were sought, the moxa rolls were put on the needle end and ne participants feel comfortably warm, numb, sore and swelling	
	Any co-interventions in	all groups? Not reported	
Outcomes	Symptom severity:		
	1) cured: The IBS symp situation appeared in f 2) improved: The IBS sy constipation improved 3) no effect: The IBS sym	mptom improved, the frequency of defecation decreased, stool get normal or	
	Quality of life: Not assessed		
	Time points of symptom measurement:		
	The responder rates of IBS symptom score were reported after 10 weeks treatment		
	Author's conclusion: "Acupuncture and Tui Na are all effective treatment methods of IBS. The combina- tion of warming needle and spinal Tui Na method to treat IBS, showed significant synergy that can im- prove the clinical efficacy of IBS treatment"		
	Additional outcomes reported in the trial but not extracted: NA.		
	[The evaluator was the doctor who treated the participants himself]		
	Adverse effects: Not reported		
Notes	Funding sources: [No funding]		
	Randomization: [Referring to a random number table]		
	Allocation: [No specific method was used to conceal the allocation]		
	Reason for identical numbers assigned to the three groups: [The participants were originally assigned referring a random number table. When Li W found unequal sample size of the 3 groups, he further recruited participants to make the sample size equal finally. The participants further recruited were not randomly assigned]		
	Information in square brackets was provided by the first author Li W, in a telephone survey on July 29, 2011		
	We dichotomized cured, improved and no effect into the 2 categories of improved or not improved ac- cording to the methods section of the review		
Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence genera- tion (selection bias)	High risk	"Sixty IBS cases of liver-qi stagnation with spleen deficiency were randomized into a treatment and control group by single-blind method"	
		[We used drawing of lots. If the number of patient of one group started to ex- ceed 30, they assigned the additional patients into the other group]	

Allocation concealment High risk [No specific method was used to conceal the allocation] (selection bias)

Acupuncture for treatment of irritable bowel syndrome (Review)

Li 2009 (Continued)			
Blinding (performance bias and detection bias) All outcomes	High risk	[Quality of life scales were evaluated by participants themselves. The IBS symptom score for TCM symptom were evaluated by the investigator who knew the group assignment of the participants	
Incomplete outcome data	Unclear risk	Dropouts were not reported or described in the article	
(attrition bias) Short term		[In order to ensure the number of 30 participants for both groups, the partici- pants who withdrew during the 4 weeks treatment were eliminated from the trial and new participants were recruited. The first author could not recall the number of the dropouts	
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for IBS symptom score for TCM symptom at the completion of the 4 week treatment period and quality of life scale before and after treatment, together with results for each outcome measured according to the Observation targets in Chen YQ 2006 Master's thesis	
Other bias	Unclear risk	ITT analysis was not reported and the author also could not recall the number of dropouts, so we could not estimate whether the "missingness" would affect the treatment effect in this trial	
		"There was no significant difference in gender, age and duration between the two groups." . The P values were 0.417 for gender, 1.0 for age and 0.919 for du- ration between the two groups. There are no P values reported for the com- parison of quality of life scale and each symptom score between two groups at baseline, but based on the mean(SD) of quality of life score and symptom score for each item, the two groups appeared comparable at baseline	

Liu 1997

Liu 1997	
Methods	Design: Parallel group Blinding: No Attempt to confirm patient blinding?: No Dropouts/withdrawals: At the end of Tx, explicitly no dropouts for all groups, but the number of base- line participants is unreliable; At 26 week follow-up, 27 drop-outs from intervention group (acupunc- ture and psychological Tx), 25 dropouts from acupuncture group and 25 dropouts from psychological group Total duration: 20 to 120 days for acupuncture treatment, 1 to 12 weeks for psychotherapy treatment, followed by an evaluation; 26 weeks follow-up evaluation after end of treatment, but only for the pa- tients who responded to treatment, and therefore not a randomized comparison Type of analysis reported: NA
Participants	Setting: [The trial was done in China] Mean age (+/-SD or Range): 16 to 64 years for entire population, 19 to 64 years for acupuncture plus psychotherapy group, 16 to 62 years for acupuncture group, 22 to 58 years for psychotherapy group Men/Women (n/n): 89/61 Recruitment method: Not reported Minimum duration of IBS symptoms: Not reported
	Actual duration of IBS symptoms before enrollment: 3 months to 32 years Diagnosis of IBS required to be eligible? (if yes, describe how patients were verified to have IBS): Only stated that "all patients had IBS" Predominant symptoms or stool patterns of participants: Not reported
	Evaluation to rule out organic gastrointestinal disease: Reported that organic gastrointestinal diseases were ruled out but not stated how they were ruled out Hospital inpatients? (Y/N; if Y list number inpatients): Not reported Were people with a history of acupuncture treatment excluded?: Not reported

Acupuncture for treatment of irritable bowel syndrome (Review)



Liu 1997 (Continued)	
Interventions	ACUPUNCTURE + PSYCHOTHERAPY N allocated to acupuncture+ psychotherapy: 50 Style of acupuncture: Chinese Points selection: Flexible formula for specific symptoms Points stimulated: ST-36, PC-6, CV-12, ST-25, LR-3, BL-20, BL-23, GV-4 / CV-8 (Moxibustion) Description of acupuncturist qualifications: No description
	Total length of treatment period (weeks): 3 to 21 weeks Number of sessions target (mean): 10 to 60 for acupuncture treatment Times per week: 3 Number of points used (mean): 3 to 4 Insertion depth: Not reported Was De Qi reportedly sought?: Yes Duration (mins): 30 Method of stimulation: Manipulation
	Psychotherapy procedure: 1 to 2 sessions per week, 2 sessions comprise 1 course (Each session was performed ahead of acupuncture)
	CONTROL GROUP A INTERVENTION (sham, if used): Acupuncture only N allocated to control group A: 50 Points selection: Flexible formula for specific symptoms Points stimulated: ST-36, PC-6, CV-12, ST-25, LR-3, BL-20, BL-23, GV-4 / CV-8 (Moxibustion) Total length of treatment period: 3 to 21 weeks Number of sessions target (mean): 10 to 60 Times per week: 3 Number of points used (mean): 3 to 4 Insertion depth: Not stated Was De Qi sought?: Yes Duration (mins): 30 Method of stimulation: Manipulation
	CONTROL GROUP B INTERVENTION (sham, if used): Psychotherapy only N allocated to control group B: 50 Total length of treatment period: Not stated Number of sessions target(mean): Not stated Times per week: 1 to 2 Duration (mins): Not reported
	Any co-interventions in all groups?: Not reported
Outcomes	Symptom severity: Symptom Improvement (3-point scoring system) at end of treatment: 1) cured: 2) improved 3) no effect Follow up (6 months after the treatment): 1) symptoms remain improved 2) symptoms reoccurred
	Measured as % responders for each groups
	Quality of life: Not assessed
	Time points of symptom measurement: the end of treatment, 26 weeks after treatment Author's conclusion: Acupuncture plus psychotherapy is better than either alone in the treatment of IBS Addtional outcomes reported in the trial but not extracted: None Adverse effects: Not stated
Notes	Additional information in square brackets was provided by the first author, Liu GZ in a telephone survey
	Comments: Low methodological quality, small sample size

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Liu 1997 (Continued)

Funding resources: [There was no funding supporting the trial]

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	[Random number table]
Allocation concealment (selection bias)	Unclear risk	[The random numbers were concealed in sealed envelopes. The author did not specify whether or not the envelopes were numbered and opaque]
Blinding (performance bias and detection bias) All outcomes	High risk	[The outcome assessor was blinded to the intervention]
Incomplete outcome data (attrition bias) Short term	High risk	Patient dropouts and withdrawals from treatment were not reported
Selective reporting (reporting bias)	Low risk	The only outcome collected was the improvement at the end of treatment and follow up: At the end of treatment: Improvement (3-point scoring system. 1) cured: 2) im- proved 3) no effect Follow up (6 months after the treatment): 1) symptoms remain improved 2) symptoms reoccurred The number and response rate for each scoring item were reported at end of treatment and at 6 months follow up (only for responded patients)
		Outcomes were reported for overall IBS symptom severity (response to treat- ment in three categories) at the completion of the treatment periods. This cat- egorization is a standard way of assessing treatment success in Chinese trials of IBS.
Other bias	High risk	The author did not report whether groups were similar at baseline regarding the most important prognostic indicators. Timing of the outcome assessment in all groups appears to vary, because the number of treatment sessions for patients is dependent on their disease condition. Intention-to-treat analysis was not reported.

Liu 2010

Liu 2010	
Methods	Design: Parallel Blinding: Yes
	Attempt to confirm patient blinding?: No Dropouts/withdrawals: No [There were no dropouts]
	Total duration: 4 weeks treatment followed by an evaluation, no follow up reported Type of analysis reported: Not reported
Participants	Setting: Outpatient of digestion medicine and inpatient of the department of internal medicine in Hebei Provincial Hospital of TCM, China
	Mean age (+/- Standard Deviation or Range): 45.75 (7.91) years for acupuncture plus Chinese medicine group, 46.16 (8.13) years for bifidobacteria group, 45.71 (7.89) years for Chinese medicine group, 46.08 (8.07) years for acupuncture group

Acupuncture for treatment of irritable bowel syndrome (Review)



Liu 2010 (Continued)	
	Men/Women (number/number): 118/182
	Acupuncture/Geshanxiaoyao Decoction: 61/89, Bifidobacterium: 19/31, Geshanxiaoyao Decoction: 17/33, Acupuncture: 21/29
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Rome III criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis
	Actual duration of IBS symptoms before enrollment: 24.56 (3.10) months for acupuncture plus Chinese medicine group, 23.79 (2.92) months for bifidobacteria group, 24.37 (2.84) months for Chinese medicine group, 25.13 (2.89) months for acupuncture group
	Diagnosis of IBS required for eligibility?: Rome III criteria for diarrhea-predominant irritable bowel syn- drome and criteria of diarrhea caused by stagnation of the liver-Qi attacking the spleen and spleen stomach deficiency (<i>Guide for clinical trials of new drugs of Chinese medicine</i>)
	Evaluation to rule out organic gastrointestinal disease: Not reported Predominant symptoms or stool patterns of participants: The study only recruited patient with diar- rhea-predominant IBS
	Hospital inpatients? Yes, but the number of inpatients was not reported. Were people with a history of acupuncture treatment excluded? Not stated
Interventions	ELECTRO-ACUPUNCTURE PLUS CHINESE MEDICINE Number allocated to acupuncture: 150 Style of acupuncture: Chinese Points selection: Fixed formula Points stimulated: 3 fixed bilateral acupoints including LV 3, SP 6 and HT 3 Description of acupuncturist qualifications: Not stated
	Total length of treatment period (weeks): 4 Number of sessions target (mean): 28 All the authors of Chinese language trials did not specify whether they treated patients in the weekend. We assume patients were treated on weekends, because the authors described that the treatment was 7 times a week
	Times per week: 7 Number of points used (mean): 3 (bilateral 6) Insertion depth: Not reported Was De Qi reportedly sought?: [Yes] Duration (minutes): [15 min] Method of stimulation: [Manipulation and Electro-acupuncture]
	Chinese medicine: Formula name: Geshan Xiaoyao Fang Ingredients and preparation: auriculate swallowwort root 15 g, Chinese thorowax 12 g, root of herba- ceous peony 20 g, Chinese angelica root 12 g, the rhizome of large-headed atractylodes 10 g, Tuckahoe 15 g, cortex albiziae 20 g, Flos Albiziae 12 g, caulis polygoni multiflor 20 g. The above herbs were de- cocted into 300 ml and divided into 2 bags with 150 ml/bag. Total length of treatment period: 4 weeks Number of sessions target (mean): 28 days per week: 7 Times per day: 2 (medicine were taken at 9 am and 9 pm) Dosage: 150 ml / time
	CONTROL GROUP A INTERVENTION: Bifidobacteria (Product name: Li Zhu Chang Le). This product is a Bifidobacterium longum composition. Number allocated to control group A: 50 Total length of treatment period: 4weeks Number of sessions target (mean): 28 days per week: 7 Times per day: 2



Liu 2010 (Continued)	Dosage: 1 pill / time
	CONTROL GROUP B: Chinese medicine alone
	Number allocated to control group B: 50 Formula name: Geshan Xiaoyao Fang (as described above)
	Total length of treatment period: 4 weeks Number of sessions target (mean): 28 days per week: 7 Times per day: 2 (medicine were taken at 9 am and 9 pm) Dosage: 150ml / time
	CONTROL GROUP C: Electro-Acupuncture alone Number allocated to control group C: 50 Style of acupuncture: Chinese Points selection: Fixed formula Points stimulated: 3 fixed bilateral acupoints including LV 3, SP 6 and HT 3 Description of acupuncturist qualifications: Not reportedTotal length of treatment period (weeks): 4 Number of sessions target (mean): 28 Times per week: 7 Number of points used (mean): 3 (bilateral 6) Insertion depth: Not reported Was De Qi reportedly sought? [Yes] Duration (minutes): [15 min]
	Method of stimulation: [Manipulation and Electro-acupuncture]
	Any co-interventions in all groups?: Not reported
Outcomes	 Symptom severity: Curative effect for Clinical IBS symptom (4-point scoring system) referring to <i>the Guide for clinical trials of new drugs of Chinese medicine</i> at end of treatment: 1) cured 2) improved 3) effective 4) no effect. 1) cured: the symptom and signs almost disappeared. The symptom scale decreased by > = 95% 2) markedly effective: the symptom and signs obviously improved. The symptom scale decreased by > = 70% 3) effective: the symptom and signs improved. The symptom scale decreased by > = 30%
	 4) no effect: the symptom and signs did not improved or aggravated. The symptom scale decreased by = < 30% Computation formula: (symptom score before treatment - symptom score after treatment) / symptom score before treatment * 100%
	Quality of life:
	"Index of curative effect for quality of life: The IBS-QOL questionnaire was used. The questionnaire items includes emotional state, mental status, sleep state, whether get up early, energy situation, phys ical condition, appetite and diet, influence to the social role and work. The score for every item ranged from highest 100 to lowest 0. Trial only reported IBS-QOL scores (mean and SD) for each item, not the total score before and after 4 weeks treatment, so quality of life was not extracted for this trial
	Time points of symptom measurement:
	The curative effect for Clinical IBS symptom were reported after 4 weeks treatment. The IBS-QOL scores (mean and SD) for each item were reported before and after 4 weeks treatment.
	Author's conclusion: "Geshan Xiaoyao Fang combined with acupuncture could improve the IBS-QOL of IBS patients." (Extracted from Liu QQ 2010 trial). "Geshan Xiaoyao Fang combined with acupuncture had good curative effect for diarrhea-predominant IBS" (Extracted from Wang ZK 2010a trial). "Elec- tro-acupuncture treatment of diarrhea-predominant irritable bowel syndrome is safe and effective, it is worth for further research and promotion."
	Additional outcomes reported in the trial but not extracted: The Clinical IBS symptom scores for each item such as diarrhea, abdomen ache/distention, borborygmus, lack of appetite, tire and asthenia, fa- tigue and not wanting talk were reported before and after 4 weeks treatment



Liu 2010 (Continued)	Adverse effects: "The results of routine blood, urine and stool test did not significantly change before and after treatment, which suggested there's no obvious toxic side effect in treatment group and con- trol groups."
Notes	Funding sources: Hebei provincial sci-tech support plan (No. 072761825)
	Reason for identical numbers assigned to the two groups: [We did not use any specific method to en- sure the same number of participants assigned to the two groups. It might be a coincidence]
	Information in square brackets was provided by one of the first authors, Liu QQ, in a telephone survey onJuly 29, 2011 and by the other first author Wang ZK, in a telephone survey on September 26, 2011
	We dichotomized cured, improved, effective and no effect into the 2 categories of improved or not im- proved according to the methods section of the review.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	[Random number table method was used. We assigned 300 participants into a treatment group (150 participants) and 3 control groups, 50 participants in each]
Allocation concealment (selection bias)	High risk	[The allocation procedure was open. We did not use any method to conceal the allocation]
Blinding (performance bias and detection bias) All outcomes	High risk	[The evaluators were not blind to the participant assignment]
Incomplete outcome data (attrition bias) Short term	Unclear risk	[There were no dropouts in the study]. We scored this as unclear because it is unlikely that there would be zero drop-outs in a 4 week trial of 300 participants
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for clinical IBS symptoms at the completion of the 4 week treatment period, together with results for each outcome mea- sured based on the "Observational targets" section in both Liu 2010 and Wang 2010a articles
Other bias	Unclear risk	Although ITT analysis was not reported, there were reportedly no drop-outs, although this does not seem plausible, as described above
		"The two groups were comparable at baseline in terms of sex, age, disease du- ration (P > 0.05)"
		There are no P values reported for the comparison of symptom score or IBS- QOL score among 4 groups at baseline, but the groups appeared comparable at baseline

.owe 2000	
Methods	Design: Parallel
	Blinding: Yes
	Attempt to confirm patient blinding? Not reported
	Dropouts/withdrawals: Not reported
	Total duration: 3 months (from baseline)
	Type of analysis reported: Not reported



.owe 2000 (Continued)	
Participants	Setting: Canada. Not stated. Mean age (+/-Standard Deviation or Range): Range of 18 to 73 years for entire population; age not pro- vided for each arm, text states "two groups were well-matched for age" Men/Women (number/number): 80% female in entire population; sex not provided for each arm
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Not reported
	Actual duration of IBS symptoms before enrollment: Not reported
	Diagnosis of IBS required for eligibility? Not reported Evaluation to rule out organic gastrointestinal disease: Not reported Predominant symptoms or stool patterns of participants: Not reported
	Hospital inpatients? Not reportedWere people with a history of acupuncture treatment excluded? Not reported
Interventions	ACUPUNCTURE Number allocated to acupuncture: 28 Style of acupuncture: Chinese Points selection: Formula Points stimulated: Names of points were not reported
	Total length of treatment period (weeks): 4 Number of sessions target (mean): 8 Times per week: 2 Number of points used (mean): 6 (usually needled bilaterally) Insertion depth: Not stated Was De Qi reportedly sought? Not reported Duration (minutes): 20 min
	Method of stimulation: Not reported CONTROL GROUP A INTERVENTION: Sham acupuncture with blunt needles
	Number allocated to control group A: 22
	Points selection: Formula
	Points stimulated: Names of points were not stated Total length of treatment period: 4 weeks Number of sessions target (mean): 8 Times per week: 2
	Number of points used: 9 Insertion depth: 0 Was De Qi sought? No Duration (minutes): 20 Method of stimulation: None
	Any co-interventions in all groups? No
	Total follow-up period: 8 weeks
Outcomes	Symptom severity: Individual patient-determined treatment success rate Time points of symptom measurement: At the end of treatment: Acupuncture group: 57%, Sham group: 45% (P = 0.57)
	Quality of life: IBS-36 Time points of symptom measurement: (P value for pre-post treatment change); Acupuncture group; P < 0.0001, Sham; P = 0.0005 Type of outcome data reported: Difference post treatment value between acupuncture and control groups. Change from baseline reported in publication

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Lowe 2000 (Continued)	Author's conclusion:	
	Acupuncture has no method-specific therapeutic benefit for the treatment of IBS	
	Additional outcomes reported in the trial but not extracted: Barostat rectal thresholds, McGill pain score.	
	Adverse effects: Not reported	
Notes	Weak points: small sample size, lack of described information	
	Improvement in pain scores and quality of life in both the treatment and control groups suggests that aspects of sham and true acupuncture may positively influence these measures - or that these are the effects of enrollment in the clinical trial	
	In the patients who did not attain their individual "success goals" there is no data presentation to indi- cate whether they experienced worsening of symptoms or adverse effects during the trial	
	Funding sources: Not reported	
	Additional information provided by William Depew on September 20, 2011	

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "This randomized double-blind study compared acupuncture to a vali- dated placebo acupuncture technique in 50 patients with ROME positive IBS"
		Comment: No specific description of sequence generation in the abstract text
		[The study employed computer generated random number sequencing to de- termine assignment. The sequence and codes were held by a third party unas- sociated with the clinical conduct of the trial. The code was revealed only after the final consecutive case was finished the entire trial observation period]
Allocation concealment (selection bias)	Low risk	Quote: "This randomized double-blind study compared acupuncture to a vali- dated placebo acupuncture technique in 50 patients with ROME positive IBS"
		Comment: No specific description of allocation concealment in the abstract text.
		[The study employed computer generated random number sequencing to de- termine assignment. The sequence and codes were held by a third party unas- sociated with the clinical conduct of the trial. The code was revealed only after the final consecutive case was finished the entire trial observation period]
Blinding (performance bias and detection bias) All outcomes	Unclear risk	Quote: "This randomized double-blind study compared acupuncture to a vali- dated placebo acupuncture technique (tapping a blunt needle on the skin and taping in place at the same points)"
		Comment: There's no description about whether the patients in this trial were required to have <i>never</i> used acupuncture (i.e. to be acupuncture naive) or not. There were no tests for checking the success of the blinding either. In addition, there was no reference for a study of the validation of this sham procedure, and we are unfamiliar with any study that has validated this procedure
Incomplete outcome data (attrition bias) Short term	Unclear risk	Comment: Dropouts/withdrawals were not reported or described
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for individual patient-determined treatment suc- cess and IBS-36 (only P value) at the completion of the 4 week treatment pe-

Acupuncture for treatment of irritable bowel syndrome (Review)



Lowe 2000 (Continued)

		riod, together with results for each outcome measured, according to the Ab- stract
Other bias	Unclear risk	Quote: "The two groups were well matched for age, symptom scores, Beck de- pression and State-Trait anxiety scores." However, there were no statistical tests or P values reported for these comparisons, probably because of the ab- stract format.
		Comment: ITT analysis was not reported and the drop-outs were not reported either, so we couldn't know the degree of "missingness" and whether it would be to affect the estimate of the treatment effect in this trial.
		Comment: It's an abstract and the reporting about methods and outcomes was unclear and incomplete.

Methods	Design: Parallel Blinding: No Attempt to confirm patient blinding?: No Dropouts/withdrawals: 1/10 was lost to follow up from the acupuncture group; 3/20 were lost to follow up in the usual GP care group Total duration: 3 months treatment followed by an evaluation, no follow-up reported Type of analysis reported: Intention to treat analysis
Participants	Setting: "The acupuncture was provided at independent clinics" in UK Mean age (+/-SD or Range): 37.5 (13.5) years in GP care only group, 34 (7.1) years in acupuncture plus GP group Men/Women (n/n): 7/23
	Recruitment method: "Patients were identified via the databases of four GP practices in Birmingham." "Patients were invited to participate by postal questionnaire which included screening questions"
	Minimum duration of IBS symptoms: Rome II criteria (disease criteria to be fulfilled for the last 3 months with symptom onset at least 6 months prior to diagnosis)
	Actual duration of IBS symptoms before enrollment: Mean duration 79.8 (81.8 SD) months in GP care only group, 44 (25.6 SD) months in acupuncture plus GP group
	Diagnosis of IBS required to be eligible? (if yes, describe how patients were verified to have IBS): Yes, Rome II criteria were used. Evaluation to rule out organic gastrointestinal disease: Reported Predominant symptoms or stool patterns of participants: Not reported
	Hospital inpatients? (Y/N; if Y list number inpatients): No Were people with a history of acupuncture treatment excluded?: Unclear, exclude those with current acupuncture treatment
Interventions	ACUPUNCTURE plus GP care N allocated to acupuncture: 10 Style of acupuncture: Chinese Points selection: Flexible formula for specific symptoms Points stimulated: Across styles, there was considerable overlap in the treatments given, the most commonly used point overall was ST36, which was needled in 48% of treatments. Other commonly used points were PC6 (used in 20% of treatments), LI4 (19%), SP6, SP8, KI3 and CV12 (17%), and LR3 and LI 11 (16%) Description of acupuncturist qualifications: "Acupuncture was provided by five generalist acupunctur- ists who were registered with the British Acupuncture Council, and had at least five years' experience."

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Reynolds 2008 (Continued)	Total length of treatment period (weeks): 12 Number of sessions target (mean): The target number of sessions was 10 over 12 weeks (mean = 8 ses- sions) Times per week: less than 1 Number of points used (mean): 6 (usually needled bilaterally) Insertion depth: 2 to 5mm Was De Qi reportedly sought?: Yes Duration (mins): 20 min Method of stimulation: Manipulation CONTROL GROUP A INTERVENTION (sham, if used): GP care only N allocated to control group A: 20 Total length of treatment period: 12 weeks Number of sessions target (mean): Not reported Times per week: Not reported Number of points used: Not applicable Insertion depth: Not applicable Was De Qi sought?: Not applicable Duration (mins): Not reported Method of stimulation: Not applicable Any co-interventions in all groups? : "Both groups could seek care elsewhere according to need"
Outcomes	Symptom severity: "Our primary outcome measure was the IBS Symptom Severity Score (SSS) at three months. Scored from 0 to 500 (< 75 = no IBS, 75 to 175 = mild case, 175 to 300 = moderate and 300+ = severe). A minimum clinically significant change is considered to be 50 points." Measured as score for baseline and end of treatment for each group
	Quality of life: "Our secondary outcomes at three months were \dots EQ-5D." (range 0 to 1)
	Time points of symptom measurement:
	IBS Symptom Severity Score (SSS) of two groups were reported at baseline and three months (Figure 2 and Main outcomes section). The difference (with 95% CI) on IBS-SSS between groups using adjusted method and multiple imputation method (ITT) were reported at three months (Main outcomes section)
	EQ-5D was reported only at baseline
	Author's conclusion: "The results of our pilot analysis suggest that more definitive research into acupuncture for IBS is merited." Additional outcomes reported in the trial but not extracted: An adjusted between group difference with 95% confidence interval of IBS Global Impact Score was reported at three month. Hospital Anxiety and Depression scale was reported only at baseline.
	Adverse effects: "No serious adverse events were reported. The acupuncturists reported some mild adverse effects of the acupuncture treatment (mild dizziness during treatment and temporary worsening of symptoms followed by improvement). Some patients also reported mild adverse effects, but they all continued to attend for treatment"
Notes	Comments:
	Weak points: Small sample size
	In first paragraph of Outcomes measures section, the author stated that "A minimum clinically signifi- cant change is considered to be 50 points" for IBS Symptom Severity Score. In Figure 2, we measured the IBS-SSS score at baseline and at 3 months with a ruler for each patient to get response rate for both groups. For SDs of IBS-SSS post-treatment score, we used the SDs of the baseline measure as an esti- mate of the SDs of the endpoint
	Reynolds et al reported IBS-SSS means at endpoint for participants in each group, omitting from the analysis the four study participants lost to follow-up. The investigators also reported the mean IBS-SSS difference between groups at endpoint using multiple imputation to estimate the missing values of participants lost to follow-up. We used this mean difference data to conduct a generic inverse variance

Reynolds 2008 (Continued)

meta-analysis, with results very similar to those obtained from the meta-analysis of mean endpoint values. The meta-analytic estimates were -93.56 [-166.37, -20.74] for the analysis using the multiple imputation data and -106.51 [-200.72, -12.31] for the analysis using the available case data (negative values indicate improvement in symptoms in the acupuncture treatment group, compared to the group not receiving treatment)

Funding resources: The study was funded by the Innovation and Research Priming Fund, University of York. Julie Reynolds was supported by a Medical Research Council studentship. Hugh MacPherson is supported by a Department of Health post-doctoral fellowship in complementary and alternative medicine

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"The randomization sequence was computer generated by an independent re- searcher and concealed allocation was achieved by remote central telephone contact"
Allocation concealment (selection bias)	Low risk	"The randomization sequence was computer generated by an independent re- searcher and concealed allocation was achieved by remote central telephone contact"
Blinding (performance bias and detection bias) All outcomes	High risk	This study was not blinded
Incomplete outcome data (attrition bias) Short term	Low risk	"One patient (1/10) was lost to follow up from the acupuncture group: a female with no treatment preference, a moderate severity score at baseline, who had only received two treatments. Three (3/20) were lost to follow up in the usu- al GP care group, they were all male, had more severe baseline IBS severity scores and a preference for acupuncture." Given the small percentage lost to follow-up, and the use of multiple methods to assess sensitivity to missing out- comes data (last observation carried forward, assigning the overall mean to missing values, and multiple imputation based on the standard best subset re- gression method) we judged that incomplete outcome data were appropriate- ly addressed
Selective reporting (re- porting bias)	High risk	This trial reported outcomes for overall IBS symptom severity (IBS-SSS) at the completion of the treatment period. However two outcome measures (i.e. Hospital Anxiety and Depression scale and EQ-5D) reported at baseline were not reported at the end of treatment, which raises the possibility that there could have been outcome reporting bias
Other bias	Low risk	"Analysis was by intention-to-treat, and multiple imputation was used for missing data." Baseline patient characteristics in the two groups were comparable, but no P values were reported. "There were some differences at baseline in Non- Colonic Symptom Score, working status, anxiety and depression, differences that can be expected in a sample of this size." The two groups were compara- ble on mean baseline IBS symptom severity scores, the outcome extracted for our meta-analysis

Schneider 2006

Methods	Design: Parallel Blinding: Yes

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chneider 2006 (Continued)	Attempt to confirm patient blinding?: Not reported Dropouts/withdrawals: During 5 weeks treatment, 1/22 drop-outs in acupuncture group and 1/21 in sham acupuncture group Total duration: 5 weeks treatment followed by an evaluation, additional follow-up evaluation at 3
	months after end of treatment Type of analysis reported: NA
Participants	Setting: at the outpatient clinic of the Department of Gastroenterology, University Medical Hospital, Germany
	Mean age (+/-SD or Range): 47.63 (14.71) years in acupuncture group, 47.14 (16.01) years for sham acupuncture group Men/Women (n/n): 9/34 Recruitment method: "Patients were recruited at the outpatient clinic of the Department of Gastroen- terology, University Medical Hospital, via gastroenterology specialists in private practice in the Heidel berg area, and via advertisements in local newspapers" Minimum duration of IBS symptoms: 12 weeks out of the last 12 months
	Duration of IBS symptoms before enrollment: < 2 years in 3 patients, 2 to 10 years in 14 patients, > 10 years in 25 patients Diagnosis of IBS required to be eligible? (if yes, describe how patients were verified to have IBS): Rome II criteria Evaluation to rule out organic gastrointestinal disease: Reported Predominant symptoms or stool patterns of participants: Alternating symptoms (17/43) and diarrhea (17/43) were more common than constipation (5/43) or normal stool habits (4/43)
	Hospital inpatients? (Y/N; if Y list number inpatients): No Were people with a history of acupuncture treatment excluded?: Assumedly no, only patients who re- ceived AC treatment within the last three months were excluded
Interventions	ACUPUNCTURE N allocated to acupuncture: 22 Style of acupuncture: Chinese Points selection: Fixed formula Points stimulated: Liver 3, Stomach 36, Spleen 6, Conception 12, Stomach 21, Stomach 25, Heart 7, Du Mai 20. Description of acupuncturist qualifications: "AC was performed by an experienced female acupunctur ist (SB) and by a trained female research assistant (CW)"
	Total length of treatment period (weeks): 5 Number of sessions target (mean): 10 Times per week: 2 Number of points used (mean): 8 points (14 needles for both sides) Insertion depth: Not reported Was De Qi reportedly sought?: Yes Duration (mins): Not reported Method of stimulation: Manipulation
	CONTROL GROUP A INTERVENTION (sham, if used): Sham acupuncture with a blunted telescopic placebo needle N allocated to control group A: 21 Points selection: fixed formula Points stimulated: 2 cm adjacent to the real AC point, the AC point LG 20 was not needled Total length of treatment period: 5 weeks Number of sessions target (mean): 10 Times per week: 2 Number of points used: 7 points (13 needles for both sides) Insertion depth: Non-insertion Was De Qi sought?: No Duration (mins): Not reported Method of stimulation: Not applicable

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Outcomes	Symptom severity: Not assessed
Outcomes	Symptom sevency: Not assessed
	Quality of life: "Patients completed two QOL questionnaires that were used to assess any effects of symptomatic improvement by AC on their overall QOL: (a) FDDQL assesses disease related impact of bowel symptoms on quality of life, measured with 43 items on eight subscales: daily activity disease related anxiety, diet, sleep, discomfort, health perception, coping with disease, and impact of stress. Scores for subscales are added to a global QOL scale ranging from 0 to 100. FDDQL has an internal consistency of 0.94 (Cronbach's alpha); (b) the health related quality of life questionnaire (SF-36) is a validated global measure of health related quality of life unrelated to specific diseases that has been wide ly used in a variety of diseases. It uses 36 items to assess eight scales (bodily function, bodily role, bodi ly pain, general health, vitality, social function, emotional role, and physical well being)." Measured as score for each time points for each group
	Time points of symptom measurement: "Both questionnaires (FDDQL and SF-36) were given prior to treatment (t1), immediately after 10 AC treatments (t2=five weeks after the first treatment), and three months after the last treatment (t3). Improvement in the global score of FDDQL immediately after the 10 treatments (t2) was the primary end point of the study. The other subscales of the FDDQL and the SF-36 were defined as secondary end points. Outcome measures at t3 were also defined as secondary end points"
	Author's conclusion: Acupuncture in IBS is primarily a placebo response
	Adverse effects: Not reported
Notes	Comments: Carefully worked, especially for the blinding of evaluator
	Weak points: small sample size
	Means and SEMs of quality of life could be measured with a ruler in Fig.1 for short term and long term. SDs were calculated following the formula: SD = SEM * sqrt (N)
	No information about response rate was reported
	The first author Antonius Schneider was contacted by email to request information about randomiza- tion. He replied with detailed information on January 27, 2011
	Funding sources: "The trial was financially supported by the German Medical Acupuncture Association (DAGFA). The funding source had no involvement in design, performance, or analysis of the study"

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"Block randomization was done by a central telephone centre" [Computer generated random list (randomization with Excel)]
Allocation concealment (selection bias)	Low risk	"Block randomization was done by a central telephone centre so that neither the patient nor the investigator (AS) knew whether the patient would receive AC or SAC. The randomization result was told to the acupuncturist (SB, CW) di- rectly after informed consent"
Blinding (performance bias and detection bias) All outcomes	Low risk	"In the control group, patients received SAC with a blunted telescopic place- bo needle (Asia Med) that simulates an AC procedure without penetrating the skin. The efficiency of this method has been shown in various studies. Each SAC procedure was performed 2 cm adjacent to the real AC point to avoid acu- pressure effects. In the SAC group, the AC point LG 20 was not needled as fixa- tion of the plaster is not possible on the head due to hair."



Schneider 2006 (Continued)

Schneider 2006 (conunued)		"Only one patient in the entire study population had received AC in the past. She was able to identify SAC and declined to participate in the study after the third treatment"
		No attempt to confirm patient blinding, although this may not be necessary in this case because the Streitberger needle has been previously validated. The Asia Med needle is the same as the Streitberger needle, which has previously been validated in other studies (Streitberger 1998; Kleinhenz 1999; Streitberg- er 2004) and the Streitberger needles were placed close to the genuine points. Therefore, the sham was likely to be indistinguishable from the true acupunc- ture
Incomplete outcome data (attrition bias) Short term	Low risk	"Only one patient in the entire study population had received AC in the past. She was able to identify SAC and declined to participate in the study after the third treatment. Another female patient declined to participate after five treatments due to time constraints. Thus two patients dropped out during the course of the study"
		During 5 weeks treatment, 1/22 drop-outs in acupuncture group and 1/21 drop-outs in sham acupuncture group
Selective reporting (re- porting bias)	Low risk	"Both questionnaires (FDDQL and SF-36) were given prior to treatment (t1), immediately after 10 AC treatments (t2 = five weeks after the first treatment), and three months after the last treatment (t3). Improvement in the global score of FDDQL immediately after the 10 treatments (t2) was the primary end point of the study. The other subscales of the FDDQL and the SF-36 were de- fined as secondary end points. Outcome measures at t3 were also defined as secondary end points"
		Global scale and subscales of FDDQL were reported at baseline, 5 weeks and 17 weeks Differences of FDDQL global and sub scales between two groups were reported at 5 weeks and 17 weeks Only statistical comparison of SF-36 scales between and within the two groups were reported in Results section.
		Results were reported for each outcome measured according to the Methods section, and a health-related quality of life outcome (the FDDQL) was reported at end of treatment and at follow up
Other bias	Low risk	Although ITT analysis was not reported, there were very few drop-outs, so the degree of "missingness" would be unlikely to affect the estimate of the treat- ment effect in this trial "There were no differences in age or sex between the groups at entry" The "Symptoms at study entry" section/paragraph includes the following text: "The BDQ illustrates the degree and severity of bowel symptoms among the patientsNo differences were seen between those who received AC or SAC af- ter assessment"

Shi 2010

Methods	Design: Parallel
	Blinding: No
	Attempt to confirm patient blinding?: No
	Dropouts/withdrawals: During the 4 weeks treatment, 1/32 drop-outs in acupuncture group and 1/38 in pinaverium bromide group
	Total duration: 4 weeks treatment immediately followed by an evaluation, 1 week follow up reported
	Type of analysis reported: Intention to treat analysis

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Shi 2010 (Continued)	
Participants	Setting: Outpatients of digestion medicine and integrative medicine in Hunan Provincial Hospital of TCM and Xiangya Hospital of Central South University, China.
	Mean age (+/- Standard Deviation or Range): 38.51 (14.65) years for acupuncture group, 38.68 (15.72) years for pinaverium bromide group
	Men/Women (number/number): 13/57
	Acupuncture: 7/25, Medication: 6/32
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Rome III criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis
	Actual duration of IBS symptoms before enrollment: 10.62 (9.23) months for acupuncture group, 10.80 (7.86) months for pinaverium bromide group
	Diagnosis of IBS required for eligibility?: Rome III criteria for diarrhea-predominant irritable bowel syn- drome
	Evaluation to rule out organic gastrointestinal disease: Reported - colonoscopy or barium enema ex- amination Predominant symptoms or stool patterns of participants: The study only recruited patient with diar- rhea-predominant IBS
	Hospital inpatients?: No Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	ELECTRO-ACUPUNCTURE Number allocated to acupuncture: 32 Style of acupuncture: Chinese Points selection: Flexible formula for specific symptoms Points stimulated: 7 main fixed acupoints and 3 optional points. The fixed points included ST 25, ST 36, BL 20, BL 21, BL 23, BL 25 and ST 37. The optional points were PC 6, LV 3 and EX-HN1 (for the diagnosis of psychological symptoms such as depression, anxiety and insomnia, etc) Description of acupuncturist qualifications: "All manipulations were practiced by professional acupuncturists with more than 5 years of clinical experience"
	Total length of treatment period (weeks): 4 Number of sessions target (mean): 28 (daily session, for 28 days) Times per week: 7
	Number of points used (mean): 7 fixed bilateral acupoints plus 3 optional points Insertion depth: 1 cun oblique insertion for ST 25 and BL 20; 1~1.5 cun for ST 36, ST 37, BL 23 and BL 25; 0.5 to 1 cun for PC 6 and LV 3; 0.5 to 0.8 cun horizontal insertion for EX-HN 1 Was De Qi reportedly sought?: Yes, before electro-stimulation Duration (minutes): 15 min
	Method of stimulation: Manipulation and Electro-acupuncture
	Electro-acupuncture procedure: After De Qi, G6805-2A electro-acupuncture device was connected, ST-25, PC-6, LR-3 with continuous-wave, ST-36, ST-37, BL-20, BL-23, BL-25, with the dense-sparse wave, EX-HN1 Sishencong with intermittent waves. According to the patients ´ sensitivity and tolerance to ad- just the stimulation intensity and frequency. There was a close observation during electro-acupuncture treatment, to prevent accident.
	CONTROL GROUP A INTERVENTION: Pinaverium bromide (Product name: Deshute) Number allocated to control group A: 38 Total length of treatment period: 4 weeks Number of sessions target (mean): 28 days per week: 7 Times per day: 3 Dosage: 50mg / time



Shi 2010 (Continued)	Any co-interventions in all groups? Not reported		
Outcomes	Symptom severity:		
	1. Overall IBS symptom VAS score: Overall IBS symptom VAS score was rated on a 10-point visual ana- log scale		
	2. SRI: SRI is an efficacy index based on overall IBS symptom VAS score. A responder was defined as a patient whose SRI was more than 60% SRI = (symptom score before treatment - symptom score after treatment) / symptom score before treatment		
	Quality of life: Not assessed		
	Time points of symptom measurement:		
	The Overall IBS symptom VAS scores (mean and SD) were reported at baseline, 1 week, 2 week, 3 week, 4 week and 5 weeks (i.e. 1 week after the treatment). [The VAS scores were presented as mean and SD] The responder rates of IBS symptom based on SRI for both groups were: "Based on patients SRI level, at the end of treatment, effective rate in treatment group is 81,3%, in the control group 52,6%" (end of Section 4.3)		
	Author's conclusion: "Electro-acupuncture was safe and effective for diarrhea-predominant IBS and is worth further researching and popularizing. This study found evidence to support the superiority of electro-acupuncture compared with pinaverium bromide"		
	Additional outcomes reported in the trial but not extracted: The diarrhea VAS score were reported at baseline, 1 week, 2 week, 3 week, 4 week and 5 week (Figure 1). The abdominal pain/discomfort VAS score were reported at baseline, 1 week, 2 week, 3 week, 4 week and 5 week (Figure 2). The evaluators who administered the VAS tests to the patients were blind to the participant assignment.		
	Adverse effects: "One participant in electro-acupuncture group withdrew due to acupuncture syncope and one participant in pinaverium bromide group withdrew due to itch and skin rash caused by allergic reaction. There's no obvious adverse effect in other participants"		
Notes	Funding sources: [No funding]		
	Information in square brackets was provided by the first author Shi XH and second author Luo JK, in telephone surveys on Jul 21y and 22, 2011		
	Since the numerical data were not in the text, we measured the Overall IBS symptom VAS scores (mean and SD) with a ruler at each time points for both groups using Figure 3		
Risk of bias			

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	[We referred to a random number table]
Allocation concealment (selection bias)	Low risk	[We used sequentially numbered, opaque, sealed envelopes]
Blinding (performance bias and detection bias) All outcomes	High risk	[The evaluators were blind to the participant assignment]
Incomplete outcome data (attrition bias) Short term	Low risk	"One participant in electro-acupuncture group withdrew due to acupuncture syncope and one participant in pinaverium bromide group withdrew due to itch and skin rash caused by allergic reaction"

Acupuncture for treatment of irritable bowel syndrome (Review)

Shi 2010 (Continued)		
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall IBS symptom VAS score and responder rate at the completion of the 4 week treatment period, together with results for each outcome measured according to the observational targets section
Other bias	Low risk	ITT analysis was reported. It was not clear how the data were imputed for the 2 drop-outs. However, this trial had a large, statistically significant effect size (i.e. RR 1.54, 95% CI 1.09 to 2.18) and any assumptions made about the treatment response of these 2 drop-outs would not impact the overall results of this trial or the results of the meta-analysis that included this trial The two treatment groups were comparable at baseline in terms of the demo- graphic characteristics, diarrhea VAS score, abdominal pain/discomfort VAS score and overall IBS symptom VAS scores

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Methods	Design: Parallel Blinding: No Attempt to confirm patient blinding?: No
	Dropouts/withdrawals: Three cases, 1 from the treatment group and 2 from the control group, were dropped out due to loss in the follow-up. A total of 60 patients completed the study"
Participants	Setting: "The 63 patients enrolled were outpatients of special IBS-D clinics of Chinese Medicine Hospi- tal of Jiangsu province from July 2009 to November 2010"
	Mean age (+/- Standard Deviation or Range): "The 31 patients in the treatment group were…aged 18 to 61 years (38.81 +/-11.80 years on average)…The 32 patients in the [drug] treatment group were…aged 18 to 59 years (38.59 +/- 11.45 years on average)"
	Men/Women (number/number): Acupuncture group: 13 males; 18 females; Drug group: 20 males; 12 fe males
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Rome III criteria for the last 3 months with symptom onset at leas 6 months prior to diagnosis
	Actual duration of IBS symptoms before enrollment: The duration for acupuncture group was 1 to 20 years (4.23 +/- 3.96 on average); the duration for western medication group was 1 to 38 years (5.63 +/-7.35 on average)
	Diagnosis of IBS required for eligibility?: Diagnosis of IBS-D based on the Rome III classification
	Evaluation to rule out organic gastrointestinal disease: Yes Predominant symptoms or stool patterns of participants: Diarrhea-predominant IBS
	Hospital inpatients?: No Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	ACUPUNCTURE (plus mild moxibustion) Number allocated to acupuncture: 31 Style of acupuncture: Chinese Points selection: Fixed formula Points stimulated: "Points selection: bilateral Tianshu (ST25), Zusanli (ST36), Shangjuxu (ST37), Sany- inxiao (SP6), Taichong (LR3), Baihui (DU20), and Yintang (EX-HN 3) points"
	Description of acupuncturist qualifications: Not reported
	Total length of treatment period (weeks): 4

Acupuncture for treatment of irritable bowel syndrome (Review)



Sun 2011 (Continued)	Number of sessions target (mean): 20 Times per week: 5 Number of points used (mean): 7 fixed points. Insertion depth: 12 to 25 mm Was De Qi reportedly sought? Yes Duration (minutes): 30 min Method of stimulation: "Twist lifting-inserting gentle reinforcing-purging manipulation was applied af- ter de qi (getting qi)" CONTROL GROUP A INTERVENTION: Western medication "Dicetel (Pinaverium bromide tablets, product of Solvay Pharmaceuticals, France, batch No. H20080424) was given orally 50 mg, 3 times a day for a to- tal of 4 weeks" Number allocated to control group A: 32 Total length of treatment period: 4 weeks Number of sessions target (mean): 28 days per week: 7 Times per day: 3 times/day Dosage: 50 mg Any co-interventions in all groups? Not reported
Outcomes	Symptom severity:
	 Overall IBS symptom score (4-point scoring system) referring to the <i>Guiding Principle of Clinical Research on New Drugs of Traditional Chinese Medicine</i> at end of treatment: 1) cured 2) improved 3) effective 4) no effect. 1) clinically remitted: the symptom disappeared. Effect index = ≥ 95%. 2) markedly effective: symptoms and signs markedly relieved. 70% ≤ Effect index < 95% 3) effective: the symptom and signs improved. 30% ≤ Effect index < 70% 4) no effect: the symptom did not importantly improve. Effect index = < 30% Computation formula: Effect index = (symptom score before treatment - symptom score after treatment) / symptom score before treatment * 100%
	Quality of life:
	Drossman (IBS-QOL), adapted with Han modification, which consisted of 34 items in 8 domains. "Every item was differentiated into 5 grades (none, mild, moderate, lean to severe, severe) which were recorded as 5, 4, 3, 2, and 1 scores, respectively. The records were made before and after the 4 week treatment."
	Time points of symptom measurement:
	The responders for overall IBS symptom were reported at pre-treatment, week 1, week 2, and week 4
	The quality of life was reported at baseline and at end of treatment (week 4)
	Author's conclusion: "acupuncture treatment could effectively alleviate the degree and frequency of symptoms' attack in IBS-D patients…with the efficacy better than that of pinaverium bromide, show-ing a preponderance in improving patients' QOL"
	Additional outcomes reported in the trial but not extracted: Scores on the individual domains of the quality of life measure, before and after treatment
	Adverse effects: "No any adverse reaction occurred"
Notes	Funding sources: "Supported by the funds of Chinese Medicine Hospital of Jiangsu Province, China (No. Y09039-1) and Chinese Medicine Leading Talent Project of Jiangsu Province, China (No. LJ 200905)"
	Reason for identical numbers assigned to the two groups: [A block randomization was used, where there are equal numbers in the blocks. The block sizes were 8, with four assigned to acupuncture and 4 assigned to drug treatment in each block. There were 63 total randomized, and 8 blocks of 8 were used (with 1 assignment left over)]



Sun 2011 (Continued)

Trusted evidence. Informed decisions. Better health.

Allocation concealment: Not reported in the article. [Sequentially numbered, opaque envelopes]. This method does not entirely meet the Cochrane Handbook standard for allocation concealment, which requires that the envelopes by "sealed"

Evaluation blinding: [IBS-QOL was evaluated by participants themselves. Participants' symptoms were asked by staff who did not know the group assignment]

Information in square brackets was provided by the author Wu XL, in a telephone survey, followed by an e-mail for clarification, sent from Wu to Ke Cheng on December 2, 2011

We dichotomized cured, improved, effective and no effect into the 2 categories of improved or not improved according to the methods section of the review.

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"The patients were assigned by a random number table to two groups, the acupuncture treatment group and the drug control group."
		[A block randomization was used, where there are equal numbers in the blocks. The block sizes were 8, with four assigned to acupuncture and 4 as- signed to drug treatment in each block. There were 63 total randomized, and 8 blocks of 8 were used (with 1 assignment left over)]
Allocation concealment (selection bias)	High risk	[We used envelope method. The envelopes were sequentially numbered, opaque, unsealed]. We scored this trial as High Risk because the Cochrane Handbook requires that all 3 criteria (i.e. sequentially numbered, opaque, sealed) must be met for the envelope method to score a Low Risk of bias
Blinding (performance bias and detection bias) All outcomes	High risk	[IBS-QOL was evaluated by participants themselves. Participants' symptoms were asked by staff who did not know the group assignment]
Incomplete outcome data (attrition bias) Short term	Low risk	"Three cases, 1 from the treatment group and 2 from the control group, were dropped out due to loss in the follow-up. A total of 60 patients completed the study"
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall IBS symptom responders and quality of life at the completion of the 4 week treatment period, which are standard outcomes and the outcomes described as being measured in the Methods sec- tion
Other bias	Unclear risk	ITT was not used
		"The scores of symptoms in the two groups were insignificantly different be- fore treatment (t = 1.824, P = 0.074)The IBS-QOL scores in the two groups were similar at pre-treatment (t = 1.327, P = 0.19)"

Xiong 2008a

Methods

Design: Parallel Blinding: No Attempt to confirm patient blinding?: No Dropouts/withdrawals: During the 4 weeks treatment, no drop-outs mentioned in any of the three groups (acupuncture + Chinese herbal medicine group, Chinese herbal medicine group and acupuncture)



Xiong 2008a (Continued)	Total duration: 4 weeks of treatment, evaluation after treatment, but the specific time point of evalua- tion was not stated Type of analysis reported: NA
Participants	Setting: Not very specific. "120 patients from outpatient or inpatient were randomly divided into 3 groups." China Mean age (+/-SD or Range): 39.15 (15) years for Chinese herbal medicine group, 38.3 (16) years for acupuncture group, 38.6 (15) years for acupuncture + Chinese herbal medicine group Men/Women (n/n): 48/72 Recruitment method: "120 patients from outpatient or inpatient were randomly divided into 3 groups" Minimum duration of IBS symptoms: 1 year
	Actual duration of IBS symptoms before enrollment: 53.7 (5.1) months for Chinese herbal medicine group, 52.6 (5.4) months for acupuncture group, 54.2 +/- 4.9 months for acupuncture + Chinese herbal medicine group Diagnosis of IBS required to be eligible? (if yes, describe how patients were verified to have IBS): Rome II criteria and diagnostic criteria of liver-qi stagnation and spleen deficiency in "The principle of clinical research for new drug of traditional Chinese herbal medicine (in Chinese)" were used Evaluation to rule out organic gastrointestinal disease: Reported. "Subjects with other organic gas- trointestinal disease were excluded" Predominant symptoms or stool patterns of participants: Diarrhea Hospital inpatients? (Y/N; if Y list number inpatients): Yes, the number of inpatients was not reported
Interventions	Were people with a history of acupuncture treatment excluded?: Not reported ACUPUNCTURE+ CHINESE HERBAL MEDICINE N allocated to acupuncture: 42 Charles of acupunctures Chinese
	Style of acupuncture: Chinese Points selection: Fixed formula Points stimulated: CV-12, bilateral ST-25, LI-4, ST-36, LR-3
	Ingredient of Chinese herbal medicine: Thorowax root 12g, white peony root 15g, bitter orange imma- ture 6g, turmeric root, tuckahoe 15g, tangerine peel 6g, divaricate saposhnikovia root 9g, bighead atractylodes rhizome 15g, aucklandia root 6g, natgrass galingale rhizome 6g, oriental waterplantain rhizome 15g, licorice root 3g
	Description of acupuncturist qualifications: No description.
	Total length of treatment period (weeks): 4 Number of sessions target (mean): 28 Times per week: 7 Number of points used (mean): 9 points Insertion depth: Not reported Was De Qi reportedly sought?: Yes Duration (mins): 30 min Method of stimulation: Manipulation.
	Moxibusiton procedure: Moxibustion with warming needle procedure were performed on bilateral ST-25 and ST-36
	CONTROL GROUP A INTERVENTION (sham, if used): Chinese herbal medicine only group N allocated to control group A: 40 Total length of treatment period: 4 weeks Number of sessions target (mean): 28 Times per week: 7 Ingredient of Chinese herbal medicine: same as described above
	CONTROL GROUP B INTERVENTION (sham, if used): Acupuncture only group N allocated to control group B: 38 Points selection: Fix Points stimulated: CV-12, bilateral ST-25, LI-4, ST-36, LR-3



Xiong 2008a (Continued)	
	Total length of treatment period: 4 weeks Number of sessions target (mean): 28 Times per week: 7 Number of points used: 9 points Insertion depth: Not stated
	Was De Qi sought?: Yes Duration (mins): 30 Method of stimulation: Manipulation
	Moxibusiton procedure: Moxibustion with warming needle procedure were performed on bilateral ST-25 and ST-36
	Any co-interventions in all groups?: Not reported
Outcomes	Quality of life: Not assessed
	Symptom severity: Symptom Improvement (4-point scoring system) at end of treatment: 1) cured: 2) improved 3) effective 4) no effect
	"Measurements: Abdominal pain, diarrhea, abdominal distension, stool frequency and properties were assessed According to "The principle of clinical research for new drug of traditional Chinese herbal medicine (in Chinese)", no symptom (0 score) means well-formed motions, soft stool, 1 to 2 times a day; mild symptom (1 score) means life and work are not influenced, mushy stool or stool with a little mucus, 2 to 3 times a day; moderate symptom (2 score) means the symptom influences life and work but can be tolerated, moderate mucous stool, 4 to 5 times a day; severe symptom (3 score): the symptom interfere with life and work and can not be tolerated, rice-water stool or stool with lots of mucous, above 6 times a day"
	"1) Cured: disappearance of main symptoms, well-formed stool, disappearance of mucus, improve- ment of score > = 95%;
	2) Improved: disappearance of most main symptoms, well-formed stool, significantly decrease of mu- cus, improvement of symptom score > = 70% and < 95%;
	3) Effective: improvement of main symptoms, watery stool, decrease of mucus, improvement of symptom score > = 30%;
	4) No effect: no improvement of symptoms, improvement of symptom score < 30%
	Improvement of symptom score = (total symptom score before treatment - total symptom score after treatment) / total symptom score before treatment
	Total symptom score = abdominal pain score + abdominal distension score + diarrhea score + stool fre- quency score + stool properties score"
	Time points of symptom measurement: Before and after the treatment
	Additional outcomes reported in the trial but not extracted: Table 2, 3, 4 and 5 showed the number of responders of improvement of diarrhea, abdominal pain, abdominal distension and stool frequency respectively for each group after treatment
	Author's conclusion: "The findings indicate that the combination of acupuncture and Chinese herbal medicine shows promise in the treatment of patients with diarrhea-predominant IBS"
	Adverse effects: Not reported
Notes	We extracted data for acupuncture + Chinese herbal medicine versus Chinese herbal medicine only
	Additional information in square brackets was provided by the first author Xiong, in an telephone conversation with Ke Cheng from September 18, 2009, and translated by Ke Cheng in an e-mail to Eric Manheimer and L. Susan Wieland on February 28, 2011



Xiong 2008a (Continued)

Funding resources: When the first author Xiong was telephoned, she stated there was no funding supporting the trial

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	"120 patients from outpatient or inpatient were randomly divided into 3 groups."
		[The generation of the sequence was referred to a random number table]
Allocation concealment (selection bias)	High risk	[No, we only used the random number table, we didn't use envelope method or central allocation]
Blinding (performance bias and detection bias) All outcomes	High risk	Open study. Treatment allocation was not blinded
Incomplete outcome data (attrition bias) Short term	High risk	"Both trials didn't explicitly report the 100% follow up in the paper or report to me by the author."
Selective reporting (re- porting bias)	Low risk	Outcomes were reported for overall IBS symptom severity (response to treat- ment in four categories) at the completion of the treatment period. This cate- gorization is a standard way of assessing treatment success in Chinese trials of IBS. Abdominal pain, diarrhea, abdominal distension, stool frequency and properties were assessed at baseline and the number of responders for each symptom were described after treatment
Other bias	High risk	Intention-to-treat analysis was not reported
		There was a good balancing of the three groups at baseline for the age and du- ration of IBS, but there was no information about other baseline variables re- lated to symptom severity

Xue 2009

Design: Parallel Blinding: No
Attempt to confirm patient blinding?: No
Dropouts/withdrawals: Not reported in the article. [3/103 withdrew from the acupuncture group, 5/105 withdrew from the sulfasalazine group]
Total duration: 23 to 49 days treatment followed by an evaluation, no follow up reported. ("A course of treatment included 10 sessions (days) acupuncture and there was a break of 3 days between two courses. Participants were administrated totally 2 to 4 courses")
Type of analysis reported: Not specified
Setting: Outpatient and inpatient in Gansu Provincial Tumor Hospital and Gansu Provincial Hospital of Traditional Chinese Medicine, China
Mean age (+/- Standard Deviation or Range): 37.77 (9.70) years for acupuncture group, 38.40 (9.65) years for sulfasalazine group
Men/Women (number/number): 99/101

Acupuncture for treatment of irritable bowel syndrome (Review)



Xue 2009 (Continued)	
	Acu/Moxa: 48/52, Medication: 51/49
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Rome II criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis
	Actual duration of IBS symptoms before enrollment: 5.17 (3.19) months for acupuncture group, 5.12 (3.66) years for sulfasalazine group
	Diagnosis of IBS required for eligibility?: Rome II criteria
	Evaluation to rule out organic gastrointestinal disease: type-B ultrasonic, endoscopy, colonoscopy to exclude other digestive disease Predominant symptoms or stool patterns of participants: Not reported
	Hospital inpatients? Yes, the number of inpatients was not reported Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	ACUPUNCTURE + Moxibustion Number allocated to acupuncture: 103 [210 participants were randomly assigned to two groups, 105 for both groups originally, which was different with that documented in article (i.e. 100 participants). During the trial, 3 withdrew in acupuncture group, 5 withdrew in sulfasalazine group. To reach an equal sample size in both groups, two participants who should have been assigned to acupuncture group were not recruited] Style of acupuncture: Chinese Points selection: Fixed Points stimulated: Six fixed acupoints including ST 25, CV 8, ST 36, PC 6, LI 4 and LV 3
	Description of acupuncturist qualifications: Not reported
	Total length of treatment period (weeks): 23 to 49 days ("A course of treatment included 10 sessions (days) acupuncture and there was a break of 3 days between two courses. Participants were adminis- trated totally 2 to 4 courses.") Number of sessions target (mean): 20 to 40 Times per week: 7 Number of points used (mean): 6 fixed acupoints, 5 of which are bilateral Insertion depth: Not reported Was De Qi reportedly sought? Not reported Duration (minutes): [20 min] Method of stimulation: Manipulation with even method (Ping bu ping xie)
	Moxibustion procedure: Moxibustion was administrated on CV 8 for 5 min
	CONTROL GROUP A INTERVENTION:
	Sulfasalazine (Company: Shanghai Sanwei Pharmaceutical Co., Ltd)
	Number allocated to control group A:
	105 (as described above)
	Total length of treatment period: 20 to 40 days Medication – Sulfasalazine (SASP), 10mg/kg, once daily, course of 10 days, total 2 to 4 courses
	Number of sessions target (mean): 20 to 40
	Days per week: 7
	Times per day: 1
	Dosage: 10 mg/kg
	Any co-interventions in all groups? Not reported

Acupuncture for treatment of irritable bowel syndrome (Review)

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Xue 2009 (Continued)	
Outcomes	Symptom severity:
	Overall IBS symptom (3-point scoring system) at end of treatment: 1) cured 2) improved 3) no effect. The outcome was measured as responder rates for both groups, but there's no clear description of the evaluation basis or procedure
	Quality of life: Not assessed
	Time points of symptom measurement: The responder rates of overall IBS symptom were reported at the end of treatment (23 to 49 days) for both groups
	Author's conclusion: "Evident Acupuncture treatment effect on patients with IBS." (Abstract)
	Additional outcomes reported in the trial but not extracted: The SOD, MDA and NO in patient plasma were reported at baseline and end of treatment The evaluators who asked the patients the symptom score questions were blind to the participant assignment
	Adverse effects: Not reported.
Notes	[The number of participants assigned to either group was originally 105, which was different from that documented in the article (i.e., the article stated that 100 participants were assigned to each group), but the investigator could not describe clearly the method used to ensure exactly the same number of participants assigned to either treatment group. During the trial, 3 withdrew in acupuncture group, 5 withdrew in sulfasalazine group, leaving 102 in the acupuncture group and 100 in the sulfasalazine group. To reach an equal sample size in both groups, 2 participants who were assigned to the acupuncture group were not included in the analysis. So finally, there were 100 participants analyzed in the acupuncture group]
	Funding sources: Natural science foundation of Gansu Province, China (Grant No. 3ZS041 - A25 - 071)
	Information in square brackets was provided by one of the study personnel Kou Suotang, in telephone surveys on July 25 and 28, 2011
	We dichotomized cured, improved and no effect into the 2 categories of improved or not improved ac- cording to the methods section of the review

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	[We referred to a random number table]
Allocation concealment (selection bias)	Unclear risk	[We used sequentially numbered, opaque, sealed envelopes]
Blinding (performance bias and detection bias) All outcomes	High risk	[The evaluators were blind to the participant assignment]
Incomplete outcome data (attrition bias) Short term	Low risk	[The number of participants assigned to either group was originally 105, which was different with that documented in article (i.e. 100 participants). During the trial, 3 withdrew in acupuncture group, 5 withdrew in sulfasalazine group. To reach an equal sample size in both groups, 2 participants who should be as- signed to acupuncture group were finally not been recruited. So finally, there were 103 participants assigned to acupuncture group and 105 participants as- signed to sulfasalazine group]
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall IBS symptom at the completion of the treatment period, together with results of SOD, MDA and NO in patients' plas-

Acupuncture for treatment of irritable bowel syndrome (Review)



Xue 2009 (Continued)		ma before and after treatment according to the Observation item and meth- ods section
Other bias	Unclear risk	Although ITT analysis was not reported, there were few drop-outs, so the de- gree of "missingness" would be unlikely to affect the estimate of the treatment effect in this trial. The two groups were comparable at baseline

Zeng 2010	
Methods	Design: Parallel Blinding: No Attempt to confirm patient blinding?: No Dropouts/withdrawals: During the 4 weeks treatment, 4/33 drop-outs in acupuncture group and 1/32 intrimebutine maleate group
	Total duration: 4 weeks treatment immediately followed by an evaluation, 2 month follow up reported
	Type of analysis reported: Not specified
Participants	Setting: Acupuncture outpatient and inpatient in Hangzhou Hospital of TCM from 2008 to 2009, China
	Mean age (+/- Standard Deviation or Range): 35.15 (7.23) years for acupuncture group, 34.70 (6.45) years for trimebutine maleate group
	Men/Women (number/number): 27/38 Acu/Moxa: 13/20, Medication: 14/18
	Recruitment method: Not reported
	Minimum duration of IBS symptoms: Rome III criteria for the last 3 months with symptom onset at least 6 months prior to diagnosis
	Actual duration of IBS symptoms before enrollment: 15.24 (7.85) months for acupuncture group, 14.86 (7.77) months for trimebutine maleate group
	Diagnosis of IBS required for eligibility?: Rome III criteria for diarrhea-predominant irritable bowel syn- drome
	Evaluation to rule out organic gastrointestinal disease: Reported – exclusion criterion was no organic bowel disease and no other biochemical abnormalities
	Predominant symptoms or stool patterns of participants: The study only recruited patients with diar- rhea-predominant IBS
	Hospital inpatients?: Yes, the number of inpatients was not reported Were people with a history of acupuncture treatment excluded?: Not reported
Interventions	ACUPUNCTURE + MOXIBUSTION Number allocated to acupuncture: 33 Style of acupuncture: Chinese Points selection: Flexible formula Points stimulated: 5 main fixed acupoints and 6 optional points. The fixed points are GV 7, CV 12, CV 4, CV 6 and ST 37. The optional points were LV 3, SP 9, BL 20, BL 21, BL 23 and BL 25 Description of acupuncturist qualifications: Not reported Total length of treatment period (weeks): ~4 weeks (30 days) Number of sessions target (mean): 30 Times per week: 7 Number of points used (mean): 5 fixed acupoints plus 6 optional points Insertion depth: Not reported Was De Qi reportedly sought?: Not reported Duration (minutes): 20 min

Acupuncture for treatment of irritable bowel syndrome (Review)



tion (selection bias)	
Bias Random sequence genera-	Authors' judgement Support for judgement Low risk [Referred to a random number table]
	Authors' judgement Support for judgement
Risk of bias	
	We dichotomized improved, effective and no effect into the 2 categories of improved or not improved according to the methods section of the review
	Information in square brackets was provided by the second author Bao YH, in a telephone survey onJu- ly 22, 2011
Notes	Funding sources: [No funding]
	Adverse effects: Not reported
	Additional outcomes reported in the trial but not extracted: The Bristol score (mean and SD) were re- ported at baseline, 4 weeks (30 days) and 2 months follow up (Table 2). [the evaluators were blind to the participant assignment]
	Author's conclusion: "Moxibustion in dog days could improve the symptom of diarrhea-predominant IBS patients"
	The numbers of responders for symptom improvement were reported at 4 weeks
	The Overall abdominal symptom scale (mean and SD)were reported at baseline, 4 weeks (30 days) and 2 months follow up
	Time points of symptom measurement:
	Quality of life: Not assessed
	 Symptom improvement(3-point scoring system) at end of treatment: 1) markedly effective 2) effective 3) no effect. The evaluation of symptom improvement was based on an intestinal symptoms scale. markedly effective: the decrease of intestinal symptom scale > 80%, 2) effective: the decrease of intestinal symptom scale is between 50% and 80%; 3) no effect: the decrease of intestinal symptom scale < 50% or worsening symptoms. Measured as number of responders for both groups
	1. Overall abdominal symptom scale.
Outcomes	Symptom severity:
	Any co-interventions in all groups? Not reported
	CONTROL GROUP A INTERVENTION: trimebutine maleate (product name: Niweifu) Number allocated to control group A: 32 Total length of treatment period: ~4 weeks (30 days) Number of sessions target (mean): 30 days per week: 7 Times per day: 3 Dosage: 100mg / time
	Moxibustion procedure: After acupuncture, moxibustion was administrated mainly on CV 12, CV 6, CV 4, BL 20, BL 21 and BL 23. The moxa was made into a hemisphere with diameter of 4 cm and height of 2.5 cm, then it was put into a moxibustion box and ignited. The duration of moxibustion on each acupoint was 30 min (The duration of moxibustion was not reported in the article. The information was provided by the 3 rd author Chu JM, in a telephone survey on July 26, 2011)
Zeng 2010 (Continued)	Method of stimulation: Manipulation with even method (Ping bu ping xie)

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Zeng 2010 (Continued)

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Allocation concealment (selection bias)	Low risk	[We used sequentially numbered, opaque, sealed envelopes]
Blinding (performance bias and detection bias) All outcomes	High risk	[The evaluators were blind to the participant assignment]
Incomplete outcome data (attrition bias) Short term	Low risk	"4 participants in acupuncture group withdrew and 1 participant in trimebu- tine mleate group withdrew during the treatment." There is no information about the missing outcome data
Selective reporting (re- porting bias)	Low risk	This trial reported outcomes for overall abdominal symptom scale and the number of responders at the completion of the 30 days treatment period, to- gether with results for each outcome measured according to the observational targets
Other bias	Unclear risk	Although ITT analysis was not reported, there were few drop-outs, so the de- gree of "missingness" would be unlikely to affect the estimate of the treatment effect in this trial
		The two treatment groups were comparable at baseline in terms of the vari- ables Bristol scale and overall abdominal symptom scale. The two treatment groups were also comparable at baseline in terms of demographic characteris- tics and duration of IBS

Characteristics of excluded studies [ordered by study ID]

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Study	Reason for exclusion	
Chan 1997	Non-randomized pilot study with 7 patients	
Chen 2005	Co-intervention of ear acupressure was only given to acupuncture group. This trial is the same as the Li 2005 trial with regard to acupuncture method, treatment duration, age range, sex ratio and disease duration of participants, however it has a different control intervention and different re- sults. It is a possible duplicate of the Li 2005 trial	
Chu 2009	According to the description on Clinicaltrials.gov (http://clinicaltrials.gov/ct2/show/NCT00900965), this study, which was ongoing at the time of the original 2006 Cochrane review and has since been completed, involves only one acupuncture treatment and does not measure any of the outcomes relevant to this review (i.e. its purpose is to investigate the effects of acupuncture on rectal sensa- tion and brain activity in patients with IBS)	
Ding 2004	The first author Ding was telephoned. She said she could not recall if the random sequence was used and also said that some participants were treated with both acupuncture and western medi- cine	
Fireman 2001	One of the co-authors, Rafael Carasso, stated (to Eric Manheimer during a telephone interview on December 20, 2011) that participants were assigned to comparison groups according to the order in which they came into the clinic, which is not a randomization procedure	
Fu 1993	The second author Cai was telephoned. He said the participants were assigned based on their ad- mission order	
Gabuzian 1994	This study did not appear to be eligible based on the translated abstract, but we did not translate the full report to confirm this (language barrier to Russian)	

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Study	Reason for exclusion According to a telephone survey of the first author Guo GL on Jul 22, 2011, the participants were assigned by odd or even hospital record number. This trial was published first in Apr 2010 in Hebei Journal of Traditional Chinese Medicine, and as a duplicate publication in Jun 2010 in Modern Journal of Integrated Traditional Chinese and Western Medicine		
Guo 2010			
Hou 2011	According to a telephone survey of the first author Hou BG on Dec 1, 2011, the participants were as- signed according to their preference		
Huang 1999	The trial compared acupuncture plus Chinese medicine plus psychological counselling versus Chi- nese medicine alone.		
Huang 2006	According to the survey from the first author, they referred to a random number table, but some patients were assigned according to their will		
	The authors compared acupuncture with a different form of Chinese medicine which is Tuina (i.e. Acupuncture plus Tuina versus Acupuncture versus Tuina).		
Kang 2006	The trial compared ear-acupressure versus western medication. Ear-acupressure alone does not meet our inclusion criteria for type of intervention		
Kunze 1990	We could not determine whether this study was eligible based on the translated abstract (the full publication was in German and there was a language barrier to German.)		
Li 2005	Co-intervention of ear acupressure was only given to acupuncture group. Note this is a possible d plicate of the Chen 2005 trial		
Li 2007	The authors didn't report a measure of overall IBS symptoms, overall IBS response, or health-relat- ed quality of life but only had an improvement measure for each of the individual symptoms (e.g. diarrhea, abdominal pain etc.)		
Liao 2000	The author was telephoned and it was confirmed that this was not a real randomized trial		
Liu 1995	The author could not recall any of the methods used in the trial so we cannot assume that this wa a true randomized trial		
Long 2006	The trial compared acupuncture plus bifidobacteria versus bifidobacteria plus patented Chinese medicine versus acupuncture alone, which does not meet our inclusion criteria for type of interven tion		
Ma 2009	The trial compared warming needle moxibustion plus acupoint injection therapy versus western medicine		
Ouyang 2003	The trial compared ear-acupressure plus pinaverium bromide versus ear-acupressure alone versus pinaverium bromide alone. Ear-acupressure alone does not meet our inclusion criteria for type of intervention		
Ouyang 2006	According to the description on Clinicaltrials.gov (http://clinicaltrials.gov/ct2/show/NCT00219505) this study, which was ongoing at the time of the original review, was terminated and not completed		
Qian 2011	This trial clearly states in the text, on page 258, column 2, last line, that the patients were not blind- ed as to whether they got true or sham acupuncture, which is the purpose of using a sham control. Therefore this trial cannot be compared or combined with other sham-controlled trials.		
Shi 2011	According to a telephone survey of the first author Shi ZM on Jul 22, 2011, the participants were as- signed casually, not using any specific randomization method. This trial was published first in Oct 2010 in Hebei Journal of Traditional Chinese Medicine, and as duplicate publications in Apr 2011 in		

Acupuncture for treatment of irritable bowel syndrome (Review)

Study	Reason for exclusion			
	Liaoning Journal of Traditional Chinese Medicine and in Jul 2011 in Chinese Acupuncture & Moxi- bustion			
Song 2005	According to a telephone survey of the first author Song JY on Jul 23, 2011, the participants were inpatient and were administrated with western medicine. If the symptoms of patients were not im proved, they were treated with acupuncture after consultation			
Song 2011	The trial compared ear-acupressure plus otilonium bromide versus otilonium bromide alone. Ear- acupressure alone does not meet our inclusion criteria for type of intervention			
Sun 2009	The trial compared acupuncture plus Chinese medicine versus western medicine			
Wang 2002	Complicated intervention: acupuncture with moxa on the tip of the needle, plus intra-rectum ene- ma with Chinese herb solution			
Wang 2007	According to the survey from the first author, patients were assigned according to their will. This trial compared acupuncture versus western medicine plus Traditional Chinese herbal medicine			
Wang 2008a	The authors didn't report an outcome for overall IBS symptom severity or health-related quality of life but only had an improvement measure for each of the individual symptoms (e.g. diarrhea, ab- dominal pain etc.)			
Wang 2008b	We are unable to contact the only author for information about randomization			
Wang 2009	The authors used a kind of trigger point therapy which does not meet our inclusion criteria for types of intervention			
Wang 2011	The trial used eye-acupuncture, which is a type of micropuncture that is not considered traditional Chinese acupuncture, and has not been approved popularly			
Wu 2004	The trial compared acupuncture plus ear-acupressure vs. acupuncture alone, which does not me our inclusion criteria for types of intervention			
Wu 2008	The trial compared acupuncture on ST25 versus acupuncture on SP15.			
Xiao 2003	According to a telephone survey with the first author Xiao FT on Nov 25, 2011, the participants were assigned based on the doctor's judgment. The outcome of the trial was recurrence rate which does not meet our inclusion criteria for types of outcome measures			
Xiao 2004	Evaluation of the therapeutic effect of acupoint transcutaneous electrical nerve stimulation (TENS			
Xiong 2008b	According to a telephone survey of the first author Xiong JW on Jul 22, 2011, the participants were assigned by alternation. Note: Although the author last name and year is the same as the Xiong X 2008 trial included in our review, they are not the same author or trial			
Xu 2009	According to a telephone survey of the first author Xu MF on Jul 21, 2011, the participants were as- signed by their preference			
Yang 2000	Intervention was Vit B1 point (ST-36) injection and Chinese massage			
Yu 2007	According to the survey from the first author, the patients were assigned according to alternation or rotation.			
	The authors compared acupuncture with a different form of Chinese medicine which is Chinese herbal medicine (i.e. acupuncture plus Chinese medicine versus Chinese medicine versus western medicine.)			

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Study	Reason for exclusion
Yu 2011	The trial compared ear-acupressure plus Chinese medicine versus Chinese medicine alone versus ear-acupressure alone. Ear-acupressure alone does not meet our inclusion criteria for type of intervention
Zhang 1996	The trial used tapping acupuncture which was not inserted into human body, so it does not meet our inclusion criteria for types of interventions. This trial was published first in 1996 in Chinese Acupuncture & Moxibustion, and as a duplicate publication in 1997 in Journal of Jiamusi Medical College (former name of Heilongjiang Medicine and Pharmacy)
Zhang 2000	According to the telephone survey from the first author Zhang, he said he didn't use any random method of allocation
Zhang 2010	The trial compared acupuncture versus Chinese medicine
Zhang 2011a	According to a telephone survey of the first author Zhang Y on Jul 22, 2011, the participants were assigned by rotation
Zhang 2011b	According to a telephone survey of the first author Zhang HC on Dec 1, 2011, some participants were assigned according to the patients' preference
Zhao 2003	The trial compared needle acupuncture versus moxibustion
Zheng 2003	According to the telephone survey from the first author Zheng, he stated they used random method of allocation, but he could not recall the details of the generation of random sequence. He also could not explain why the patient numbers of the two groups were not comparable (41 for Bifidobacterium capsule+pinaverium plus acupuncture group, 28 for Bifidobacterium cap- sule+pinaverium group)

Characteristics of ongoing studies [ordered by study ID]

Anastasi	
Trial name or title	Acupuncture/Moxibustion for Irritable Bowel Syndrome (IBS)
Methods	
Participants	Patients with IBS (N=171)
Interventions	Double Blind RCT comparing
	1: Standard Acu/Moxa (fixed protocol)
	2: Individualized Acupuncture/Moxibustion
	3. Sham Acupuncture/Placebo Moxibustion
Outcomes	Primary outcome: Patient-rated abdominal pain/discomfort and IBS-Secondary symptom rating: intestinal gas, bloating and stool consistency [Time Frame:Six months]
Starting date	Start date: July 2009; Expected date of completion: March 2013
Contact information	Ann M Chung
	212-922-7038
	amc20@nyu.edu

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Anastasi (Continued)

Notes

ClinicalTrials.gov identifier: NCT00945074 Location: New York University, Division of Special Studies in Symptom Management Sponsor: New York University, National Institutes of Health (NIH) Other Study ID numbers: 7833948;R01NR010730

MacPherson	
Trial name or title	Acupuncture for Irritable Bowel Syndrome (IBS)
Methods	
Participants	Patient with IBS (N=220)
Interventions	RCT comparing acupuncture plus usual GP care versus usual GP care alone.
Outcomes	Primary outcome measures:
	 IBS Symptom Severity Score (IBS SSS) at 3 months Cost-effectiveness study: European quality of life questionnaire (EQ-5D) at 12 months Medication use at baseline, 3 months, 6 months, 9 months and 12 months Health services used at baseline, 3 months, 6 months, 9 months and 12 months Days lost from work at baseline, 3 months, 6 months, 9 months and 12 months
Starting date	Start date: 01/10/2008, Expected date of completion: 30/09/2011
Contact information	Dr Hugh MacPherson
	+44 (0)1904 321394
	hm18@york.ac.uk
Notes	Location:United Kingdom
	Sponsor details: c/o Mrs S Final, Heslington, York, United Kingdom, YO10 5DD (smf3@york.ac.uk, +44 (0)1904 430000)
	Link to record in ISRCTN Register: ISRCTN08827905

DATA AND ANALYSES

Comparison 1. Acupuncture versus sham acupuncture

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptom severity (mean outcomes) post- treatment scores (short-term: up to 3 mos, closest to 8 wks)	3	231	Std. Mean Difference (IV, Random, 95% CI)	-0.09 [-0.35, 0.17]

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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
2 Symptom severity (responders) (short-ter- m:up to 3 mons, closest to 8 wks)	3	262	Risk Ratio (M-H, Ran- dom, 95% CI)	1.09 [0.87, 1.37]
3 Symptom severity (generic inverse vari- ance) post-treatment scores (short-term: up to 3 mos, closest to 8 wks)	4	281	Std. Mean Difference (Random, 95% CI)	-0.11 [-0.35, 0.13]
4 Quality of life (mean outcomes) post-treat- ment scores (short-term: up to 3 mos, closest to 8 wks)	3	253	Std. Mean Difference (IV, Random, 95% CI)	-0.03 [-0.27, 0.22]
5 Quality of life (responders) (short-term:up to 3 mons, closest to 8 wks)	1	153	Risk Ratio (M-H, Ran- dom, 95% CI)	1.25 [0.58, 2.68]
6 Quality of life (mean outcomes) post-treat- ment scores (long-term: >3 mos, closest to 6 mos)	1	41	Std. Mean Difference (IV, Random, 95% CI)	0.07 [-0.54, 0.69]

Analysis 1.1. Comparison 1 Acupuncture versus sham acupuncture, Outcome 1 Symptom severity (mean outcomes) post-treatment scores (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	Acu	puncture	Sham a	acupuncture		Std. Mean Difference		Weight		Std. Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)		Rar	ndom, 95% CI			Random, 95% Cl	
Anastasi 2009	12	2.1 (1.3)	15	1.7 (1.4)			+	\rightarrow	11.57%	0.25[-0.51,1.01]	
Forbes 2005	22	11.4 (5.1)	29	10.9 (4.2)					21.87%	0.11[-0.45,0.66]	
Lembo 2009	78	174.5 (98.3)	75	193.6 (83.4)					66.57%	-0.21[-0.53,0.11]	
Total ***	112		119						100%	-0.09[-0.35,0.17]	
Heterogeneity: Tau ² =0; Chi ² =1	79, df=2(P=0.4	1); I ² =0%									
Test for overall effect: Z=0.65(P=0.52)										
			Favours	acupuncture	-1	-0.5	0 0.5	1	Favours sh	am acupuncture	

Analysis 1.2. Comparison 1 Acupuncture versus sham acupuncture, Outcome 2 Symptom severity (responders) (short-term:up to 3 mons, closest to 8 wks).

Study or subgroup	Acupuncture	Acupuncture Sham acupuncture		Risk Ratio		Weight	Risk Ratio
	n/N	n/N	M-H	, Random, 95% CI			M-H, Random, 95% CI
Forbes 2005	11/27	10/32		+		11.09%	1.3[0.66,2.59]
Lembo 2009	46/78	43/75	-	—— <mark>—</mark> ——		72.19%	1.03[0.79,1.35]
Lowe 2000	16/28	10/22		+		16.72%	1.26[0.72,2.2]
Total (95% CI)	133	129				100%	1.09[0.87,1.37]
Total events: 73 (Acupunctur	e), 63 (Sham acupuncture)						
Heterogeneity: Tau ² =0; Chi ² =	0.7, df=2(P=0.7); l ² =0%						
Test for overall effect: Z=0.76	(P=0.45)		1 1				
	Favours sl	nam acupuncture	0.5 0.7	1 1	.5 2	Favours acupuncture	

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Analysis 1.3. Comparison 1 Acupuncture versus sham acupuncture, Outcome 3 Symptom severity (generic inverse variance) post-treatment scores (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	Acupunc- ture	Sham acupunc- ture	Std. Mean Difference	S	td. Mean Difference	v	Veight	Std. Mean Difference
	Ν	Ν	(SE)		V, Random, 95% Cl			IV, Random, 95% CI
Anastasi 2009	12	15	0.3 (0.389)	_			9.85%	0.25[-0.51,1.01]
Forbes 2005	22	29	0.1 (0.283)		•	· 1	L8.59%	0.11[-0.45,0.66]
Lembo 2009	78	75	-0.2 (0.162)	_		ŗ	56.66%	-0.21[-0.53,0.11]
Lowe 2000	28	22	-0.3 (0.316)		•		14.9%	-0.26[-0.88,0.36]
Total (95% CI)							100%	-0.11[-0.35,0.13]
Heterogeneity: Tau ² =0; Chi ² =2.	05, df=3(P=0.56); I ² =0%							
Test for overall effect: Z=0.91(P	=0.36)							
		Favor	rs acupuncture	-1 -0.5	0 0.5	1 F	avors sha	m acupuncture

Analysis 1.4. Comparison 1 Acupuncture versus sham acupuncture, Outcome 4 Quality of life (mean outcomes) post-treatment scores (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	Acu	puncture	Shama	acupuncture		Std. M	lean Difference	Weight	Std. Mean Difference
	N	Mean(SD)	Ν	Mean(SD)		Rai	ndom, 95% Cl		Random, 95% CI
Forbes 2005	27	64.6 (19.8)	32	65.6 (19.8)				23.2%	-0.05[-0.56,0.46]
Lembo 2009	78	73.1 (17.3)	75	73.8 (17.6)		_		60.58%	-0.04[-0.36,0.28]
Schneider 2006	21	64 (82.1)	20	60 (64.2)			+	16.23%	0.05[-0.56,0.67]
Total ***	126		127				•	100%	-0.03[-0.27,0.22]
Heterogeneity: Tau ² =0; Chi ² =	0.08, df=2(P=0.9	6); I ² =0%							
Test for overall effect: Z=0.22	(P=0.83)								
		Fave	ours sham	acupuncture	-1	-0.5	0 0.5	¹ Favours	acupuncture

Analysis 1.5. Comparison 1 Acupuncture versus sham acupuncture, Outcome 5 Quality of life (responders) (short-term:up to 3 mons, closest to 8 wks).

Study or subgroup	Acupuncture	Sham acupuncture			Ris	k Rat	io			Weight	Risk Ratio
	n/N	n/N			M-H, Rar	ndom	, 95% CI				M-H, Random, 95% Cl
Lembo 2009	13/78	10/75				-				100%	1.25[0.58,2.68]
Total (95% CI)	78	75								100%	1.25[0.58,2.68]
Total events: 13 (Acupuncture),	, 10 (Sham acupuncture)										
Heterogeneity: Not applicable											
Test for overall effect: Z=0.57(P	=0.57)										
	Favours sl	nam acupuncture	0.1	0.2	0.5	1	2	5	10	Favours acupuncture	2

Analysis 1.6. Comparison 1 Acupuncture versus sham acupuncture, Outcome 6 Quality of life (mean outcomes) post-treatment scores (long-term: >3 mos, closest to 6 mos).

Study or subgroup	Acu	Acupuncture Sha		acupuncture	Std. Mean Difference	Weight	Std. Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)	Random, 95% CI		Random, 95% CI	
Schneider 2006	21	63 (77.4)	20	57 (79.7)		100%	0.07[-0.54,0.69]	
Total ***	21		20		•	100%	0.07[-0.54,0.69]	
Heterogeneity: Not applicable								
Test for overall effect: Z=0.24(P=0.81)							
		Fav	ours sham	acupuncture	-5 -2.5 0 2.5 5	Favours ad	cupuncture	

Comparison 2. Acupuncture versus pharmacological medication

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptom severity (responders) (short-term: up to 3 mons, closest to 8 wks)	5	449	Risk Ratio (M-H, Ran- dom, 95% Cl)	1.28 [1.12, 1.45]
2 Symptom severity (mean outcomes) post- treatment scores (short-term: up to 3 mos, closest to 8 wks)	3	190	Std. Mean Difference (IV, Random, 95% CI)	-0.72 [-1.02, -0.43]

Analysis 2.1. Comparison 2 Acupuncture versus pharmacological medication, Outcome 1 Symptom severity (responders) (short-term: up to 3 mons, closest to 8 wks).

Study or subgroup	Acupuncture	Pharmacologi- cal medicatio		Risk Ratio		Weight	Risk Ratio	
	n/N	n/N		M-H, Random, 95%	CI		M-H, Random, 95% CI	
Chen 2011	25/29	22/30		+-		17.66%	1.18[0.91,1.53]	
Shi 2010	26/32	20/38		-+-		11.43%	1.54[1.09,2.18]	
Sun 2011	27/30	24/30		-		22.94%	1.13[0.91,1.39]	
Xue 2009	81/100	55/100		+		24.94%	1.47[1.2,1.8]	
Zeng 2010	27/29	24/31		*		23.03%	1.2[0.97,1.49]	
Total (95% CI)	220	229		•		100%	1.28[1.12,1.45]	
Total events: 186 (Acupunctu	ıre), 145 (Pharmacological n	nedicatio)						
Heterogeneity: Tau ² =0.01; Ch	ii ² =5.83, df=4(P=0.21); l ² =31.	43%						
Test for overall effect: Z=3.73	(P=0)							
	Favour	s pharmacological	0.01 0.	.1 1	10 100	^D Favours acupuncture		

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Analysis 2.2. Comparison 2 Acupuncture versus pharmacological medication, Outcome 2 Symptom severity (mean outcomes) post-treatment scores (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	Acu	Acupuncture		macolog- medicatio	Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	Random, 95% CI		Random, 95% Cl
Shi 2010	32	2.7 (1.1)	38	4.1 (1.9)	-	35.31%	-0.93[-1.43,-0.44]
Sun 2011	30	2.9 (1.7)	30	4.5 (2.7)	-#-	31.92%	-0.7[-1.22,-0.17]
Zeng 2010	29	4.2 (2.1)	31	5.6 (2.9)	-	32.77%	-0.52[-1.03,-0]
Total ***	91		99		•	100%	-0.72[-1.02,-0.43]
Heterogeneity: Tau ² =0; Chi ² =	1.31, df=2(P=0.5	2); I ² =0%					
Test for overall effect: Z=4.79	(P<0.0001)						
			Favours	acupuncture	-5 -2.5 0 2.5	5 Favours m	edication

Comparison 3. Acupuncture versus Bifidobacterium

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptom severity (responders) (short- term: up to 3 mons, closest to 8 wks)	2	181	Risk Ratio (M-H, Ran- dom, 95% CI)	1.07 [0.90, 1.27]

Analysis 3.1. Comparison 3 Acupuncture versus Bifidobacterium, Outcome 1 Symptom severity (responders) (short-term: up to 3 mons, closest to 8 wks).

Study or subgroup	Acupuncture	Bifidobac- terium		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		М-Н, Я	andom, 95% Cl			M-H, Random, 95% CI
An 2010	41/47	26/34			+		61.84%	1.14[0.92,1.42]
Liu 2010	33/50	34/50			+		38.16%	0.97[0.74,1.28]
Total (95% CI)	97	84			•		100%	1.07[0.9,1.27]
Total events: 74 (Acupuncture	e), 60 (Bifidobacterium)							
Heterogeneity: Tau ² =0; Chi ² =	0.88, df=1(P=0.35); I ² =0%							
Test for overall effect: Z=0.81	(P=0.42)							
	Favour	s Bifidobacterium	0.01	0.1	1 1	0 100	Favours acupuncture	9

Comparison 4. Acupuncture versus psychotherapy

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptom severity (responders) (short- term: up to 3 mons, closest to 8 wks)	1	100	Risk Ratio (M-H, Ran- dom, 95% CI)	1.05 [0.87, 1.26]

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Analysis 4.1. Comparison 4 Acupuncture versus psychotherapy, Outcome 1 Symptom severity (responders) (short-term: up to 3 mons, closest to 8 wks).

Study or subgroup	Acupuncture	Psychotherapy		Risk Ra	tio		Weight	Risk Ratio
	n/N	n/N	M-H	, Random	, 95 %	CI		M-H, Random, 95% CI
Liu 1997	42/50	40/50		+			100%	1.05[0.87,1.26]
Total (95% CI)	50	50		•			100%	1.05[0.87,1.26]
Total events: 42 (Acupuncture), 40 (P	sychotherapy)							
Heterogeneity: Not applicable								
Test for overall effect: Z=0.52(P=0.6)								
	Favo	urs psychotherapy	0.1 0.2	0.5 1	2	5 10	Favours acupuncture	

Comparison 5. Acupuncture plus psychotherapy versus psychotherapy alone

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Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptom severity (responders) (short- term: up to 3 mons, closest to 8 wks)	2	182	Risk Ratio (M-H, Ran- dom, 95% CI)	1.16 [1.04, 1.29]

Analysis 5.1. Comparison 5 Acupuncture plus psychotherapy versus psychotherapy alone, Outcome 1 Symptom severity (responders) (short-term: up to 3 mons, closest to 8 wks).

Study or subgroup	or subgroup Acupuncture No acupunc- Risk Ratio as adjunct ture		Weight	Risk Ratio	
	n/N	n/N	M-H, Random, 95% CI		M-H, Random, 95% CI
Liu 1997	48/50	40/50		48.99%	1.2[1.03,1.39]
Xiong 2008a	40/42	34/40	**	51.01%	1.12[0.97,1.3]
Total (95% CI)	92	90	•	100%	1.16[1.04,1.29]
Total events: 88 (Acupunctur	e as adjunct), 74 (No acupur	ncture)			
Heterogeneity: Tau ² =0; Chi ² =	0.42, df=1(P=0.52); I ² =0%				
Test for overall effect: Z=2.76	i(P=0.01)				
	Favou	rs no acupuncture	0.1 0.2 0.5 1 2 5 10	Favours acupunctu	re

Favours no acupuncture Favours acupuncture

Comparison 6. Acupuncture plus another traditional Chinese medicine (TCM) treatment versus the other TCM treatment alone

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptom severity (responders) (short- term: up to 3 mos, closest to 8 wks)	4	466	Risk Ratio (M-H, Random, 95% CI)	1.17 [1.02, 1.33]
2 Quality of life (mean outcomes) (short- term: up to 3 mos, closest to 8 wks)	1	60	Std. Mean Difference (IV, Random, 95% CI)	0.48 [-0.03, 0.99]

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Analysis 6.1. Comparison 6 Acupuncture plus another traditional Chinese medicine (TCM) treatment versus the other TCM treatment alone, Outcome 1 Symptom severity (responders) (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	y or subgroup Adjunct Other Risk Ratio acupuncture TCM alone				Weight	Risk Ratio			
	n/N	n/N		М-Н, Б	andom, 959	% CI			M-H, Random, 95% CI
Chen 2009	29/30	28/30			•			28.64%	1.04[0.92,1.16]
Li 2009	60/62	44/62			+			23.37%	1.36[1.16,1.61]
Liu 2010	135/150	37/50			+			22.64%	1.22[1.02,1.45]
Xiong 2008a	40/42	34/40			•			25.35%	1.12[0.97,1.3]
Total (95% CI)	284	182			•			100%	1.17[1.02,1.33]
Total events: 264 (Adjunct ac	upuncture), 143 (Other TCM	alone)							
Heterogeneity: Tau ² =0.01; Ch	i ² =9.56, df=3(P=0.02); l ² =68.6	52%							
Test for overall effect: Z=2.3(P=0.02)		1						
	Favour	s no acupuncture	0.01	0.1	1	10	100	Favours acupuncture	!

Analysis 6.2. Comparison 6 Acupuncture plus another traditional Chinese medicine (TCM) treatment versus the other TCM treatment alone, Outcome 2 Quality of life (mean outcomes) (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	Adjunct acupuncture		Other	Other TCM alone		Std. Mean Difference			Weight	Std. Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)		Ran	dom, 95% (:1			Random, 95% CI
Chen 2009	30	175.6 (7.4)	30	171.6 (8.9)						100%	0.48[-0.03,0.99]
Total ***	30		30							100%	0.48[-0.03,0.99]
Heterogeneity: Not applicable											
Test for overall effect: Z=1.83(P=0.07)											
			Favours no	acupuncture	-100	-50	0	50	100	Favours ac	upuncture

Comparison 7. Acupuncture versus no specific treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Symptom severity (responders) (short- term: up to 3 mons, closest to 8 wks)	2	181	Risk Ratio (M-H, Ran- dom, 95% CI)	2.11 [1.18, 3.79]
2 Symptom severity (mean outcomes) post- treatment scores (short-term: up to 3 mos, closest to 8 wks)	2	181	Std. Mean Difference (IV, Random, 95% CI)	-1.29 [-2.59, 0.01]
3 Quality of life (mean outcomes) post-treat- ment scores (short-term: up to 3 mos, clos- est to 8 wks)	1	155	Std. Mean Difference (IV, Random, 95% CI)	0.34 [0.02, 0.66]
4 Quality of life (responders) (short-term: up to 3 mons, closest to 8 wks)	1	155	Risk Ratio (M-H, Ran- dom, 95% CI)	1.43 [0.65, 3.14]

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Analysis 7.1. Comparison 7 Acupuncture versus no specific treatment, Outcome 1 Symptom severity (responders) (short-term: up to 3 mons, closest to 8 wks).

Study or subgroup	Acupuncture	No treatment		Risk Ratio		Weight	Risk Ratio	
	n/N	n/N	M-H, Random, 95% Cl				M-H, Random, 95% Cl	
Lembo 2009	46/78	27/77				62.96%	1.68[1.18,2.4]	
Reynolds 2008	9/9	5/17			—	37.04%	3.11[1.53,6.33]	
Total (95% CI)	87	94		-	•	100%	2.11[1.18,3.79]	
Total events: 55 (Acupunctur	e), 32 (No treatment)							
Heterogeneity: Tau ² =0.11; Ch	i ² =2.31, df=1(P=0.13); l ² =56	75%						
Test for overall effect: Z=2.51	(P=0.01)					_		
	Fav	ours no treatment	0.1 0.2	0.5 1 2	5 10	Favours acupuncture		

Analysis 7.2. Comparison 7 Acupuncture versus no specific treatment, Outcome 2 Symptom severity (mean outcomes) post-treatment scores (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	Acu	puncture	Not	reatment		Std. M	lean Difference		Weight	Std. Mean Difference
	N	Mean(SD)	Ν	Mean(SD)		Rai	ndom, 95% CI			Random, 95% Cl
Lembo 2009	78	174.5 (98.3)	77	237 (76)			÷		56.63%	-0.71[-1.03,-0.38]
Reynolds 2008	9	128 (79.5)	17	287 (73.1)		-	₽		43.37%	-2.05[-3.05,-1.04]
Total ***	87		94				•		100%	-1.29[-2.59,0.01]
Heterogeneity: Tau ² =0.75; Chi	² =6.12, df=1(P=	0.01); l ² =83.67%								
Test for overall effect: Z=1.94(P=0.05)				1					
			Favours	acupuncture	-10	-5	0 5	10	Favours no	treatment

Analysis 7.3. Comparison 7 Acupuncture versus no specific treatment, Outcome 3 Quality of life (mean outcomes) post-treatment scores (short-term: up to 3 mos, closest to 8 wks).

Study or subgroup	Acupuncture		No treatment		Std. Mean Difference	Weight	Std. Mean Difference
	Ν	Mean(SD)	Ν	Mean(SD)	Random, 95% Cl		Random, 95% CI
Lembo 2009	78	73.8 (17.3)	77	67.9 (17.3)	-	100%	0.34[0.02,0.66]
Total ***	78		77		•	100%	0.34[0.02,0.66]
Heterogeneity: Not applicable							
Test for overall effect: Z=2.1(P=0.04)							
			Favours	no treatment	-2 -1 0 1 2	Favours ac	cupuncture

Analysis 7.4. Comparison 7 Acupuncture versus no specific treatment, Outcome 4 Quality of life (responders) (short-term: up to 3 mons, closest to 8 wks).

Study or subgroup	Acupuncture	No treatment			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		м-н,	Random, 95	5% CI			M-H, Random, 95% Cl
Lembo 2009	13/78	9/77			-			100%	1.43[0.65,3.14]
Total (95% CI)	78	77			•			100%	1.43[0.65,3.14]
	Fav	ours no treatment	0.01	0.1	1	10	100	Favours acupuncture	

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Study or subgroup	Acupuncture n/N	No treatment n/N			Risk Ratio	-		Weight	Risk Ratio M-H, Random, 95% Cl
Total events: 13 (Acupuncture), 9	•			,					
Heterogeneity: Not applicable									
Test for overall effect: Z=0.88(P=0	0.38)								
	Fa	vours no treatment	0.01	0.1	1	10	100	Favours acupuncture	!

APPENDICES

MEDLINE (OVID)	EMBASE (OVID)	PUBMED	CENTRAL	Cochrane IBD/FBD Group Special- ized Reg- ister	Cochrane Comple- mentary and Al- ternative Medicine (CAM) Field Spe- cialized Register	Chinese CNKI Database	Chinese VIP Data- base	SinoMed Database (an up- date of the Chi- nese Bio- medical Database (CBM))	Cumula- tive In- dex to Nursing and Allied Health (CINAHL)
1 (colonic disease or colon disease).mp. [mp=proto- col supplementary concept, rare disease supplementary concept, title, original ti- tle, abstract, name of sub- stance word, subject head- ing word, unique identifier] (669) 2 irritable bowel syn- drome.mp. or exp irritable colon/ (6817) 3 ((irritable or functional or spastic) and (bowel or colon)).mp. (15406) 4 1 or 2 or 3 (16050) 5 acupuncture thera- py.mp. or exp acupuncture/ (11126) 6 exp alternative medicine/ or exp acupuncture/ or acupunctur*.mp. (157863) 7 electroacupuncturemp. or exp electroacupuncture/ (2581) 8 acupoints.mp. (1103) 9 electrostimulation ther- apy/ or exp nerve stimu- lation/ or percutaneous electrical nerve stimula- tion.mp. or exp transcuta-	1 colonic disease.mp. or exp colon disease/ 2 irritable bowel syn- drome.mp. or exp irritable colon/ 3 ((irritable or functional or spastic) and (bowel or colon)).mp. [mp=title, ab- stract, subject headings, heading word, drug trade name, original title, device manufacturer, drug manu- facturer name] 4 1 or 2 or 3 5 acupuncture therapy.mp. or exp acupuncture/ 6 exp alternative medi- cine/ or exp acupuncture/ or acupunctur*.mp. 7 electroacupuncture.mp. or exp electroacupuncture/ 8 acupoints.mp. 9 electrostimulation ther- apy/ or exp nerve stimu- lation/ or percutaneous electrical nerve stimula- tion.mp. or exp transcuta- neous nerve stimulation/ 10 auriculoacupunc- ture.mp. 11 5 or 6 or 7 or 8 or 9 or 10	acupunc- ture AND (ibs OR irritable bowel syn- drome) Search date: No- vember 30, 2011 Results: 82	(irritable OR func- tional OR spas- tic) AND (bowel OR colon) AND (acupunc- tur* OR elec- troacupunc- ture OR elec- tro-acupunc- ture OR acupoints OR "per- cutaneous electri- cal nerve stimula- tion" OR auricu- loacupunc- ture) search all text Search date: No- yember	The IBD/ FBD regis- ter is on- line at: http:// ibd-fb- d.cochrane.o uploads/ Group %20Spec %20Regis- ter.pdf (MEDLINE records) and http:// ibd-fb- d.cochrane.o uploads/ Group %20Spec %20Regis- ter.pdf	(acupunc- tur* OR rfiles/ troacupunc- ture OR elec- tro-acupunc- ture OR acupoints OR "per- cutaneous electri- cal nerve stimula- tion" OR rg/sita/- loacupunc- ture)	or (Zhu ti='Jie	bet (i.e., Pinyin): U=(((Chang yi ji +(Chang*Ji re))+(Jie chang*(Gong neng+Nian Ye xing +Guo min +Jing lu- an)))*(Zhen ci+Zhen jiu+Dian zhen+Wen zhen)) U means all text. Search date: No- vember 25, 2011 Results:	Strategy in Chinese phonet- ic alpha- bet (i.e., Pinyin): ((Chang yi ji OR Chang*Ji re) OR (Jie chang AND (Gong neng OR Nian ye xing OR Guo min OR Jing lu- an))) AND (Zhen ci OR Zhen jiu OR Di- an zhen OR Wen zhen) search in Que sheng (Que sheng in- cludes ti- tle, MeSH, keywords, abstract.	((irrita- ble OR functional OR spas- tic) AND (bowel OR colon) AND (acupunc- tur* OR elec- troacupune ture OR elec- tro-acupune ture OR acupoints OR "per- cutaneous electri- cal nerve stimula- tion" OR auricu- loacupunc- ture)) and SU acupunc- ture Search modes -

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C 👌	(Continued)							
Acupuncture for treatment of irritable bowel syndrome (Review) Copyright © 2012 The Cochrane Collaboration. Published by John Wiley & Sons, Ltd	neous nerve stimulation/	12 randomized controlled	Results: 28	ister%20-		an zhen'	journal	Boolean/
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 Search date: November 22, 2011

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 2011

 29 4 and 11 and 28 (281)
 Results: 633

Search date: November 28, 2011

Results: 274



Footnotes

Appendix 2. Characteristics of randomized controlled trials in acupuncture for irritable bowel syndrome (IBS)

Study	N a	Country	Diagnos- tic criteria used for IBS	Criteria for improve- ment in overall IBS symptoms	Criteria for improve- ment in IBS-related quality of life	Time point for out- come as- sess-ment ^b	Acupuncture treatment	Control treatment(s)
An 2010	81 c	China	Rome II	Categorical d	_ e	4 wks (EoT)	Fixed formula with moxibus- tion at 2 fixed points; 24 ses- sions over 4 wks	Combined Bifidobacterium, lactobacillus and Enterococ- cus faecium capsules (2 pills 3x/d)
Anastasi 2009	29	USA	Rome II/III	CGI	-	4 wks (EoT)	Flexible formula with mox- ibustion at all points; 8 ses- sions over 4 wks	Sham acupuncture super- ficial needling 2-3 cm from true acupoints
Chen 2009	60 c	China	Rome II	Categorical	FDG QoL scale	4 wks (EoT)	Flexible formula with moxi- bustion at 4 fixed pts; 12 ses- sions over 4 wks + Chinese herbal formula	Chinese herbal formula alone (3 pills, 3x/d)
Chen 2011	60	China	Rome III – IBS-D	Categorical	-	3 wks (EoT)	Flexible formula with mox- ibustion at 3-5 pts; 15 ses- sions over 3 wks	Montmorillonite (1 bag, 2x/ day) + loperamide (4mg, 3x/ day) + pinaverium bromide (50mg, 3x/d) ^f
Forbes 2005	59	UK	Rome I and Manning	Global symptom score based on patient diary g	EuroQol	13 wks (EoT)	Individualized; 10 sessions over 10 wks	Sham acupuncture pene- trating needles at non-acu- points
Lembo 2009 h	230	USA	Rome II	IBS-AR and IBS-SSS	IBS-QoL	3 wks (EoT)	Flexible formula; 6 sessions over 3 wks	 Sham acupuncture - val- idated Streitberger placebo needles placed at non-acu- points in general vicinity of the true acupoints Wait list with continua-
Li 2009	186	China	Rome III	Categorical	-	10 wks (EoT)	Flexible formula with moxi- bustion at all points; 60 ses-	tion of usual care Tuina spinal massage alone (60 sessions over 10 wks)

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(Continued)							sions over 10 wks + Tuina spinal massage	
Liu 1997	150	China	Not speci- fied	Categorical	-	3-21 wks (EoT)	Flexible formula with moxi- bustion at 1 acupoint; 10-60	 Acupuncture alone Psychotherapy alone
							sessions over 20-120 days + psychotherapy	2) Psychotherapy alone
Liu 2010	300	China	Rome III – IBS-D	Categorical	-	4 wks (EoT)	Fixed formula EA; 28 sessions over 4 wks + Chinese herbal	1) EA alone
			0-01				formula	2) Chinese herbal formula alone (150ml, 2x/d)
								3) Bifidobacterim longum alone (1 pill, 2x/d)
Lowe 2000	50	Canada	Rome (ver- sion not stated)	Dichoto- mous mea- sure of symptom relief	IBS-36	4 wks (Eot)	Fixed formula; 8 sessions over 4 wks	Sham acupuncture tap- ping blunt needle on the skin and then taping the needle in place
Reynolds 2008	30	UK	Rome II	IBS-SSS	-	3 mos (EoT)	Flexible formula; 8 sessions over 3 mos	Usual care
Schneider 2006	43	Germ-any	Rome II	Not mea- sured	FDDQL	5 wks (EoT)	Fixed formula; 10 sessions over 5 wks	Sham acupuncture validat ed Streitberger placebo nee dles placed 2 cm from true acupoints
Shi 2010	70	China	Rome III – IBS-D	Categorical	-	5 wks (1 wk after (EoT) ⁱ	Flexible formula EA; 28 ses- sions over 4 wks	Pinaverium bromide (50mg, 3x/d)
Sun 2011	63	China	Rome III – IBS-D	Categorical	-	4 wks (EoT)	Fixed formula; 20 sessions over 4 wks	Pinaverium bromide (50mg, 3x/d)
Xiong 2008a	120	China	Rome II	Categorical	-	4 wks (EoT)	Fixed formula with moxibus- tion at 2 pts; 28 sessions over 4 wks + Chinese herbal for- mula	Chinese herbal formula alone
Xue 2009	210	China	Rome II	Categorical	-	3-7 weeks (EoT)	Fixed formula with moxibus- tion at 1 point; 20-40 ses- sions over 3-7 wks	Sulfasalazine (10 mg, 1x/d)

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Acup	(Continued)								
puncture for	Zeng 2010	65	China	Rome III – IBS-D	Categorical	-	30 d (EoT)	Flexible formula with mox- ibustion at 6 points; 30 ses- sions over 30 d	Trimebutine maleate (100mg, 3x/d)
÷.		L				<u>cl::</u>			

IBS, irritable bowel syndrome; EoT, end of treatment; wks, weeks; d, day; CGI, Clinical Global Impression Scale (Whitehead 2000); acupoints, acupuncture points; FDG QoL, Functional Digestive Diseases Quality of Life Scale, an IBS quality of life questionnaire which had been used in previous Chinese studies but that has not been validated; IBS-D, study restricted eligibility to IBS-D (IBS with diarrhea) subtype patients; EuroQol, EuroQol Group's rating scale (Johnson 1998); IBS-AR, IBS-Adequate Relief question (Mangel 1998); IBS-SSS, IBS Severity Scoring System (Francis 1997); IBS-QoL, IBS Quality of Life Measure (Bijkerk 2003); EA, electroacupuncture; IBS-36 (Groll 2002); FDDQL, Functional Digestive Diseases Quality of Life Questionnaire (Chassany 1999).

^a Number randomized.

^b The time point listed is the number of weeks after randomization.

^c For these 2 trials, the author did not record nor recall the numbers randomized, nor the numbers of dropouts, and the numbers analyzed are reported here.

^d For all 11 trials conducted in China, a symptom scale was used to assess the severity of the patients' overall IBS-related symptoms (e.g., abdominal pain, defecation difficulties, diarrhea) both at baseline and after treatment. For 8 of these trials (Xiong 2008a; Chen 2009; An 2010; Liu 2010; Shi 2010; Zeng 2010; Chen 2011; Sun 2011), a percentage improvement from baseline scores was then calculated (i.e., (baseline symptom score – symptom score after treatment)/ baseline symptom score), and this percentage change from baseline was then grouped into 2 (Shi 2010), 3 (An 2010; Zeng 2010), or 4 (Xiong 2008a; Chen 2009; Liu 2010; Chen 2011; Sun 2011) categories, which were then converted into 2 categories for the meta-analysis, as described in the Methods section. For the other 3 trials (Liu 1997; Xue 2009; Liu 2010), it was not clear how the symptom scale scores were converted into the categorical data. For these 11 trials, the criteria for improvement is listed as "Categorical" in this table.

^e A dash (-) indicates that the outcome not measured.

^f The Montmorillonite was given to all patients and the loperamide and pinaverium bromide was added if the diarrhea did not stop.

g This symptom diary is based on the Bristol scale (Heaton 1991).

^h In this trial, patients were randomized to five arms. The first arm was a wait-list control. Participants in the remaining 4 arms were randomized to sham or true acupuncture, with or without an augmented practitioner-patient interaction. There was no main effect of practitioner-patient interaction; therefore we combined the two acupuncture groups (augmented and limited encounter) and the two sham acupuncture groups (augmented and limited encounter) in order to compare the effects of acupuncture and sham acupuncture.

ⁱ This trial also included an EoT measurement point, for which the results were very similar to the 1 wk post EoT measurement point.

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Appendix 3. Acupuncture adequacy assessments of included studies*

Study	Choice of acupoints	Total num- ber of ses- sions	Treatment duration	Treatment frequency	Needling technique	Experience	Assessment of likelihood of physiological activity of sham control	Guess of study
An 2010	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention	Don't know
Anastasi 2009	Adequate	Adequate	Adequate	Adequate	Adequate	Adequate	Potential for weak physiological activity. This is a shallow-insertion sham acupuncture and the "sham points were 2 to 3 cm away from the traditional true point location and not on a meridian." The superficial insertion of the needles at non-acupoints might cause a phys- iological response, although weaker than that of deeper insertion of needles at true acu- points. Also, as the exact size of an acupunc- ture point is not known, the true acupoints might be stimulated by this superficial needle placement 2 to 3 cm from the true acupoints.	Don't know
Chen 2009	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention	Don't know
Chen 2011	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo controls	Don't know
Forbes 2005	Adequate	Adequate [†]	Adequate	Inadequate (Once per week treat- ment seems not frequent enough.)	Adequate	Adequate	Potential for weak physiological activity. From the description, it sounds like a nee- dle insertion sham at non-acupoints, which might cause a physiological response, al- though weaker than that of insertion of nee- dles at true acupoints	One acupunc- turist (XS) i unaware o this study o its results. The other acupunc- turist (LL) guesses th must be conducted in England but doesn' know who the author are
Lembo 2009	Adequate	Adequate	Adequate	Adequate	Adequate	Adequate	Unlikely to have physiological activity. Use of non-insertion Streitberger sham device in the	Don't knov

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(Continuea)							relative vicinity of the genuine acupoints is an adequate control. While perhaps unlikely, it is possible that the true acupoints may be stim- ulated by the sham needle placement in the vicinity of the true acupoints	
Li 2009	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention	Don't know
Liu 1997	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention	Don't know
Liu 2010	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention	Don't know
Lowe 2000	Don't know [‡]	Adequate	Adequate	Adequate	Don't know	Inadequate [§]	Unlikely to have physiological activity: It seems adequate as this sham only involved "tapping a blunt needle on the skin and tap- ing in place at the same point". Although the shams were placed at the same true acu- points, the blunt needles were only briefly tapped, so would be unlikely to have a thera- peutic effect	Don't know
Reynolds 2008	Adequate	Adequate	Adequate	Inade- quate ^{††}	Adequate ^{‡‡}	Adequate	No sham/placebo intervention group	Don't know
Schneider 2006	Adequate	Adequate	Adequate	Adequate	Adequate	Adequate	Unlikely to have physiological activity: This is a non-insertion sham using a placebo acupuncture device. It seems an adequate control. While unlikely, it is possible that the true acupoints may be stimulated by the sham needle placement 2-3 cm from the true acupoints	Don't know
Shi 2010	Adequate	Adequate	Adequate	Adequate	Adequate	Adequate	No sham/placebo intervention	Don't know
Sun 2011	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo controls	Don't know
Xiong 2008a	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention group.	Don't know
Xue 2009	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention	Don't know
Zeng 2010	Adequate	Adequate	Adequate	Adequate	Adequate	Don't know	No sham/placebo intervention	Don't know

(Continued)

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Footnotes

* Two independent acupuncturists assessed acupuncture as adequate in terms of the choice of acupuncture points, number of sessions, needling technique and experience of acupuncturists for all trials.

† 10 sessions of treatment may be ok. However, treatment once per week does not seem frequent enough.

‡ Names of acupoints were not provided.

§ "Physiotherapist with Level I acupuncture accreditation" does not seems highly experienced.

^{††} Although the total number of treatments (10 treatments) is adequate, the frequency of the treatment is probably not adequate (over 3 months).

^{‡‡} Use of two different styles of acupuncture treatment is problematic. It is a weakness of the study design.

Appendix 4. Adverse events in the acupuncture and control groups*

Study	Assessment Meth- ods for Adverse Events	Minor Adverse Events per Group, n/n†		Assessment Methods for – Side Effects	Comments‡	
		Acupuncture	Control	of Acupunc- ture		
An 2010	NR	0	NR	NR	"No adverse events were found during the treatment"	
Anastasi 2009	The study facilita- tor (SF) was trained and responsible for structured elic- itation of adverse events	0	0	NR	"No adverse events were reported"	
Chen 2009	NR	0	0	NR	"No adverse effects appeared in all participants"	
Chen 2011	NR	NR	NR	NR	NR	
Forbes 2005	NR	0	0	NR	"No direct adverse events were elicit- ed"	
Lembo 2009	NR	1/78	Sham, 2/75; Waitlist, 0/77	NR	"Three adverse events were report- ed during the acupuncture vs.sham acupuncture phase of the study All of these events were considered to be un- related to the study procedure"	
Li 2009	NR	NR	NR	NR	NR	
Liu 1997	NR	NR	NR	NR	NR	
Liu 2010	"The results of rou- tine blood, urine and stool test did not significantly change before and after treatment, which suggested there's no obvious	0	0	NR	"The results of routine blood, urine and stool test did not significantly change before and after treatment, which suggested there's no obvious toxic side effect in treatment group and control groups."	

Acupuncture for treatment of irritable bowel syndrome (Review)

(Continued)

(continued)	toxic side effect in treatment group and control groups"				
Lowe 2000	NR	NR	NR	NR	NR
Reynolds 2008	"An open text ques- tion at one and three months gathered adverse events. We also col- lected data on safe- ty and treatment process from logs completed by the acupuncturists"	NR	NR	NR	"No serious adverse events were re- ported. The acupuncturists report- ed some mild adverse effects of the acupuncture treatment (mild dizzi- ness during treatment and temporary worsening of symptoms followed by improvement). Some patients also re- ported mild adverse effects, but they all continued to attend for treatment"
Schneider 2006	NR	NR	NR	NR	NR
Shi 2010	NR	1/32	1/38	NR	"One participant in electro-acupunc- ture group withdrew due to acupunc- ture syncope and one participant in pinaverium bromide group withdrew due to itch and skin rash caused by al- lergic reaction. There's no obvious ad- verse effect in other participants"
Sun 2011	NR	0	0	NR	"No any adverse reaction occurred"
Xiong 2008a	NR	NR	NR	NR	NR
Xue 2009	NR	NR	NR	NR	NR
Zeng 2010	NR	NR	NR	NR	NR

Footnotes

* Definition of adverse events and side effects of acupuncture and methods of assessments varied among trials. NR = not reported; RCT = randomized controlled trial.

† For the only two RCTs (Lembo 2009; Shi 2010) that reported number of adverse events per group, numbers reported in this table are the numbers of patients in each group who had adverse events. The one serious adverse event observed in the acupuncture groups in Lembo 2009 was not interpreted as treatment-related. The one adverse event observed in the acupuncture group in Shi 2010 was syncope due to acupuncture.

[‡] The frequency, severity or duration of the minor side effects associated with the needling was not systematically reported in any RCT.

WHAT'S NEW

Date	Event	Description	
11 January 2012	New citation required and conclusions have changed	Substantively updated review with new conclusions and authors	
11 January 2012 New search has been performed		New literature searches conducted between November 15 and November 28, 2011. New studies added.	

Acupuncture for treatment of irritable bowel syndrome (Review)



CONTRIBUTIONS OF AUTHORS

Study concept and design: Eric Manheimer, L. Susan Wieland, Brian Berman, Ke Cheng, Lixing Lao, Xueyong Shen Data extraction: Eric Manheimer, Ke Cheng, Shih Min Li, L. Susan Wieland Acupuncture adequacy assessments: Lixing Lao, Shen Xueyong Analysis and interpretation of data: Eric Manheimer Drafting the manuscript: Eric Manheimer, with L. Susan Wieland contributing to the Methods section Critically revised manuscript for important intellectual content and provided final approval of the version to be published: Brian Berman, Ke Cheng, Lixing Lao, Shih Min Li, Eric Manheimer, Xueyong Shen, L. Susan Wieland Obtained funding: Brian Berman, Eric Manheimer

DECLARATIONS OF INTEREST

None known

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INDEX TERMS

Medical Subject Headings (MeSH)

Acupuncture Therapy [*methods]; Irritable Bowel Syndrome [*therapy]; Randomized Controlled Trials as Topic

MeSH check words

Humans