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Acupuncture and acupressure for premenstrual syndrome (Review)

Armour M, Ee CC, Hao J, Wilson TM, Yao SS, Smith CA

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TABLE OF CONTENTS

HEADER	1
ABSTRACT	1
PLAIN LANGUAGE SUMMARY	2
SUMMARY OF FINDINGS	4
BACKGROUND	9
OBJECTIVES	9
METHODS	9
Figure 1.	11
RESULTS	13
Figure 2.	15
Figure 3.	16
DISCUSSION	18
AUTHORS' CONCLUSIONS	19
ACKNOWLEDGEMENTS	19
REFERENCES	20
CHARACTERISTICS OF STUDIES	23
DATA AND ANALYSES	33
Analysis 1.1. Comparison 1 Acupuncture vs sham acupuncture, Outcome 1 Overall premenstrual symptoms- Mood	34
Analysis 1.2. Comparison 1 Acupuncture vs sham acupuncture, Outcome 2 Overall premenstrual symptoms - Physical	34
Analysis 1.3. Comparison 1 Acupuncture vs sham acupuncture, Outcome 3 Adverse events.	34
Analysis 1.4. Comparison 1 Acupuncture vs sham acupuncture, Outcome 4 Response rate.	35
Analysis 1.5. Comparison 1 Acupuncture vs sham acupuncture, Outcome 5 Quality of Life.	35
Analysis 2.1. Comparison 2 Acupuncture vs no treatment, Outcome 1 Overall premenstrual symptoms	36
Analysis 2.2. Comparison 2 Acupuncture vs no treatment, Outcome 2 Adverse events.	36
Analysis 3.1. Comparison 3 Acupressure vs sham acupressure, Outcome 1 Overall premenstrual symptoms.	36
Analysis 3.2. Comparison 3 Acupressure vs sham acupressure, Outcome 2 Quality of Life.	37
APPENDICES	37
WHAT'S NEW	44
HISTORY	44
CONTRIBUTIONS OF AUTHORS	44
DECLARATIONS OF INTEREST	44
SOURCES OF SUPPORT	45
DIFFERENCES BETWEEN PROTOCOL AND REVIEW	45
INDEX TERMS	45



[Intervention Review]

Acupuncture and acupressure for premenstrual syndrome

Mike Armour¹, Carolyn C Ee², Jie Hao³, Tanya Marie Wilson⁴, Sofia S Yao⁵, Caroline A Smith¹

¹NICM Health Research Institute, Western Sydney University, Penrith, Australia. ²National Institute of Complementary Medicine (NICM), Western Sydney University, Penrith, Australia. ³School of Science and Health, Western Sydney University, Sydney, Australia. ⁴Atticus Health, Hastings, Australia. ⁵School of Science and Health, University of Western Sydney, Campbelltown, Australia

Contact address: Mike Armour, NICM Health Research Institute, Western Sydney University, Building 5, Campbelltown Campus, Penrith, NSW, 2751, Australia. m.armour@westernsydney.edu.au, 17382885@student.uws.edu.au.

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ABSTRACT

Background

Acupuncture has a history of traditional use in China for women's health conditions including premenstrual syndrome (PMS), but its effectiveness for this condition remains unclear. This review examined the available evidence supporting the use of acupuncture or acupressure to treat PMS.

Objectives

To evaluate the effectiveness and safety of acupuncture or acupressure for women with PMS or premenstrual dysphoric disorder (PMDD).

Search methods

We searched the Cochrane Gynaecology and Fertility Specialised Register, Cochrane Central Register of Studies Online (CENTRAL CRSO), MEDLINE, Embase, AMED, PsycINFO, CINAHL (from inception to 21 September 2017), two clinical trial databases (from their inception to 21 September 2017), and four electronic databases in China (from their inception to 15 October 2017): Chinese Biomedical Literature database (CBM), China National Knowledge Infrastructure (CNKI), VIP information/ Chinese Scientific Journals database and WANFANG. Reference lists from included articles were handsearched.

Selection criteria

We included studies if they randomised women with PMS and associated disorders (PMDD and late luteal phase dysphoric disorder/ LPDD) to receive acupuncture or acupressure versus sham, usual care/waiting-list control or pharmaceutical interventions mentioned by the International Society for Premenstrual Disorders (ISPMD). If acupuncture or acupressure were combined with another therapy, these studies were also included where the additional therapy was the same in both groups. Cross-over studies were eligible for inclusion, but only data from the first phase could be used.

Data collection and analysis

Two review authors independently selected the studies, assessed eligible studies for risk of bias, and extracted data from each study. Study authors were contacted for missing information. The quality of the evidence was assessed using GRADE. Our primary outcomes were overall premenstrual symptoms and adverse events. Secondary outcomes included specific PMS symptoms, response rate and quality of life.

Main results

Five trials (277 women) were included in this review. No trials compared acupuncture or acupressure versus other active treatments. The number of treatment sessions ranged from seven to 28. The quality of the evidence ranged from low to very low quality, the main limitations being imprecision due to small sample sizes and risk of bias related to detection bias and selective reporting.

Acupuncture versus sham acupuncture

Acupuncture may provide a greater reduction in mood-related PMS symptoms (mean difference (MD) -9.03, 95% confidence interval (CI) -10.71 to -7.35, one randomised controlled trial (RCT), n = 67, low-quality evidence) and in physical PMS symptoms (MD -9.11, 95% CI -10.82 to -7.40, one RCT, n = 67, low-quality evidence) than sham acupuncture, as measured by the Daily Record of Severity of Problems scale (DRSP). The evidence suggests that if women have a mood score of 51.91 points with sham acupuncture, their score with acupuncture would be between 10.71 and 7.35 points lower and if women have a physical score of 46.11 points, their score with acupuncture would be between 10.82 and 7.4 points lower. There was insufficient evidence to determine whether there was any difference between the groups in the rate of adverse events (risk ratio (RR) 1.74, 95% CI 0.39 to 7.76, three RCTs, n = 167, $l^2 = 0\%$, very low-quality evidence).

Specific PMS symptoms were not reported

There may be little or no difference between the groups in response rates. Use of a fixed-effect model suggested a higher response rate in the acupuncture group than in the sham group (RR 2.59, 95% CI 1.71 to 3.92; participants = 100; studies = 2; $l^2 = 82\%$), but owing to the high heterogeneity we tested the effect of using a random-effects model, which provided no clear evidence of benefit for acupuncture (RR 4.22, 95% CI 0.45 to 39.88, two RCTs, n = 100, $l^2 = 82\%$, very low-quality evidence).

Acupuncture may improve quality of life (measured by the WHOQOL-BREF) compared to sham (MD 2.85, 95% CI 1.47 to 4.23, one RCT, n = 67, *l*ow-quality evidence).

Acupuncture versus no treatment

Due to the very low quality of the evidence, we are uncertain whether acupuncture reduces PMS symptoms compared to a no treatment control (MD -13.60, 95% CI -15.70 to -11.50, one RCT, n = 14).

No adverse events were reported in either group.

No data were available on specific PMS symptoms, response rate or quality of life outcomes.

Acupressure versus sham acupressure

We found low-quality evidence that acupressure may reduce the number of women with moderate to severe PMS symptoms at the end of the trial compared to sham acupressure (RR 0.64 95% CI 0.52 to 0.79, one RCT, n = 90, low-quality evidence). The evidence suggests that if 97 women out of 100 in the sham acupressure group had moderate to severe PMS symptoms, the number of women in the acupressure group with moderate to severe symptoms would be 50 to 76 women.

Acupressure may improve both physical (MD 24.3, 95% CI 17.18 to 31.42, one RCT, n = 90, low-quality evidence) and mental (MD 17.17, 95% CI 13.08 to 21.26, one RCT, n = 90, low-quality evidence) quality of life.

No data were available on adverse events, specific symptoms or response rates.

Authors' conclusions

The limited evidence available suggests that acupuncture and acupressure may improve both physical and psychological symptoms of PMS when compared to a sham control. There was insufficient evidence to determine whether there was a difference between the groups in rates of adverse events. There is no evidence comparing acupuncture or acupressure versus current ISPMD recommended treatments for PMS such as selective serotonin reuptake inhibitors (SSRIs). Further research is required, using validated outcome measures for PMS, adequate blinding and suitable comparator groups reflecting current best practice.

PLAIN LANGUAGE SUMMARY

Acupuncture and acupressure for premenstrual syndrome (PMS)

Review question

Cochrane authors reviewed the evidence on the effectiveness and safety of acupuncture or acupressure in women with premenstrual syndrome (PMS) or premenstrual dysphoric disorder.

Background

We wanted to know whether using acupuncture or acupressure therapy was better than receiving sham acupuncture, no treatment or currently recommended pharmaceutical medications for PMS such as serotonin reuptake inhibitors (SSRIs - a type of anti-depressant).

Study characteristics

Acupuncture and acupressure for premenstrual syndrome (Review)

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We found five randomised controlled trials including 277 women that examined the effect of acupuncture or acupressure in women with PMS. The evidence is current to September 2017. We also ran a smaller search in November 2017 to see if we had missed any recently published studies. Three trials compared acupuncture with sham acupuncture, one compared acupuncture with no treatment and one compared acupressure with sham acupressure.

Key results

Acupuncture may reduce overall mood and physical PMS symptoms when compared to sham. Acupressure may reduce the number of women having moderate to severe PMS symptoms when compared to sham acupressure. There was not enough evidence to determine the safety of acupuncture or acupressure.

Quality of the evidence

The quality of the evidence ranged from low to very low. The main limitations were imprecision due to small sample sizes (too few women in the study) and risk of bias related to blinding (where researchers or participants knew what treatment they were getting).

SUMMARY OF FINDINGS

Summary of findings for the main comparison. Acupuncture compared to sham for premenstrual syndrome

Acupuncture compared to sham acupuncture for premenstrual syndrome

Patient or population: women with PMS

Setting: outpatient clinic or community

Intervention: acupuncture

Comparison: sham acupuncture

Outcomes Anticipated absolute effects (95% CI)		olute effects [*]	(95% CI) pan	№ of partici- pants (studies)	Certainty of the evidence (GRADE)	Comments
	Risk with sham acupuncture	Risk with Acupuncture			· ·	
Overall premenstrual symptoms - Mood at two month follow-up. As- sessed with: Daily Record of Sever- ity of Problems (DRSP) scale. Scale range for DRSP (mood) 0-60. Lower scores indicate less severe premenstrual symptoms	The mean over- all premenstru- al symptoms- Mood was 51.91	MD 9.03 lower (10.71 lower to 7.35 lower)	-	67 (1 RCT)	⊕⊕⊝⊝ LOW ¹	Acupuncture may reduce the num- ber of mood PMS related symp- toms women have, compared to sham acupuncture.
Overall premenstrual symptoms - Physical at two month follow-up.As- sessed with: DRSP. Scale range for DRSP (physical) 0-66. Lower scores indicate less severe premenstrual symptoms	The mean over- all premenstru- al symptoms - Physical was 46.11	MD 9.11 lower (10.82 lower to 7.40 lower)	-	67 (1 RCT)	⊕⊕⊝⊝ LOW 1	Acupuncture may reduce the num- ber of physical PMS related symp- toms women have, compared to sham acupuncture.
Adverse events at up to 2 months' follow-up	24 per 1000	42 per 1000 (10 to 189)	RR 1.74 (0.39 to 7.76)	167 (3 RCTs)	⊕ooo VERY LOW ²³	There was insufficient evidence to determine whether there was any difference between the groups in the rate of adverse events.
Specific PMS symptoms - not mea- sured	-	-	-	-	-	No studies reported on this out- come
Response rate at end of treatment	327 per 1000	1000 per 1000 (147 to 1000)	RR 4.22 (0.45 to 39.88)	100 (2 RCTs)	⊕⊝⊝⊝ VERY LOW ⁴⁵⁶	There was insufficient evidence to determine whether there was

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						any difference in response rate be- tween the groups.
Quality of Life assessed with: WHOQOL-BREF.	The mean qual- ity of life score	MD 2.85 higher (1.47 higher to	-	67 (1 RCT)	⊕⊕⊝⊝ LOW ¹	Acupuncture may improve quality of life, compared to sham acupunc-
Scale range 0-100.	was 73.8	4.23 higher)				ture.
Higher scores indicate greater quali- ty of life.						
*The risk in the intervention group (its 95% CI).	and its 95% confider	nce interval) is base	d on the assumed r	isk in the compar	son group and the	relative effect of the intervention (and
CI: Confidence interval; MD: mean diff	erence; RCT: randor	nised controlled tri	al; RR: risk ratio			
Moderate certainty: We are moderate substantially different Low certainty: Our confidence in the						
Very low certainty: We have very little Downgraded two levels due to very se Downgraded one level due to serious eporting.	rious imprecision: sr risk of bias: unclear	nall sample size (67 risk of blinding of	' participants) both participants a	nd assessors in o	ne of the three tria	ls. This may have affected adverse event
Very low certainty: We have very little Downgraded two levels due to very se Downgraded one level due to serious eporting. Downgraded two levels due to very se Downgraded two levels due to very se	rious imprecision: sr risk of bias: unclear rious imprecision: sr rious imprecision: sr	nall sample size (67 risk of blinding of nall sample size (16 nall sample size (10	7 participants) both participants a 67 participants) and 10 participants)	nd assessors in o rare events. 95%	ne of the three tria	ls. This may have affected adverse event covers both benefit and harm
Very low certainty: We have very little Downgraded two levels due to very se Downgraded one level due to serious eporting. Downgraded two levels due to very se Downgraded two levels due to very se Downgraded one level due to serious r	rious imprecision: sr risk of bias: unclear rious imprecision: sr rious imprecision: sr isk of bias: unclear ri	nall sample size (67 risk of blinding of nall sample size (16 nall sample size (10 sk of blinding of bo	7 participants) both participants a 67 participants) and 10 participants) th participants and	nd assessors in o rare events. 95% assessors in one c	ne of the three tria confidence interval f the two included t	ls. This may have affected adverse event
Very low certainty: We have very little Downgraded two levels due to very se Downgraded one level due to serious eporting. Downgraded two levels due to very se Downgraded two levels due to very se Downgraded one level due to serious r	rious imprecision: sr risk of bias: unclear rious imprecision: sr rious imprecision: sr isk of bias: unclear ri	nall sample size (67 risk of blinding of nall sample size (16 nall sample size (10 sk of blinding of bo	7 participants) both participants a 67 participants) and 10 participants) th participants and	nd assessors in o rare events. 95% assessors in one c	ne of the three tria confidence interval f the two included t	ls. This may have affected adverse event covers both benefit and harm
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Very low certainty: We have very little Downgraded two levels due to very se Downgraded one level due to serious preporting. Downgraded two levels due to very se Downgraded two levels due to very se Downgraded one level due to serious r Downgraded one level due to serious i Summary of findings 2. Acupunct	rious imprecision: sr risk of bias: unclear rious imprecision: sr rious imprecision: sr isk of bias: unclear ri nconsistency: signifi ture compared to	nall sample size (67 risk of blinding of nall sample size (16 nall sample size (10 sk of blinding of bo cant heterogeneity no treatment fo	7 participants) both participants a 67 participants) and 00 participants) th participants and f (1 ² = 82%) due to di	nd assessors in o rare events. 95% assessors in one c ifferent definition:	ne of the three tria confidence interval f the two included t	ls. This may have affected adverse event covers both benefit and harm

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	Risk with no treatment	Risk with Acupuncture				
Overall premenstrual symptoms at end of treatment. Assessed with the Menstrual Symptom Severity List (MSSL).	The mean over- all premenstru- al symptom score was 17.54	MD 13.60 lower (15.70 lower to 11.50 lower)	-	14 (1 RCT)	⊕⊝⊝⊝ VERY LOW ¹²	Due to the very low quality of the evidence, we are uncertain whether acupuncture reduces PMS symptoms compared to a no treat-
Scale range 0-33.						ment control
Lower scores indicate less severe symptoms.						
Adverse events at end of treatment	0 per 1000	0 per 1000 (0 to 0)	not estimable	20 (1 RCT)	⊕⊙⊝⊝ VERY LOW ¹³	Neither group reported any ad- verse events
Specific PMS symptoms - not mea- sured	-	-	-	-	-	No studies reported on this out- come
Response rate - not measured	-	-	-	-	-	No studies reported on this out- come
Quality of Life - not measured	-	-	-	-	-	No studies reported on this out- come

*The risk in the intervention group (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; MD: mean difference

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

¹ Downgraded one level due to serious risk of bias: unclear risk of bias in randomisation, high risk of bias in blinding of participants which is likely to affect patient reported outcome. High level of missing data.

² Downgraded two levels due to very serious imprecision: very small sample size (14 participants)

³ Downgraded two levels due to very serious imprecision: very small sample size (14 participants) for rare events.

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Summary of findings 3. Acupressure compared to sham acupressure for premenstrual syndrome

Acupressure compared to sham acupressure for premenstrual syndrome

Patient or population: women with PMS

Setting: community

Intervention: acupressure

Comparison: sham acupressure

Outcomes	Anticipated abso (95% CI)	lute effects*	Relative effect (95% CI)	№ of partici- pants (studies)	Certainty of the evidence (GRADE)	Comments
	Risk with sham acupressure	Risk with Acu- pressure				
Overall premenstrual symptoms at end of treatment. Assessed with: number of women reporting moderate or severe PMS symptoms at the end of the intervention. Lower scores indicate more women with mild PMS or no PMS symptoms.	97 per 100	62 per 100 (50 to 76)	RR 0.64 (0.52 to 0.79)	90 (1 RCT)	⊕⊕⊝⊝ LOW ¹	Acupressure may reduce the number of women with mod- erate to severe PMS symp- toms at the end of the trial compared to sham acupres- sure
Adverse events - not measured	-	-	-	-	-	No studies reported on this outcome
Specific PMS symptoms - not measured	-	-	-	-	-	No studies reported on this outcome
Response rate - not measured	-	-	-	-	-	No studies reported on this outcome
Quality of Life - Physical at end of treat- ment. Assessed with: SF-12 Scale range 0-100 Higher scores indicate greater quality of life	The mean qual- ity of Life - Physical score was 67.2	MD 24.30 higher (17.18 higher to 31.42 higher)	-	90 (1 RCT)	⊕⊕⊝⊝ LOW ¹	Acupressure may improve physical quality of life com- pared to sham acupressure.
Quality of Life - Mental at end of treat- ment. Assessed with: SF-12 Scale range 0-100 Higher scores indicate greater quality of life	The mean qual- ity of Life - Men- tal score was 52.34	MD 17.17 higher (13.08 higher to 21.26 higher)	-	90 (1 RCT)	⊕⊕⊙⊙ LOW 1	Acupressure may improve mental quality of life com- pared to sham acupressure.

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*The risk in the intervention group (and its 95% confidence interval) is based on the assumed risk in the comparison group and the relative effect of the intervention (and

CI: Confidence interval; RR: Risk ratio; MD: mean difference

GRADE Working Group grades of evidence

High certainty: We are very confident that the true effect lies close to that of the estimate of the effect

Moderate certainty: We are moderately confident in the effect estimate: The true effect is likely to be close to the estimate of the effect, but there is a possibility that it is substantially different

Low certainty: Our confidence in the effect estimate is limited: The true effect may be substantially different from the estimate of the effect

Very low certainty: We have very little confidence in the effect estimate: The true effect is likely to be substantially different from the estimate of effect

¹ Downgraded two levels due to very serious imprecision: small sample size (90 participants)

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its 95% CI).





BACKGROUND

Description of the condition

Premenstrual syndrome (PMS) is a clinical syndrome resulting in symptoms that occur in relation to the menstrual cycle, and can lead to interference with women's lives. Symptoms usually begin five to 11 days before menstruation, and cease when menstruation starts, or soon after (Backstorm 1991; Dickerson 2004). PMS is common, with 95% of women reported to have one or more premenstrual symptoms during their reproductive life (Hylan 1999). Symptoms are often mild, but in some women can be severe enough to substantially affect daily activities. About 5% to 8% of women thus suffer from severe PMS; most of these women also meet criteria for the associated condition of premenstrual dysphoric disorder (PMDD) (Kimberly 2008). There is evidence that raised levels of inflammatory markers may play a role in PMS (Bertone-Johnson 2014; Gold 2016).The aetiology of PMS is still unclear.

More than 200 premenstrual symptoms have been recorded and can be classified into three broad categories:

- 1. behavioural symptoms including fatigue, insomnia, dizziness, changes in sexual interest, food cravings or overeating;
- psychological symptoms including irritability, anger, depressed mood, crying and tearfulness, anxiety, tension, mood swings, lack of concentration, confusion, forgetfulness, restlessness, loneliness, decreased self-esteem, tension;
- 3. physical symptoms including headaches, breast tenderness and swelling, back pain, abdominal pain and bloating, weight gain, swelling of extremities, water retention, nausea, muscle and joint pain (Dickerson 2004; Reid 1986).

PMDD, previously known as late luteal phase dysphoric disorder (LLPDD), is a more severe form of PMS (O'Brien 2011). PMDD is characterised by intense emotional symptoms occurring between ovulation and menstruation. Symptoms include severe depression, irritability and/or mood swings which interfere with relationships, social functioning, and work or school (Bancroft 1993; Medem 2004). According to the Diagnostic and Statistical Manual of Mental Disorders (DSM-V) the diagnosis of PMDD requires (1) the presence of at least five luteal-phase symptoms (panel), at least one of which must be a mood symptom (i.e. depressed mood, anxiety or tension, affect lability, or persistent anger and irritability); (2) the timing of symptoms confirmed by two cycles of daily charting; and (3) evidence of functional impairment. Finally, symptoms must not be the exacerbation of another psychiatric condition (APA 2013).The American College of Obstetrics and Gynecology (ACOG) criteria include the presence of at least one psychological or physical symptom that causes significant impairment and confirmed by means of prospective ratings (ACOG 2001).

Description of the intervention

Acupuncture is a Chinese medical treatment that has been used for more than 2000 years in China and Japan (White 2004). Acupuncture involves the insertion of fine metal needles into the skin and underlying tissues at precise points on the body. The needle can be left alone, or can be stimulated by twisting of the needle in various ways, or stimulated by electricity (electro-acupuncture). Acupressure stimulates the same points as acupuncture using manual pressure, usually with the finger or thumb, on these points rather than the insertion of a needle (Beal 1999). Acupressure is able to be self-delivered by the person themselves, while acupuncture is usually delivered by a trained practitioner. However, acupuncture is considered to give a greater 'dose' of treatment due to the insertion and manipulation of the needle (Armour 2016), rather than pressure alone. Acupuncture treatment is composed of needling aspects (choice of points and needling techniques), specific components relating to the style of diagnosis and treatment used, and generic needling components not specific to acupuncture such as belief, time, and attention given to the patient (Langevin 2011).

How the intervention might work

The mechanism of acupuncture remains unclear although potential mechanisms have been identified from research, and it is likely that a number of concurrent pathways may contribute to the overall therapeutic effect. Acupuncture can modulate endogenous opioids (Mayor 2013), and needling of specific acupuncture points such as SP6 may alter blood flow to the uterus and modulate prostaglandin levels (Armour 2016). Recent research has shown that acupuncture acts on a variety of inflammatory markers (McDonald 2013; McDonald 2015). Because women with PMS have elevated levels of inflammatory markers (Gold 2016), this may be a primary pathway through which acupuncture may reduce PMS symptoms.

Why it is important to do this review

Many interventions have been used to treat PMS, such as selective serotonin reuptake inhibitors (SSRIs) and hormonal interventions. Current guidelines for the treatment of PMS and PMDD from the International Society for Premenstrual Disorders (ISPMD) recommend SSRI medication as a first line of treatment (Ismaili 2016). SSRI treatments are effective for many women with PMS however the side effects are frequent and can lead to discontinuation (Marjoribanks 2013). Acupuncture may be an effective non-pharmacological therapeutic option with a much lower rate of adverse events than SSRI medication (Witt 2009).

OBJECTIVES

To evaluate the effectiveness and safety of acupuncture or acupressure for women with premenstrual syndrome (PMS) or premenstrual dysphoric disorder (PMDD).

METHODS

Criteria for considering studies for this review

Types of studies

We included randomised controlled trials without restriction on language and publication types. Cross-over studies were eligible for inclusion, but we planned to include only data from the first phase. We excluded non-randomised studies (for example studies with evidence of inadequate sequence generation such as alternate days, patient numbers).

Types of participants

Inclusion

We included studies of women of any age who met the diagnostic criteria for premenstrual syndrome (PMS) or premenstrual



dysphoric disorder (PMDD). Diagnosis of PMS or PMDD had to be made prior to trial inclusion by a healthcare professional such as a general practitioner using an established diagnostic tool such as the DSM-IV (Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition) and ICD-10 (International Statistical Classification of Diseases and Related Health Problems 10th Revision), or a validated symptom screening tool.

Exclusion

We excluded studies where there was only a self-diagnosis of PMS.

Types of interventions

The included intervention group methods were body acupuncture, electro-acupuncture, scalp acupuncture, ear acupuncture, acupressure, electromagnetic acupuncture, pyonex, intradermal needling, or moxibustion. However, we excluded acupoint injection. If acupuncture or acupressure were combined with another valid therapy, we included these studies if the additional therapy was the same in both groups such as acupuncture plus selective serotonin reuptake inhibitors (SSRIs) versus SSRI alone.

Control group methods were sham acupuncture or sham acupressure, no treatment, or any conventional treatments recommended by the International Society for Premenstrual Disorders (ISPMD), including SSRIs, oral contraceptives, gonadotropin-releasing hormone agonists, danazol or estradiol.

We excluded studies which compared two different forms of acupuncture, or compared acupuncture with a herbal medicine or a conventional treatment without proven efficacy.

Types of outcome measures

Primary outcomes

- 1. Overall premenstrual symptoms, measured using a validated prospective screening tool (such as the Premenstrual Symptom Screening Tool (PSST) or Daily Record of Severity of Problems (DRST)), or by pre-defined medical diagnostic criteria. These may be reported as an overall score or by separate sub scales such as mental and physical symptoms.
- 2. Adverse events (all adverse events, specific adverse effects, withdrawals for adverse effects).

Secondary outcomes

- 1. Specific symptoms of PMS: behavioural, psychological, physical.
- 2. Response rate (according to how response to treatment was defined in individual studies).
- 3. Quality of life, measured by a validated scale such as the Short form (SF) SF-36 or SF-12.

Search methods for identification of studies

We conducted a comprehensive and exhaustive search strategy to identify all relevant studies regardless of language or publication status (published, unpublished, in press, and in progress). The search was conducted in consultation with the Gynaecology and Fertility Group Information Specialist. See Appendix 1, Appendix 2; Appendix 3; Appendix 4; Appendix 5; Appendix 6; Appendix 7 for search strategies.

Electronic searches

We searched the Cochrane Gynaecology and Fertility Specialised Register, the Cochrane Central Register of Studies (CENTRAL CRSO), MEDLINE, AMED, Embase, PsycINFO, CINAHL (from inception to 21 September 2017) and four electronic databases in China (from inception to 15 Oct 2017): Chinese Biomedical Literature database (CBM), China National Knowledge Infrastructure (CNKI), VIP information/Chinese Scientific Journals database and WANFANG database. An additional search on 2 November 2017 was undertaken using PubMed and Google Scholar to identify any recent trials not yet included in the previous databases.

Searching other resources

Other sources of trials included:

- 1. Trial registers for ongoing and registered trials: http:// www.clinicaltrials.gov and WHO International Clinical Trials Registry Platform (apps.who.int/trialsearch/).
- 2. Reference lists from included articles and previous systematic reviews.
- 3. Additional electronic resources were searched in November 2017:
 - DARE Database of Abstracts of Reviews of Effects (reference lists from non Cochrane reviews on similar topics);
 - b. ProQuest Dissertations & Theses for unpublished dissertations and theses;
 - c. Conference abstracts on the Web of Science;
 - d. OpenGrey for unpublished literature from Europe http:// www.opengrey.eu/;
 - e. LILACS database for the Portuguese and Spanish speaking world http://regional.bvsalud.org/php/index.php?lang=en.

Additional electronic resources were searched in English.

Data collection and analysis

Selection of studies

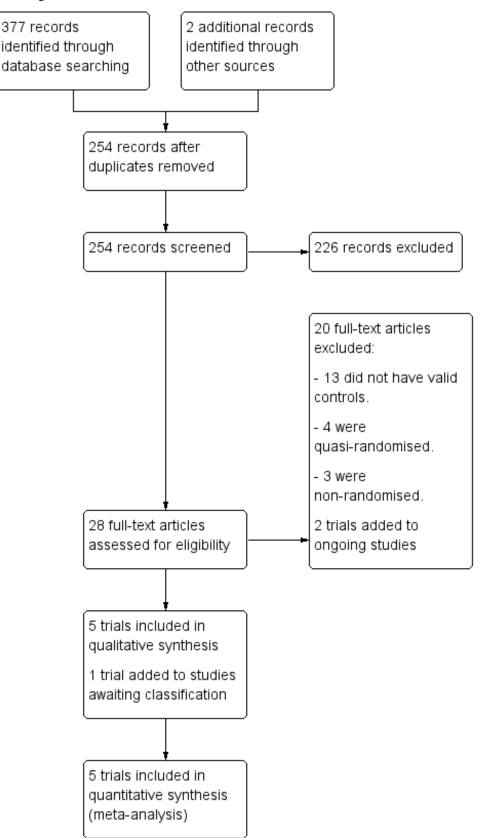
Two review authors (MA and CE) screened the titles and abstracts of articles found in the search, and discarded trials that were clearly not eligible. One review author (JH) searched for and selected the trials from the Chinese databases.

One review author (JH) translated Chinese papers. MA, CE and JH independently assessed whether trials met the inclusion criteria, with disagreements resolved by discussion. If articles contained insufficient information to make a decision about eligibility, MA or CE attempted to contact authors of the original reports to obtain further details. If details of randomisation were unclear in the reporting, we contacted all trial authors to ascertain if the study was truly randomised. We made a first contact and then sent a reminder one month later.

The selection process was documented with a "PRISMA" flow chart (Figure 1).



Figure 1. Study flow diagram.





Data extraction and management

Following an assessment for inclusion MA, CS, CE, TW and JH independently extracted data. We resolved discrepancies by discussion. For each included trial we extracted data regarding the location of the trial, the methods of the trial (as per assessment of risk of bias), the participants (age range, eligibility criteria), the nature of the interventions, and data relating to the outcomes specified above. We collected information on reported benefits and adverse effects. We extracted data and entered them onto a form sourced from the *Cochrane Handbook for Systematic Reviews of Interventions* (Higgins 2011). MA checked and entered data into Review Manager 5.3 (RevMan 2014).

Assessment of risk of bias in included studies

Two review authors (from MA, CS, CE, TW and JH) independently assessed risks of bias for each trial, using the criteria described in the *Cochrane Handbook* (Higgins 2011). The tool consists of seven items, with three potential responses: 'yes', 'no', and 'unclear'. In all cases a judgement of 'yes' indicates a low risk of bias and a judgement of 'no' indicates a high risk of bias. If insufficient detail was reported our judgement was usually 'unclear'. We also made a judgement of 'unclear' if we knew what happened in the study but the risk of bias was unknown to us, or if an entry was not relevant to the study at hand (particularly for assessing blinding and incomplete outcome data, or when the outcome being assessed by the entry had not been measured in the study).

We assessed the following characteristics: sequence generation, allocation concealment, blinding (or masking) of participants, blinding (or masking) of outcome assessors, incomplete data assessment, selective outcome reporting, and other sources of bias. We resolved disagreements that arose at any stage by discussion between the review authors or with a third party, when necessary. We generated a 'Risk of bias' assessment table for each study. We assessed other aspects of trial quality including the extent of blinding (if appropriate), whether groups were comparable at baseline, the extent of losses to followup, non-compliance and whether the outcome assessment was standardised according to the *Cochrane Handbook* (Higgins 2011). This information is presented in the Characteristics of included studies.

Measures of treatment effect

We performed statistical analysis in accordance with the *Cochrane Handbook* (Higgins 2011). We performed statistical analysis using Review Manager 5 software (RevMan 2014). For dichotomous data, we expressed results for each study as summary risk ratios (RRs) with corresponding 95% confidence intervals (CIs), using the Mantel-Haenszel method. We expressed continuous data as mean differences (MDs) with 95% CIs, or as standardised mean differences (SMDs) if outcomes were conceptually the same but measured in different ways in the different trials. A standard rule of thumb for interpreting effect sizes is that 0.2 represents a small effect, 0.5 a moderate effect and 0.8 a large effect.

Unit of analysis issues

The primary analysis was per woman randomised. We included trials with multiple arms and describe them in the Characteristics of included studies, for example, two different types of acupressure compared with sham acupressure. If there were two acupuncture or

acupressure groups, we combined data from both treatment arms into one group. For studies with a sham control and no treatment control group, we divided the shared intervention evenly between groups as described in the *Cochrane Handbook* (Higgins 2011). Where outcomes were repeated measures, we undertook analysis of outcomes at the end of the intervention.

Dealing with missing data

We analysed data on an intention-to-treat basis, as far as possible. We did not impute missing data but we did report the proportion lost to follow-up and analysed only the available data. In future updates, if more eligible studies are included, the impact of including studies with high levels of missing data in the overall assessment of treatment effect will be explored by using sensitivity analysis.

Assessment of heterogeneity

We measured inconsistency across trials in the meta-analysis using the I^2 statistic. This describes the percentage of total variation across studies that is due to heterogeneity rather than chance (Higgins 2003; Higgins 2011). The interpretation of the I^2 statistic is as follows:

- 1. 10% to 40% might not be important;
- 2. 30% to 60% may represent moderate heterogeneity;
- 3. 50% to 90% may represent substantial heterogeneity;
- 4. 75% to 100% considerable heterogeneity.

We regarded heterogeneity as substantial if an I^2 was greater than 50% and either the Tau² was greater than zero, or there was a low P value (less than 0.10) in the Chi² test for heterogeneity. If we identified substantial heterogeneity (above 50%), we used a random-effects model in the meta-analysis.

Assessment of reporting biases

We planned to investigate potential biases of publication using the funnel plot or other analytical methods (Egger 1997). If there were 10 or more studies in the same analysis we would have investigated reporting biases (such as publication bias) using funnel plots. If visual assessment had suggested asymmetry, we would have explored possible reasons.

Data synthesis

We carried out statistical analysis using the Review Manager software (RevMan 2014). We used fixed-effect meta-analysis for combining data where it was reasonable to assume that studies were estimating the same underlying treatment effect: i.e. where trials were examining the same intervention, and the trials' populations and methods were judged sufficiently similar, or if the number of studies was small (less than three).

If there was clinical heterogeneity sufficient to expect that the underlying treatment effects differed between trials, or if substantial statistical heterogeneity was detected, we used random-effects meta-analysis to produce an overall summary, if an average treatment effect across trials was considered clinically meaningful. The random-effects summary was treated as the average range of possible treatment effects and we discussed the clinical implications of treatment effects differing between trials. If the average treatment effect was not clinically meaningful, we did not combine trials. Where we used random-effects analyses, the

results were presented as the average treatment effect with 95% CIs, and the estimates of Tau^2 and $\mathsf{I}^2.$

An increase in the odds of a particular outcome, which may be beneficial (e.g. reduction in symptoms) or detrimental (e.g. adverse effects), will be displayed graphically in the meta-analyses to the right of the centre-line and a decrease in the odds of an outcome to the left of the centre-line.

Subgroup analysis and investigation of heterogeneity

Had we identified substantial heterogeneity, we planned to investigate it using sensitivity analyses and to consider whether an overall summary was meaningful, and if it was, to undertake a random-effects analysis.

Sensitivity analysis

We planned to conduct sensitivity analyses for the primary outcomes to determine whether the conclusions were robust to arbitrary decisions made regarding the eligibility and analysis. These analyses included consideration of whether the review conclusions would have differed if eligibility had been restricted to studies at low risk of bias (defined as studies rated as being at low risk of bias with respect to sequence generation and allocation concealment, and not rated as at high risk of bias in any of the domains assessed).

We did not perform any sensitivity analyses because each subgroup analysis did not include sufficient studies.

Overall quality of the body of evidence: 'Summary of findings' tables

We used the GRADE approach as outlined in the GRADE handbook in order to assess the quality of the body of evidence. We prepared a 'Summary of findings' table using GRADEpro (GRADEpro GDT 2015) and Cochrane methods. This table outlines the overall quality of the body of evidence for the main review outcomes (Overall premenstrual symptoms, adverse events, specific PMS symptoms, response rates and quality of life) for the main review comparison (acupuncture versus sham). We also generated 'Summary of findings' tables for the comparisons 'acupuncture versus no treatment'; and 'acupressure versus sham acupressure' for these same outcomes.

The GRADE approach uses five considerations (study limitations, consistency of effect, imprecision, indirectness and publication bias) to assess the quality of the body of evidence for each outcome. The evidence can be downgraded from 'high quality' by one level for serious (or by two levels for very serious) limitations, depending on assessments for risk of bias, indirectness of evidence, serious inconsistency, imprecision of effect estimates or potential publication bias. Judgements were justified, documented, and incorporated into reporting of results for each outcome.

Assessment of the quality of the acupuncture intervention delivered

We assessed the quality of the acupuncture or acupressure intervention in published journal articles using the National Institute for Complementary Medicine Acupuncture Network (NICMAN) scale (Smith 2017). The NICMAN scale was developed to assess the quality of the acupuncture delivered in a clinical trial, and is designed to be used in combination with the STandards for Reporting Interventions in Clinical Trials of Acupuncture (STRICTA) reporting guidelines (MacPherson 2010). This scale assesses and scores the following: the rationale behind the point selection, the qualifications and experience of the practitioner delivering the intervention, the adequacy of the number of treatment sessions, and the reporting of vital components of the acupuncture practice itself such as needle depth, diameter and stimulation. Higher scores indicate improved quality of the acupuncture intervention delivered, but there is no cut-off for a 'high' versus 'low' quality score on the NICMAN scale. Each trial was independently scored out of 23 by two review authors (from CS, JH, CE, TW, YY). Any discrepancies were resolved by discussion. If this was not reached a third review author (MA) decided on the final score.

RESULTS

Description of studies

Results of the search

The search retrieved 379 articles. Twenty-eight studies were potentially eligible and were retrieved in full text. Five studies met our inclusion criteria. Twenty studies were excluded. One study was placed in awaiting classification due to an inability to contact the study authors. Two studies are ongoing. See study tables: Characteristics of included studies; Characteristics of excluded studies; Characteristics of studies awaiting classification; Characteristics of ongoing studies

Included studies

Study design and setting

Two hundred and seventy-seven women with premenstrual syndrome (PMS)were included in this review. One trial was from Croatia (Habek 2002), two from China (Yu 2006; Zhang 2017), one from Korea (Shin 2009) and one from Iraq (Bazarganipour 2017). One trial (Bazarganipour 2017) reported funding support.

Two trials were undertaken in the community (Bazarganipour 2017; Shin 2009), two trials in an outpatient hospital setting (Yu 2006; Zhang 2017), and in one trial the setting was unclear (Habek 2002).

All studies were randomised controlled trials (RCTs). Two studies had a parallel two-arm design in which participants were randomised either to acupuncture or sham acupuncture group (Habek 2002; Yu 2006). Three studies had a three-arm design (Bazarganipour 2017; Shin 2009; Zhang 2017). For two of the three armed trials, only the appropriate arms of verum acupuncture and sham or waiting-list control were included in the analysis (Shin 2009; Zhang 2017). One trial included two different acupressure groups, one using the acupuncture point LI4 and one using LR3 (Bazarganipour 2017). These were combined into one active group as specified in Unit of analysis issues.

Participants

Women in two trials were aged from 18 to 45 years (Bazarganipour 2017;Yu 2006), one from 18 to 40 years (Zhang 2017),one from 20 to 35 years (Shin 2009); the other only reported the mean ages of the two groups were both about 30 years (Habek 2002). Sample sizes ranged from 20 (Shin 2009) to 90 (Bazarganipour 2017).

DSM-IV was used as diagnostic criteria in two trials (Yu 2006; Zhang 2017), ICD-10 was used in one trial (Shin 2009), PSST scores in one

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trial (Bazarganipour 2017) and a patient history form was used in the other trial (Habek 2002).

Women were screened for baseline PMS symptoms for three menstrual cycles in two trials (Shin 2009; Yu 2006), two cycles in one trial (Habek 2002), and one cycle in two trials (Bazarganipour 2017; Zhang 2017).

Interventions and comparators

Three trials reported the effect between acupuncture and sham acupuncture (Habek 2002; Yu 2006; Zhang 2017), and one between acupuncture and no treatment control (Shin 2009). One of these trials (Habek 2002), combined acupuncture with auriculo-acupuncture, which is acupuncture administered to points on the pinna of the ear, while another administered Korean hand acupuncture (Shin 2009). One trial compared acupressure and sham acupressure (Bazarganipour 2017). The type of sham acupuncture used was superficial needling on 'non-meridian' or 'non-acupuncture' points in two trials (Habek 2002; Yu 2006) and a non-penetrative sham in one trial (Zhang 2017). Sham acupressure consisted of acupressure being applied to an area unrelated to PMS symptoms (Bazarganipour 2017).

Treatment frequency and total number of sessions

Acupuncture 'dose' consists of neurophysiological and cumulative dose (Armour 2016). Neurophysiological components include needle retention time, number of acupuncture points used, location of acupuncture points and type of stimulation used (manual versus electro-acupuncture). Cumulative dose components include the frequency and total number of treatments.

The treatment frequency, timing of treatment and total number of treatments used in the included studies were very heterogenous. The number of treatments ranged from seven (Habek 2002) to 28 (Bazarganipour 2017) in total. Treatments were given throughout the menstrual cycle in two studies (Shin 2009; Zhang 2017), in the luteal phase only in two studies (Bazarganipour 2017; Habek 2002), and in the last seven days of the cycle in another study (Yu 2006). Treatment frequency varied from daily (Bazarganipour 2017), to three times per week (Zhang 2017).

The number of acupuncture points varied between studies

Duration of treatment sessions

Session durations and needle retention ranged from 15 minutes (Shin 2009) to 30 minutes (Habek 2002; Yu 2006; Zhang 2017).

Outcomes

All trials except Zhang 2017 collected their outcomes at the end of the intervention. Zhang 2017 had a one- and two-month follow-up. Data from the two-month follow-up were used.

Overall premenstrual symptoms

Daily symptom rating charts were used in three studies (Shin 2009, Yu 2006; Zhang 2017). The Premenstrual Daily Symptom Diary (PTSD), a four-point scale from 'no symptom' to 'severe symptoms' on 19 items for 31 days was used in one trial (Yu 2006). It is unclear if this was a validated outcome measure. The sum of the symptoms score change between pretreatment and the third treatment month was used, 'mean rank' but not 'mean and standard deviation' was reported as the outcome, so the data

were not included in the meta-analysis (Yu 2006). The Menstrual Symptom Severity List, a five-point scale rating symptoms from 'not present' to 'extreme' on 33 items (Mitchell 1992) was used in one trial (Shin 2009). The treatment effects were defined as the differences between the first pretreatment luteal phase (five days prior to menses) mean score and the first post-treatment luteal phase (five days prior to menses) mean score, but only end scores were reported, so we used end scores as the outcome in this review (Shin 2009). One study (Bazarganipour 2017) used the Premenstrual Symptom Screening Tool (PSST) scores as their outcome (Steiner 2003). PSST has 19 questions in two parts, 14 questions related to mood, body, and behaviour and five questions based on the potential impact of these symptoms on an individual's life. These data were presented as the number of women reporting no or mild PMS symptoms compared to those reporting moderate to severe at the end of the intervention. The number of women reporting moderate to severe PMS at the end of the review was presented in the analysis. One study (Zhang 2017), used a Daily Record of Severity of Problems scale (DRSP), a validated measure of PMS symptoms (Endicott 2006), and provided separate scores for mood and physical symptom components. These were reported separately in the meta-analysis.

Adverse events

Adverse events were reported as an outcome in all acupuncture trials but not in the acupressure trial (Bazarganipour 2017). Specific adverse events reported included haematomas, pain at the site of needling and itching after needling.

Specific symptoms of PMS: psychological, physical and functional symptoms, irritability.

No studies reported on this outcome.

Response rate

Two trials reported effectiveness rate. Treatments were considered to be successful if there were no further symptoms of PMS, if acupuncture and medication treatment of PMS became unnecessary or if PMS symptoms did not occur for at least a year after the acupuncture therapy (Habek 2002). Acupuncture was considered to be effective if the improvement rate of the scores on PTSD was greater than or equal to 30% (Yu 2006).

Quality of life

Two trials reported quality of life (QoL) outcomes. One study (Bazarganipour 2017) used the SF12 and one study (Zhang 2017) used the WHO Quality of Life-BREF scale (WHOQOL-BREF). The SF-12 is a 12-item questionnaire used to assess generic health outcomes from the patient's perspective (Ware 1996). It is based on a subset of questions from the larger SF-36 and covers the same eight domains of health outcomes, including physical functioning (PF), role-physical (RP), bodily pain (BP), general health (GH), vitality (VT), social functioning (SF), role-emotional (RE), and mental health (MH). The WHOQOL-BREF instrument comprises 26 items, which measure the following broad domains: physical health, psychological health, social relationships, and environment (Whoqol 1998).



Excluded studies

Twenty studies were excluded, see Characteristics of excluded studies;

Three studies were excluded because they were not randomised controlled trials (RCTs) (Li 2004; Li 2008; Wang 2003).

Four studies were excluded because they were quasi-RCTs (Carvalho 2013; Hong 2002; Sun 2004; Jiang 2005).

Four studies were excluded because they used combined comprehensive treatment methods as control group (Xu 2006a, Xu 2006b, Xu 2011; Zhang 1994).

Two studies were excluded because they used two acupuncture methods as control group (Gu 2008; Zhao 2012).

Two studies were excluded because they used traditional Chinese medicine as control group (Hu 2010, Li 2014).

Five studies were excluded because they compared acupuncture with western medicine but that medicine was not mentioned in Types of interventions (Cao 2013; Chen 2011; Koleini 2017; Sun 2008; Zheng 2001).

Risk of bias in included studies

See Figure 2 and Figure 3. One study (Zhang 2017) was at a low risk of bias in all domains.

Figure 2. Risk of bias graph: review authors' judgments about each risk of bias item presented as percentages across all included studies.

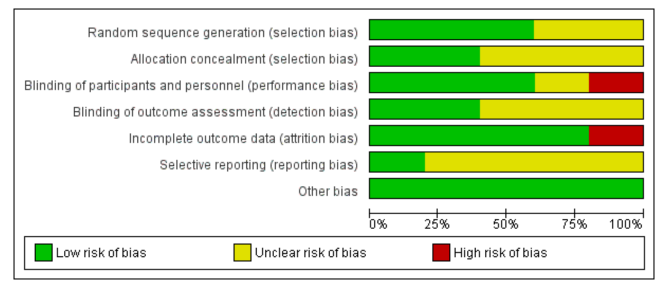
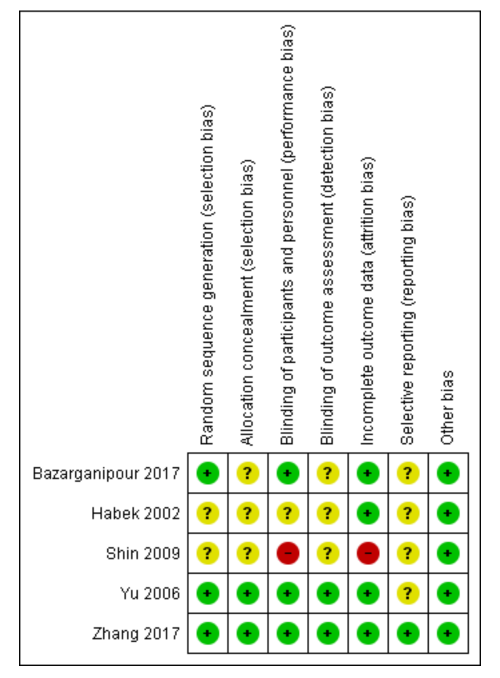




Figure 3. Risk of bias summary: review authors' judgments about each risk of bias item for each included study.



Allocation

Random sequence generation

The method of random sequence generation was associated with a low risk of bias in three trials (Bazarganipour 2017; Yu 2006; Zhang 2017). The two remaining trials were rated as unclear risk of bias as they did not report the methods of randomisation (Habek 2002; Shin 2009).

Allocation concealment

Two trials were rated as at low risk of bias related to allocation concealment (Yu 2006; Zhang 2017), while the other three were

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rated as unclear due to lack of clear reporting (Bazarganipour 2017; Habek 2002; Shin 2009).

Blinding

Sham-acupuncture was used in three trials (Habek 2002; Yu 2006; Zhang 2017). Blinding of participants was not tested in one trial (Habek 2002) and was rated as unclear. Blinding remained intact for two trials (Yu 2006; Zhang 2017), which were rated as low risk of bias. Sham acupressure was used in one trial (Bazarganipour 2017), blinding was not tested but it is unlikely that due to the pressure used that blinding would have been unsuccessful and was rated as low risk of bias. One trial using a no treatment control did not attempt to blind (Shin 2009) and was rated as high risk of bias. Cochrane Library

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Detection bias was rated as low risk of bias in two trials (Yu 2006; Zhang 2017), where the clinicians assessing the outcomes were blinded to group allocation. Blinding was rated as unclear in the other three trials (Bazarganipour 2017; Habek 2002; Shin 2009), where the blinding status of the assessors and analysts was not reported.

Incomplete outcome data

Four trials had missing data for less than 20% of participants and were rated as at low risk of attrition bias (Bazarganipour 2017; Habek 2002; Yu 2006; Zhang 2017). One trial had data missing for over 20% of participants and was rated as at high risk of attrition bias (Shin 2009).

Selective reporting

Four trials were rated as at unclear risk of selective reporting as there was no published protocol (Bazarganipour 2017; Habek 2002; Shin 2009; Yu 2006). One trial (Zhang 2017) was rated as at low risk of bias: this was a PhD thesis and included a comprehensive protocol in the methods section.

Other potential sources of bias

No other potential source of bias was identified in any study, and all studies were rated as at low risk for this domain.

NICMAN scale scores

Three studies were analysed and NICMAN scale scores assessed (Bazarganipour 2017; Habek 2002; Shin 2009). It was not practical to perform this assessment on an entire Masters or PhD thesis so no score was generated for Yu 2006 or Zhang 2017. Scores (out of 23) ranged from 14 (Habek 2002), 16 (Bazarganipour 2017) to 19 (Shin 2009). Overall, the design of the study, including the population, intervention, comparison and outcomes was considered adequate in all three studies, however there was poor reporting on the rationale behind the acupuncture point selection and the qualifications of the therapist performing the acupuncture in both Bazarganipour 2017 and Shin 2009.

Effects of interventions

See: Summary of findings for the main comparison Acupuncture compared to sham for premenstrual syndrome; Summary of findings 2 Acupuncture compared to no treatment for premenstrual syndrome; Summary of findings 3 Acupressure compared to sham acupressure for premenstrual syndrome

1 Acupuncture versus sham

One trial used 'mean rank' but not 'mean and standard deviation' as the outcome for 'Overall premenstrual symptoms', so was not included in this meta-analysis (Yu 2006).

Primary outcomes

1.1 Overall premenstrual symptoms - Mood

(Analysis 1.1)

One trial (Zhang 2017) reported on this outcome.

Acupuncture may provide a greater reduction in mood-related premenstrual syndrome (PMS) symptoms than sham acupuncture (mean difference (MD) -9.03, 95% confidence interval (CI) -10.71

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to -7.35, one RCT, n = 67, low-quality evidence), as measured by the Daily Record of Severity of Problems scale (DRSP). The evidence suggests that if women have a mood score of 51.91 points with sham acupuncture, their score with acupuncture would be between 10.71 and 7.35 points lower.

1.2 Overall premenstrual symptoms - Physical

(Analysis 1.2)

One trial (Zhang 2017) reported on this outcome. Acupuncture may provide a greater reduction in physical PMS symptoms than sham acupuncture (MD -9.11, 95% CI -10.82 to -7.40, one RCT, n = 67, lowquality evidence), as measured by the DRSP. The evidence suggests that if women have physical score of 46.11 points their score with acupuncture would be between 10.82 and 7.4 points lower.

1.3 Adverse events

(Analysis 1.3)

Three trials reported the safety of acupuncture in this comparison. Overall, there was insufficient evidence to determine if there was a difference in the adverse event rate between acupuncture and sham acupuncture (RR 1.74, 95% CI 0.39 to 7.76, three RCTs, n = 167, $I^2 = 0\%$, very low-quality evidence)

Secondary outcomes

1.4 Response rate

(Analysis 1.4)

Two trials (Habek 2002; Yu 2006) reported response rate for acupuncture versus sham acupuncture

Individually, both studies found a benefit in the intervention group, and pooling of their data supported this finding (RR 2.59, 95% CI 1.71 to 3.92; participants = 100; studies = 2; $I^2 = 82\%$). However, due to the high heterogeneity we tested the effect of using a random-effects model, which provided no clear evidence of benefit for acupuncture (RR 4.22, 95% CI 0.45 to 39.88, two RCTs, n = 100, $I^2 = 82\%$, very low-quality evidence). The high heterogeneity is most likely due to very heterogenous definitions of what was defined as a 'response rate' between the two studies as well as clinical heterogeneity in the intervention delivered. We concluded that owing to the very low quality of the evidence, there may be little or no difference between the groups for this outcome.

1.5 Quality of life

(Analysis 1.5)

One trial reported quality of life outcomes using the WHOQOL-BREF (Zhang 2017). Acupuncture may improve quality of life compared to sham acupuncture, measured at the end of the intervention (MD 2.85, 95% CI 1.47 to 4.23, one RCT, n = 67, low-quality evidence).

1.6 Specific symptoms of PMS

No study reported this outcome.

2 Acupuncture versus no treatment

Primary outcomes

2.1. Overall premenstrual symptoms

(Analysis 2.1)

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Acupuncture and acupressure for premenstrual syndrome (Review)



One trial reported overall premenstrual symptoms using the Menstrual Symptom Severity List (MSSL), and reported end scores for this outcome (Shin 2009). Due to the very low quality of the evidence, we are uncertain whether acupuncture reduces PMS symptoms compared to a no treatment control (MD -13.60, 95% CI -15.70 to -11.50, one RCT, n = 14, very low-quality evidence)

2.2. Adverse events

(Analysis 2.2)

One trial reported the safety of acupuncture. There were no adverse events reported in either group (Shin 2009).

Secondary outcomes

2.3 Specific symptoms of PMS

No study reported this outcome.

2.4 Response rate

No study reported this outcome.

3 Acupressure versus sham acupressure

Primary outcomes

3.1 Overall premenstrual symptoms

(Analysis 3.1)

One trial reported overall PMS (Bazarganipour 2017). There was evidence that acupressure may reduce the number of women with moderate to severe PMS symptoms at the end of the trial compared to sham acupressure (RR 0.64 95% CI 0.52 to 0.79, one RCT, n = 90, low-quality evidence). The evidence suggests that if 97 women out of 100 in the sham acupressure group had moderate to severe PMS symptoms, the number of women in the acupressure group with moderate to severe symptoms would be 50 to 76 women.

3.2. Adverse events

No study reported this outcome.

Secondary outcomes

3.3 Specific symptoms of PMS

No study reported this outcome.

3.4 Response rate

No study reported this outcome.

3.5 Quality of life

(Analysis 3.2)

One trial reported quality of life outcomes using the SF-12 scale (Bazarganipour 2017). The mental and physical subscale components were reported separately. There was evidence that acupressure may improve both physical (MD 24.30, 95% CI 17.18 to 31.42, one RCT, n = 90, low-quality evidence) and mental (MD 17.17, 95% CI 13.08 to 21.26, one RCT, n = 90, low-quality evidence) quality of life, compared to sham acupressure.

DISCUSSION

Summary of main results

Evidence from these five trials including 277 women found that both acupuncture and acupressure may reduce overall premenstrual syndrome (PMS) symptoms. Acupuncture may reduce the overall severity of PMS symptoms as well as the number of both mood-related and physical PMS symptoms at the end of the intervention. There was insufficient evidence to determine whether there is a difference in rates of adverse events between acupuncture and sham acupuncture. For the single acupressure trial there was a reduction in the number of women suffering from moderate or severe PMS at the end of the intervention. There are a small number of trials included within each comparison, and this limits the power of the review to detect any meaningful differences between groups.Therefore, while there may be benefits for acupuncture and acupressure, these benefits should be interpreted with caution until larger studies are undertaken.

Overall completeness and applicability of evidence

There were a small number of trials included in this review, and the outcome measures for PMS symptoms were heterogenous, limiting the ability to make meaningful meta-analysis. The completeness and applicability of the evidence is limited due to the small number of trials, with relatively small sample sizes. The reporting on specific safety outcomes was limited in some trials, either due to a lack of reporting or due to very small sample sizes making it hard to detect these relatively rare events. All trials compared acupuncture with either sham or with no-treatment. No trial compared acupuncture or acupressure with the current International Society for Premenstrual Disorders (ISPMD) recommended treatments.

Trials varied significantly in the frequency and total number of treatments given as well as the acupuncture points chosen and the timing of the intervention relative to menses, which may reflect differences in 'best practice' between different countries. This difference in both 'dosage' and timing around menses can potentially affect the outcome in menstrual conditions (Armour 2016), however due to the small number of trials a subgroup analysis on this factor could not be undertaken.

Quality of the evidence

The 'Risk of bias' tables (Figure 2; Figure 3) show that acupuncture and acupressure studies on PMS have not generally been undertaken to a high methodological standard. Only one trial was at a low risk of bias across all domains. The majority of studies had low risk of bias in randomisation and unclear risk of bias in allocation concealment, blinding of outcome assessors and selective reporting. In cases where it was unclear, we contacted the authors to provide additional details, however there was no response.

None of the trials' acupuncture interventions were rated highly on the National Institute for Complementary Medicine Acupuncture Network (NICMAN) scale, with issues in reporting the rationale for point selection being common. Acupuncture point selection, based on either biomedical or traditional understanding, depending on the framework chosen, should be used to guide point selection for trials, and without a clear rationale for point choice it is difficult to determine if studies are using treatments that reflect clinical practice.

Acupuncture and acupressure for premenstrual syndrome (Review)

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Blinding of participants was at low risk of bias in three studies, however none of these reported assessments of blinding success between groups. The lack of blinding of outcome assessors, or in the case of participant-reported outcomes, of participants themselves, may influence outcomes, and therefore, were rated as high risk of bias. Due to the small number of studies in each comparison means that while individual studies have had positive results, there is currently insufficient evidence for a consistent treatment effect.

The GRADE quality of evidence was often downgraded due to imprecision, mostly related to small sample sizes, and risk of bias particularly related to participant blinding. The overall quality of evidence for GRADE was moderate to very low for PMS symptoms and very low for adverse events (Summary of findings for the main comparison; Summary of findings 2; Summary of findings 3).

Potential biases in the review process

We attempted to minimise bias during the review process. Two review authors assessed the eligibility of studies, carried out data extraction and assessed the risk of bias. We are aware that some literature on acupuncture and acupressure may not be published in mainstream journals and therefore maybe excluded from the main databases. Our search was comprehensive and we included studies identified in languages other than English, however we did not systematically search all other language databases, for example Korean language only databases, therefore we cannot rule out the possibility that some studies may have been missed.

Agreements and disagreements with other studies or reviews

There has been only one systematic review on this topic performed in the past five years (Jang 2014). That review included eight acupuncture studies and included four trials that did not meet the inclusion criteria for this review due to the choice of control interventions (Gu 2008; Hong 2002; Xu 2006a; Xu 2006b). Overall, the authors found that physical symptoms such as breast tenderness improved while psychological symptoms did not. Our review included several more recent trials that were not included in Jang 2014 and found benefits for acupuncture for both physical and psychological symptoms.

AUTHORS' CONCLUSIONS

Implications for practice

The limited evidence available suggests that acupuncture and acupressure may improve both physical and psychological

symptoms of premenstrual syndrome (PMS) when compared to a sham control. There was insufficient evidence to determine whether there was a difference between the groups in rates of adverse events.There is no evidence comparing acupuncture or acupressure versus current International Society for Premenstrual Disorders (ISPMD) recommended treatments for PMS, such as selective serotonin reuptake inhibitors (SSRIs). Further research is required, using validated outcome measures for PMS, adequate blinding and suitable comparator groups reflecting current best practice.

Implications for research

Very few studies included validated outcome measures for PMS and none compared acupuncture or acupressure with a currently accepted pharmaceutical or psychological intervention for PMS. We suggest future research include three-armed trials comparing acupuncture or acupressure with a suitable sham and with an active pharmaceutical or psychological intervention suggested by current clinical guidelines. Given the issues with blinding a three-armed trial, having blinded clinician-rated scales as well as subjective patient-centred outcomes should be considered. There is a need to improve the quality and reporting in future trials, especially with regards to the choice of acupuncture point selection and the rationale behind the choice of point(s) should be clear. Future trials need to improve reporting on the qualifications and experience of the practitioner delivering the acupuncture or acupressure intervention, including years of training, qualification and current registration status.

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REFERENCES

References to studies included in this review

Bazarganipour 2017 {published data only}

Bazarganipour F, Taghavi SA, Allan H, Beheshti F, Khalili A, Miri F, et al. The effect of applying pressure to the LIV3 and LI4 on the symptoms of premenstrual syndrome: a randomized clinical trial. *Complementary Therapies in Medicine* 2017;**31**:65-70. [PUBMED: 28434473]

Habek 2002 {published data only}

Habek D, Habek JC, Barbir A. Using acupuncture to treat premenstrual syndrome. *Archives of Gynecology and Obstetrics* 2002;**267**(1):23-6.

Shin 2009 {published data only}

Shin KR, Ha JY, Park HJ. The effect of hand acupuncture therapy and hand moxibustion therapy on premenstrual syndrome among Korean women. *Western Journal of Nursing Research* 2009;**31**(2):171-86.

Yu 2006 {published data only}

Yu J, Liu B, Liu Z. Clinical Observation Based on a Systematic Review of Efficacy and Safety of Acupuncture Treatment for Premenstrual Syndrome [master's thesis]. Beijing (China): China Academy of Chinese Medical Science, 2006:1-89.

Zhang 2017 {published data only}

Zhang GC. Clinical study on the treatment of acupuncture for premenstrual syndrome. Clinical Study on the Treatment of Acupuncture for Premenstrual Syndrome (PhD Thesis). Guang Zhou (China): Guang Zhou University of Chinese Medicine, 2017.

References to studies excluded from this review

Cao 2013 {published data only}

Cao L, Hou GY. Treatment of headache during menstruation with electroacupuncture and triple puncture. *Journal of Clinical Acupuncture and Moxibustion* 2013;**29**(4):21-3.

Carvalho 2013 {published data only}

Carvalho F, Weires K, Ebling M, Padilha Mde S, Ferrao YA, Vercelino R. Effects of acupuncture on the symptoms of anxiety and depression caused by premenstrual dysphoric disorder. *Acupuncture in Medicine* 2013;**31**(4):358-63.

Chen 2011 {published data only}

Chen L, Zhu W, Lin Y, Huang JP, Lin F, Yuan LP, et al. Clinical study of balance acupuncture and moxibustion for treatment of women with premenstrual headache. *Journal of Guangzhou University of Traditional Chinese Medicine* 2011;**28**(4):379-81.

Gu 2008 {published data only}

Gu AH. Observation on eight methods of intelligent turtle combined with point selection treatment based on syndrome differentiation for 30 cases with premenstrual tension syndrome. *Huna Journal of Traditional Chinese Medicine* 2008;**24**(5):30-1.

Hong 2002 {published data only}

* Hong YF. Clinical therapeutic effect of scalp acupuncture on premenstrual tension syndrome. *Chinese Acupuncture & Moxibustion* 2002;**22**(9):597-8.

Hong YF. The effect of scalp acupuncture treatment for premenstrual syndrome. *Shanghai Journal of Acupuncture and Moxibustion* 2002;**21**(3):24.

Hu 2010 {published data only}

Fang YQ, Wu J, Hu YP, Zhou R, Yang LJ, Shi Y, et al. Observation on the tranquilizing and liver-regulating acupuncture treatment for 47 PMS with liverqi stagnation type - randomised controlled study. *Guangming Journal of Chinese Medicine* 2008;**23**(9):1308-10.

* Hu YP, Zhou R. Observation on the tranquilizing and liverregulating acupuncture treatment for PMS with liver qi stagnation type. *Journal of Shanxi of Traditional Chinese Medicine* 2010;**31**(8):1047-50.

Wang J, Wu J. Observation on the tranquilizing and liverregulating acupuncture treatment for PMS with liverqi stagnation type. *Journal of Sichuan of Traditional Chinese Medicine* 2007;**25**(9):82-3.

Jiang 2005 {published data only}

Jiang W, Li Y, Sun J. Clinical study on treatment of premenstrual tension syndrome with auricular point sticking. *Chinese Acupuncture & Moxibustion* 2002;**22**(3):165-7.

* Jiang W, Shan QH. Clinical study on treatment of premenstrual tension syndrome with auricular pressure pellets. *International Journal of Clinical Acupuncture* 2005;**14**(1):27-33.

Koleini 2017 {published data only}

Koleini S, Valiani M. Comparing the effect of auriculotherapy and vitamin B6 on the symptoms of premenstrual syndrome among the students who lived in the dorm of Isfahan University of Medical Sciences. *Iranian Journal of Nursing and Midwifery Research* 2017;**22**(5):354-8. [PUBMED: 29033988]

Li 2004 {published data only}

Li YM, Yu L, Gong DF. Observation on treatment of 46 premenstrual tension syndrome cases with auricular point sticking. *Journal of Clinical Acupuncture and Moxibustion* 2004;**20**(12):45.

Li 2008 {published data only}

Li C. Acupuncture for 20 cases with premenstrual headache. *Journal of Practical Traditional Chinese Medicine* 2008;**24**(9):591.

Li 2014 {published data only}

Li L, Li JF, Chen WF, Wu YX, Huang H, Lin QM. Effect of auricular point sticking on menstrual distending pains of breasts. *Modern Clinical Nursing* 2014;**13**(2):40-2.

Sun 2004 {published data only}

Guo SY, Sun YZ. Clinical Study on the Treatment of Premenstrual Syndrome by the Back-shu and Front-mu and Network Points



Acupuncture [Master's thesis]. Harbin (China): Heilongjiang University of Chinese Medicine, 2004.

Guo SY, Sun YZ. Comparison between acupuncture and medication in treatment of premenstrual syndrome. *Shanghai Journal of Acupuncture and Moxibustion* 2004;**23**(1):5-6.

* Sun YZ, Guo SY. Comparison of therapeutic effects of acupuncture and medicine on premenstrual syndrome. *Chinese Acupuncture & Moxibustion* 2004;**24**(1):29-30.

Sun 2008 {published data only}

Sun ZH. Acupuncture for 42 premenstrual headache. *Jiangxi Journal of Traditional Chinese Medicine* 2008;**5**:50.

Wang 2003 {published data only}

Wang QQ, Chen HL. Ear electro-acupuncture for treatment of premenstrual syndrome in 49 cases. *Journal of Sichuan of Traditional Chinese Medicine* 2003;**21**(10):81-2.

Xu 2006a {published data only}

Xu YY. Clinical Study on the Treatment of Acupuncture of Backshu on Premenstrual Syndrome [Master's thesis]. Harbin: Hei Long Jiang University of Chinese Medicine, 2006.

Xu 2006b {published data only}

Xu YY, Sun YZ. Observation on the waist penetration needling therapy for premenstrual syndrome. *JCAM* 2006;**22**(5):37-8.

Xu 2011 {published data only}

Xu SL. Clinical study on treatment of premenstrual syndrome by acupuncture and moxibustion. *Journal of Acupuncture and Tuina Science* 2011;**9**(5):310-1.

Zhang 1994 {published data only}

Zhang WF, Bei YL, Zhang WZ. The effect of acupuncture in the treatment of 60 cases with premenstrual tension syndrome. *Chinese Acupuncture and Moxibustion* 1994;**Supplement**:100-1.

Zhao 2012 {published data only}

Li Z. Clinical observation on fire needle for menstrual headache due to phlegm stagnation in collaterals. *Shanghai Journal of Acupuncture and Moxibustion* 2012;**31**(3):145-6.

Zheng 2001 {published data only}

Zheng JY, Zheng JZ. The efficacy of acupuncture in the treatment of headache during menstruation: a report of 60 cases. *Hebei Journal of Traditional Chinese Medicine* 2001;**23**(8):612-3.

References to studies awaiting assessment

Kim 2005 {published data only}

Kim SC, Kim SN, Lim JA, Choi CM, Shim EK, Koo ST, et al. Effects of acupuncture treatment on the premenstrual syndrome: controlled clinical trial. *Journal of Korean Acupuncture and Moxibustion Society* 2005;**22**:41-60.

Kurebayashi 2013 {published data only}

Kurebayashi L. Auriculotherapy in the cares to the premenstrual syndrome. https://clinicaltrials.gov/ct2/show/NCT01782040.

References to ongoing studies

NCT02504515 {published data only}

Effect of homeopathy, acupuncture or anthroposophic medicine in women's quality of life. Ongoing study April 2015.

Additional references

ACOG 2001

ACOG. ACOG Practice Bulletin: premenstrual syndrome. International Journal of Gynecology & Obstetrics 2001;73:183-91.

APA 2013

American Psychiatric Association. Diagnostic and statistical manual of mental disorders-DSM-V. American Psychiatric Association. Washington DC: American Psychiatric Association, 2013.

Armour 2016

Armour M, Smith CA. Treating primary dysmenorrhoea with acupuncture: a narrative review of the relationship between acupuncture 'dose' and menstrual pain outcomes. *Acupuncture in Medicine : journal of the British Medical Acupuncture Society* 2016;**34**(6):416-24. [PUBMED: 27913451]

Backstorm 1991

Backstrom T, Hammarback S. Premenstrual syndromepsychiatric or gynaecological disorder. *Annals of Medicine* 1991;**23**(6):625-33.

Bancroft 1993

Bancroft J. The premenstrual syndrome -a reappraisal of the concept and the evidence. *Psychological Medicine* 1993;**Supplement 24**:1-47.

Beal 1999

Beal MW. Acupuncture and acupressure: applications to women's reproductive health care. *Journal of Nurse-Midwifery* 1999;**44**(3):217-30.

Bertone-Johnson 2014

Bertone-Johnson ER, Ronnenberg AG, Houghton SC, Nobles C, Zagarins SE, Takashima-Uebelhoer BB, et al. Association of inflammation markers with menstrual symptom severity and premenstrual syndrome in young women. *Human Reproduction* (*Oxford, England*) 2014;**29**(9):1987-94. [PUBMED: 25035435]

Dickerson 2004

Dickerson LM, Mazyck PJ, Hunter MH. Premenstrual syndrome. *American Family Physician* 2004;**57**(8):1743-52.

Egger 1997

Egger Matthias, Smith George Davey, Schneider Martin, Minder Christoph. Bias in meta-analysis detected by a simple, graphical test. *BMJ* 1997;**315**(7109):629.

Endicott 2006

Endicott J, Nee J, Harrison W. Daily Record of Severity of Problems (DRSP): reliability and validity. *Archives of Women's Mental Health* 2006;**9**(1):41-9. [PUBMED: 16172836]



Gold 2016

Gold EB, Wells C, Rasor MO. The association of inflammation with premenstrual symptoms. *Journal of Women's Health* 2016;**25**(9):865-74. [PUBMED: 27135720]

GRADEpro GDT 2015 [Computer program]

GRADE Working Group, McMaster University. GRADEpro GDT. Version accessed on 4th Febuary 2018. Hamilton (ON): GRADE Working Group, McMaster University, 2015.

Higgins 2011

Higgins JPT, Green S (editors). Cochrane Handbook for Systematic Reviews of Interventions Version 5.1.0 [updated March 2011]. The Cochrane Collaboration Available from www.cochrane-handbook.org.

Hylan 1999

Hylan TR, Sundell K, Judge R. The impact of premenstrual symptomatology on functioning and treatment-seeking behavior: experience from the United States, United Kingdom, and France. *Journal of Womens Health & Gender-Based Medicine* 1999;**8**(8):1043-52.

Ismaili 2016

Ismaili E, Walsh S, O'Brien PMS, Backstrom T, Brown C, Dennerstein L, et al. Fourth consensus of the International Society for Premenstrual Disorders (ISPMD): auditable standards for diagnosis and management of premenstrual disorder. *Archives of Women's Mental Health* 2016;**19**(6):953-8. [PUBMED: 27378473]

Jang 2014

Jang SH, Kim DI, Choi MS. Effects and treatment methods of acupuncture and herbal medicine for premenstrual syndrome/ premenstrual dysphoric disorder: systematic review. *BMC Complementary and Alternative Medicine* 2014;**14**:11.

Kimberly 2008

Yonkers KA, O'Brien PMS, Eriksson E. Premenstrual syndrome. *Lancet* 2008;**371**(9619):1200-10.

Langevin 2011

Langevin HM, Wayne PM, Macpherson H, Schnyer R, Milley RM, Napadow V, et al. Paradoxes in acupuncture research: strategies for moving forward. Evidence-Based Complementary and Alternative Medicine. England, 2011; Vol. 2011:180805.

MacPherson 2010

MacPherson H, Altman DG, Hammerschlag R, Li YP, WU TX, White A, et al. Revision of standards for reporting interventions in clinical trials of acupuncture. *Chinese Journal of Integrative Medicine* 2010;**8**(9):804-18.

Marjoribanks 2013

Marjoribanks Jane, Brown Julie, O'Brien Patrick Michael Shaughn, Wyatt Katrina. Selective serotonin reuptake inhibitors for premenstrual syndrome. *Cochrane Database of Systematic Reviews* 2013;**6**.

Mayor 2013

Mayor D. An exploratory review of the electroacupuncture literature: clinical applications and endorphin mechanisms. *Acupuncture in Medicine: Journal of the British Medical Acupuncture Society* 2013;**31**(4):409-15. [PUBMED: 23917395]

McDonald 2013

McDonald JL, Cripps AW, Smith PK, Smith CA, Xue CC, Golianu B. The anti-inflammatory effects of acupuncture and their relevance to allergic rhinitis: a narrative review and proposed model. *Evidence-based Complementary and Alternative Medicine: eCAM* 2013;**2013**:591796. [PUBMED: 23476696]

McDonald 2015

McDonald JL, Cripps AW, Smith PK. Mediators, receptors, and signalling pathways in the anti-inflammatory and antihyperalgesic effects of acupuncture. *Evidencebased Complementary and Alternative Medicine: eCAM* 2015;**2015**:975632. [PUBMED: 26339274]

Medem 2004

Medem Inc and American Psychiatric Association. Premenstrual dysphoric disorder (PMDD). http://www.medem.com/medlb/article_detaillb.cfm?article_ID=ZZZF9KNGTRC&sub_cat=2003 accessed at 25 July, 2004.

Mitchell 1992

Mitchell ES, Woods NF, Lentz MJ. Recognising PMS when you see it: criteria for PMS sample selection. In: Taylor DL, Woods NF editor(s). Menstruation, Health and Illness. New York (USA): Hemisphere, 1991:89-102.

O'Brien 2011

O'Brien PM, Backstrom T, Brown C, Dennerstein L, Endicott J, Epperson CN, et al. Towards a consensus on diagnostic criteria, measurement and trial design of the premenstrual disorders: the ISPMD Montreal consensus. *Archives of Women's Mental Health* 2011;**2**(14):13-21.

Reid 1986

Reid RL. Premenstrual syndrome: a time for introspection. *American Journal of Obstetrics & Gynecology* 1986;**155**(5):921-6.

RevMan 2014 [Computer program]

Nordic Cochrane Centre, The Cochrane Collaboration. Review Manager 5 (RevMan 5). Version 5.3. Copenhagen: Nordic Cochrane Centre, The Cochrane Collaboration, 2014.

Smith 2017

Smith CA, Zaslawski CJ, Cochrane S, Zhu X, Zheng Z, Loyeung B, et al. Reliability of the NICMAN Scale: an instrument to assess the quality of acupuncture administered in clinical trials. *Evidence-based Complementary and Alternative Medicine : eCAM* 2017;**2017**:5694083. [PUBMED: 28690661]

Steiner 2003

Steiner M, Macdougall M, Brown E. The premenstrual symptoms screening tool (PSST) for clinicians. *Archives of Women's Mental Health* 2003;**6**(3):203-9. [PUBMED: 12920618]



Ware 1996

Ware J, Kosinski M, Keller SD. A 12-Item Short-Form Health Survey: construction of scales and preliminary tests of reliability and validity. *Medical Care* 1996;**34**(3):220-33.

White 2004

White A, Ernst E. A brief history of acupuncture. Rheumatology. England, 2004; Vol. 43, issue 5:662-3.

Whoqol 1998

The Whoqol Group. Development of the World Health Organization WHO QoL_BREF quality of life assessment. *Psychological Medicine* 1998;**28**(3):551-8.

Witt 2009

Witt CM, Pach D, Brinkhaus B, Wruck K, Tag B, Mank S, et al. Safety of acupuncture: results of a prospective

CHARACTERISTICS OF STUDIES

Characteristics of included studies [ordered by study ID]

observational study with 229,230 patients and introduction of a medical information and consent form. *Forschende Komplementarmedizin (2006)* 2009;**16**(2):91-7. [PUBMED: 19420954]

References to other published versions of this review

Yu 2005

Yu J, Liu B, Liu Z, Welch V, Wu T, Clarke J, Smith CA. Acupuncture for premenstrual syndrome. *Cochrane Database of Systematic Reviews* 2005, Issue 2. [DOI: 10.1002/14651858.CD005290]

* Indicates the major publication for the study

Methods	Randomised, sham controlled, single-blind, parallel three-armed trial comparing acupressure vs sham acupressure.
Participants	Country: Iraq
	Site: Hormozgan University
	Participants: 97 university students with PMS
	Inclusion criteria: age 18 to 45 years, menstruation between 21 and 35 days,has moderate to severe PMS as scored by the PSST.
	Exclusion criteria: infectious skin diseases, lesions or dermatitis at the acupressure point locations, NSAID or other analgesics three hours prior to the intervention, severe depression or anxiety (based on HADS score), auditory or visual impairments, major external factors causing tension in the last six months (such as surgery or death in the family), heart disease, renal disorder, diabetes, asthma, thyroi conditions, respiratory disorders or genital diseases.
Interventions	Participants received two training sessions by a researcher (FB) who had been trained in acupressure techniques from a TCM specialist for 20 sessions over four months. Training of the students took place in student accomodation, and consisted of PMS pathology, treatment methods, and introducing acu- pressure. No description of whether self-acupressure was observed or checked after training, nor of compliance.
	In both groups, the first menstrual cycle was used as a baseline measure.
	Acupressure group 1 (n = 32)
	Acupuncture point LR3 was used in the second and third cycles. Participants were instructed to start applying pressure 14 days prior to the onset of menses.
	Acupressure group 2 (n = 32)
	Acupuncture point LI4 was used in the second and third cycles. Participants were instructed to start ap plying pressure 14 days prior to the onset of menses.
	Sham acupressure (n = 33)

Bazarganipour 2017 (Continue	d)				
	Sham point located at not in the line of the Liv	the dorsal surface of the foot between the third and the fourth toes which was /er meridian.			
	In all three groups, participants sat down for 10-15 minutes to adapt to room temperature. All windows and doors were closed and nobody was allowed to enter the room. Participants are described as being supine but it is not clear at what point they lay down. During the second and third cycle the acupres- sure was performed daily, and consisted of two minutes of pressure on each acupoint, then two min- utes of stimulation without pressure, until a light pain occurred in the acupoint, for 20 minutes in total.				
		cipant or a researcher applied the pressure during the second and third cycles, ation in the abstract vs body text. No mention of whether acupressure was given ly.			
		honed daily, in the five days leading up to menstruation, as a reminder of the and outcome collection.			
	In this review acupres sure.	ssure group 1 and group 2 were combined and compared to sham acupres-			
Outcomes	PSST questionnaire (ra	tes PMS severity), SF12 HRQOL, Hospital Anxiety and Depression Scale (HADS)			
Notes	NICMAN score: 16				
	Funding: by a grant fro	m Hormozgan University of Medical Science in Iran.			
	Conflict of Interest: not	reported			
	Date study was conduc	ted: not reported			
Risk of bias					
Bias	Authors' judgement	Support for judgement			
Random sequence genera- tion (selection bias)	Low risk	Quote: "Participants were randomly divided into the study groups using a ta- ble of random numbers" (pg 67)			
Allocation concealment (selection bias)	Unclear risk	Not reported			
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Quote: "Participants were blinded to their allocation to the intervention groups." (pg 67)			
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Patient-reported outcomes, therefore blinding of participants likely to ensure outcomes not affected. Unclear if statistician or analysts blinded, researcher teaching acupressure not blind to allocation, but this is unlikely to affect out- come			
Incomplete outcome data	Low risk	7 dropouts, similar drop-out rates between groups			
(attrition bias) All outcomes					
(attrition bias)	Unclear risk	Common outcomes reported, however no protocol published			



Habek 2002	
Methods	Randomised, sham-controlled, parallel two-arm trial
Participants	Country: Croatia Site: no details Recruitment: 35 women with PMS Inclusion: no details Exclusion: no details
Interventions	Screening: two cycles screening
	Intervention group : (n = 18), mean age = 30.6 ± 8.4 years. Traditional acupuncture plus auricu- lo-acupuncture. Traditional acupuncture points: GV20 (Baihui), bilateral LI 4 (Hegu), bilateral LR3 (Taichong), CV3 (Zhongji), CV4 (Guanyuan), CV6 (Qihai), bilateral PC6 (Neiguan), bilateral GB34 (Yan- glingquan), bilateral BL23 (Shenshu). Auriculo-acupuncture points: TF4 (auricular Shenmen). Physician (OB/GYN) inserted thin solid sterilised stainless steel needles until the de qi effect was obtained.
	Control group: (n = 17), mean age = 29.8 ± 7.3 years. The 17 women were treated with sham-superficial acupuncture without the de qi effect (inserting the same type of needles at points on the lateral thighs and arms that were not on the classically described meridians.
	Duration: 7 days (four sessions)
	Timing of administration: treatments were conducted during the luteal phase of the menstrual cycle, every second day (7 days) for a period of 30 minutes a day.
	Summary measures: 1. success rate; 2. safety.
Outcomes	1. Success rate: treatments were considered to be successful if PMS did not occur any more, if AP and medication treatment of PMS became unnecessary or if PMS symptoms did not occur for a year after the AP therapy.
	2. Safety: adverse events were recorded (haematoma)
Notes	NICMAN score: 14
	Funding: not reported
	Conflict of Interest: not reported
	Date study was conducted: not reported
Risk of bias	

Rice	Authonal independent	Connert for indeement
Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Unclear risk	Quote: "A randomized prospective, placebo-controlled trial included 35 women with PMS" (pg 24). Randomisation method not reported
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	Unclear risk	Not reported
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Not reported



Habek 2002 (Continued)

Incomplete outcome data (attrition bias) All outcomes	Low risk	No dropouts in either group
Selective reporting (re- porting bias)	Unclear risk	No protocol published
Other bias	Low risk	None noted

Shin 2009

Methods	Randomised, non-blinded, parallel 2-arms trial						
Participants	Country: Korea Site: the laboratory of Ewha Woman's University College of Nursing Recruitment: 185 women were recruited through a posted advertisement placed in a university hospi- tal, of these 30 volunteers were enrolled and randomly divided into three groups (Hand Acupuncture Therapy, Hand Moxibustion Therapy, control group) of 10 each. There were respectively 7, 8, 7 volun- teers in each group completed. Hand acupuncture group versus control group were included in this review.						
	Inclusion: women aged 20 to 35 years meeting ICD-10 criteria for PMS. Exclusion: 1. menstrual irregularity (fewer than 20 days or more than 36 days) in the previous 3 months; 2. pregnancy or breastfeeding; hysterectomy; 3. oral contraceptive use in the previous 3 months; antidepressant, benzodiazepine, or psychotropic drug use; 4. diagnosed with thyroid disorder; and a history of depression or anxiety disorder.						
Interventions	Screening: three menstrual cycles were screened prior to intervention using the Menstrual Symptom Severity Scale and used as baseline data.						
	Intervention group: hand acupuncture therapy: (n = 10)						
	Treatments were applied to the basic female corresponding hyuls on the Im Ki Mek of both hands: A5, A6, A8, A12, A16, A18; Liver Ki Mek N18; and Spleen Ki Mek F6, Needles were inserted into either hand at an approximate penetration depth of less than 1 mm. After needle insertion, the women were instruct- ed to keep their hands still for 15 minutes.						
	Control group: no treatment control group. (n = 10)						
	Women did not receive any treatment but were contacted at the time of follow-up assessment.						
	Duration: 10 sessions for 4 weeks						
	Timing of administration: 10 times treatments in the 4-week period, once every 3 days at Ewha Univer- sity Medical Center.						
	Summary measures: score change of severity of menstrual symptoms between the first pretreatment luteal phase (5 days prior to menses) and the first post-treatment luteal phase (5 days prior to menses)						
Outcomes	The Menstrual Symptom Severity developed by Mitchell, Woods, and Lentz (1992)						
Notes	NICMAN score: 19						
	Funding: not reported						
	Conflict of Interest: not reported						
	Date study was conducted: not reported						

Acupuncture and acupressure for premenstrual syndrome (Review)

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Shin 2009 (Continued)

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera-	Unclear risk	Quote: "Women were randomly divided into three groups
tion (selection bias)		(i.e., HAT, HMT, control) of 10 each" (pg 175). Method of randomisation not re- ported.
Allocation concealment (selection bias)	Unclear risk	Not reported
Blinding of participants and personnel (perfor- mance bias) All outcomes	High risk	No blinding possible due to no treatment control group. Likely to affect out- come.
Blinding of outcome as- sessment (detection bias) All outcomes	Unclear risk	Not reported
Incomplete outcome data (attrition bias) All outcomes	High risk	14 women completed (7 in the HAT group, and 7 in the control group). During the intervention phase, 6 women withdrew for a variety of reasons (e.g., work schedule, personal plans, wedding, school, vacation, and so forth).
Selective reporting (re- porting bias)	Unclear risk	No protocol published
Other bias	Low risk	None noted

Yu 2006 Methods Randomised, single-blind, sham-controlled, parallel two-arm trial Participants Country: China Site: acupuncture clinic in a multi-speciality TCM hospital Recruitment: 73 women with PMS were recruited via advertisements and from referral from affiliated psychiatric and gynaecological clinics, of these 65 were randomised. 33 into intervention group (30 completed the trial), 32 into sham-acupuncture group (30 completed the trial). Inclusion: 1.women aged 18 to 45 years with regular menstrual cycles meeting DSM-IV criteria for PMS; 2. course of disease: 3 months to 15 years; 3. voluntarily signed the informed consent. Exclusion: 1. women with organic diseases, such as diabetes, anaemia, endometriosis, thyroid function low; 2. women with abnormal personality or chronic mental disorders, such as depression, anxiety, dysthymia, panic, etc.; 3. women in pregnancy, lactation, peri-menopause or prepare for pregnancy; 4. women with alcoholism or drug abuse history in the last 6 to 12 months; 5. women with oral contraceptives or hormone treatment history in the last 3 months. Interventions Screening: three menstrual cycles used to screen for PMS symptoms before intervention Intervention group: acupuncture group (n = 33) Mean age = 32.17 ± 8.49 years. Traditional acupuncture: DU 20 (Baihui), EX-HN3 (Yintang), EX-HN 3 (Taiyang), bilateral SP 6 (Sanyinjiao), bilateral SP 10 (Xuehai). Additional points according to different symptoms: add bilateral LR3 (Taichong), CV17 (Danzhong), bilateral LR 14 (Qimen) in PMS women with emotional instability, irritability or breast pain; add bilateral EX-CA1 (Zigong), CV4 (Guanyuan), bilateral SP9 (Yinlingquan) in PMS women with headache, limbs pain or abdominal pain; add bilateral ST36



Yu 2006 (Continued)									
	(Zusanli), CV6 (Qihai) in PMS women with fatigue or sleepiness; Add bilateral PC6 (Neiguan), bilater- al HT7 (Shenmen) in PMS women with insomnia or heart palpitations; add bilateral BL23 (Shenshu), GV4 (Mingmen), bilateral KI3 (Taixi) in PMS women with waist discomfort; different depth for different points; manual manipulation every 10 minutes, retention for 30 minutes. Disposable acupuncture nee- dles (0.30 mm x 40 mm, 0.30 mm x 25 mm were used.								
	Control group: sham-	Control group: sham-acupuncture group (n = 32)							
	Mean age = 31.07 ± 7.79 years. Sham acupuncture: superficial acupuncture at non-acupuncture points 1 cm away from the above points at 0.2 to 0.3 cun depth, no manipulation.								
	Duration: three cycles	Duration: three cycles (9 sessions).							
	Timing of administration	on: 3 treatments in the 7 days before menstruation for three menstrual cycles.							
	Summary measures: 1 response rate; 4. safety	. score change after 3 cycles; 2. change in days with discomfort after 3 cycles; 3. /.							
Outcomes	 Premenstrual Daily Symptom Diary, the symptoms include: irritability or tension, anger or sho per, anxiety or nervousness, depression or sadness, crying or tearfulness, relationship pro tiredness or lack of energy, insomnia, changes in sexual interest, food craving or overeating, dif concentrating, feeling overwhelmed, headaches, breast tenderness or swelling, back pain, abd pain, muscle and joint pain, weight gain, nausea. Days with discomfort: the total days with discomforts before menstruation. Response rate = the total scores before treatment - after treatment) / the total scores before treat 100%. 								
	Effective: > or = 30%, ineffective:< 30%.								
Notes	NICMAN score: not applicable								
	Funding: not applicable (masters thesis)								
	Conflict of Interest: not applicable (masters thesis)								
	Date study was conducted: not reported								
Risk of bias									
Bias	Authors' judgement	Support for judgement							
Random sequence genera- tion (selection bias)	Low risk	Central randomisation, using a computer random number generator, details unknown to investigators							
Allocation concealment (selection bias)	Low risk Clinical study of telephone voice response system, details unknown to investi- gators								
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Participants blinded							
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk Data management by the third party, details unknown to investigators								

Yu 2006 (Continued)

Selective reporting (re- porting bias)	Unclear risk	No protocol published
Other bias	Low risk	None noted

Methods	Randomised, sham controlled, parallel three-arm trial comparing acupuncture + intradermal needling vs non-channel superficial acupuncture + intradermal needling vs sham acupuncture and sham intra- dermal needling.						
Participants	Country: China						
	Site: participants were recruited from Hai Kou Hospital of Traditional Chinese Medicine and Hai Nan Provincial Hospital of Traditional Chinese Medicine in Hai Kou, China from January 2015 to June 2016.						
	Participants: 105 women aged 18-40 years, diagnosed with PMDD						
	Inclusion criteria						
	 Diagnosed with PMDD (Association American Psychiatric, Diagnostic and statistical manual of ment disorders: DSM-IV-TR: 4th Edition (2000). 						
	2. Age between 18 and 40 years.						
	3. Mentally fully conscious and willing to comply with the study.						
	4. Signed consent form.						
	5. Hamilton Anxiety Rating Scale (HAMA) score > 7.						
	6. Regular menstruation cycles (a period of 21-35 days per cycle).						
	Exclusion criteria						
	1. Age under 18 years old or above 40 years old.						
	Have received any pharmaceutical or non pharmaceutical treatment for PMDD within the latest months.						
	3. Have other medical conditions such as chronic heart failure, liver and renal failure, haematopoiet system diseases, psychosis and cancer.						
	4. Pregnancy or potential to fall pregnant, or lactating.						
	5. Poor evaluation value on compliance or fear of acupuncture.						
	6. Smoker or alcoholic for a long time.						
	7. Consent form not signed.						
Interventions	Experimental intervention: acupuncture (n = 35)						
	 Bilateral GV20(Baihui), EX-HN3 (Yintang); oblique needle insertion at a 30° angle to the skin of 0.5 cu depth. 						
	2. Bilateral SP6 (Sanyinjiao), KI6 (Zhaohai), LI4 (Hegu), and LR3 (Taichong); Perpendicular insertion o 0.5-0.8 cun depth.						
	 Bilateral CV12 (Zhongwan), CV10 (Xiawan), CV7(Qihai), RN4 (Guanyuan), EX-CA1 (Zigong), ST25 (Tian shu), Perpendicular insertion of 1.5 cun depth. Needle manipulation every 10 minutes 						
	Duration: 30 minutes each intervention, three times interventions a week, interval between two inter- ventions > 24 hours, four weeks as a course, two courses in total. Follow-up at after 2 months. Stop in- tervention during the first, second and third days of menstruation.						
	Intradermal needle therapy: embedding needles in the skin 3 mm to 5 mm at BL15 (Xinshu),BL18 (Gan- shu) and BL23 (Shenshu) for 2-3 days.						



Zhang 2017 (Continued)

Duration: interventions twice a week, interval between two interventions > 72 hours, four weeks as a course, two courses in total. Follow-up after 2 months. Stop intervention during the first, second and third days of menstruation.

Control/Comparison intervention

1. Non-acupoint superficial + intradermal needle control group 1 (n = 35)

- 1. Non-acupoint superficial:
 - a. Bilateral 1 cun close to GV20 (Baihui), EX-HN3(Yintang); oblique needle insertion at a 30° angle to the skin of 0.5 cm depth.
 - b. Bilateral 1 cun close to SP6 (Sanyinjiao), KI6 (Zhaohai), LI4 (Hegu), and LR3 (Taichong); Perpendicular insertion of 0.5 cm depth.
 - c. Bilateral 1 cun close to CV12 (Zhongwan), CV10 (Xiawan), CV7(Qihai), CV4 (Guanyuan), EX-CA1 (Zigong), ST25 (Tianshu); Perpendicular insertion of 0.5 cm depth.

Duration: 30 minutes each intervention, three times interventions a week, interval between two interventions > 24 hours, four weeks as a course, two courses in total. Follow-up at after 2 months. Stop intervention during the first, second and third days of menstruation.

2. Intradermal needle therapy: Embedding needles in the skin 3-5 mm at BL15 (Xinshu),BL18 (Ganshu) and BL23 (Shenshu) for 2 to3 days.

Duration: twice interventions a week, interval between two interventions > 72 hours, four weeks as a course, two courses in total. Follow at after 2 months. Stop intervention during the first, second and third days of menstruation.

2. Acupoint non-invasive acupuncture + non-invasive intradermal needle control group 2 (n = 35)

1. Acupoint non-invasive acupuncture: Non-invasive sham acupuncture on bilateral GV20 (Baihui), EX-HN3(Yintang); SP6 (Sanyinjiao), KI6 (Zhaohai), LI4 (Hegu), and LR3 (Taichong); CV12 (Zhongwan), CV10 (Xiawan), CV7 (Qihai), CV4 (Guanyuan), EX-CA1 (Zigong), ST25 (Tianshu), Needle manipulation every 10 minutes

Duration: 30 minutes each intervention, three times interventions a week, interval between two interventions > 24 hours, four weeks as a course, two courses in total. Follow at after 2 months. Stop intervention during the first, second and third days of menstruation.

2. Non-invasive sham intradermal needle at BL15 (Xinshu), BL18 (Ganshu) and BL23 (Shenshu) for 2-3 days.

Duration: interventions twice a week, interval between two interventions > 72 hours, four weeks as a course, two courses in total. Follow-up after 2 months. Stop intervention during the first, second and third days of menstruation.

In this review we included the comparison of acupuncture versus control group 2 (i.e. acupuncture versus acupoint non-invasive acupuncture + non-invasive intradermal needle)

Outcomes	Primary: Hamilton Anxiety Scale (HAMA)						
	Secondary: WHO Quality of Life-BREF scale (WHOQOL-BREF), Daily Record of Severity of Problems scale (DRSP)						
	Adverse events reported (haematoma, pain at needle site, and itching at needle site).						
	Endpoints: after one and two courses of intervention and at one and two months follow-up.						
Notes	NICMAN score: Not applicable						
	Funding: Not applicable (PhD Thesis)						
	Conflict of interest: not applicable (PhD Thesis)						

Acupuncture and acupressure for premenstrual syndrome (Review)



Zhang 2017 (Continued)

Date study was conducted: January 2015 to June 2016

Risk of bias

Bias	Authors' judgement	Support for judgement
Random sequence genera- tion (selection bias)	Low risk	Quote: "A computerized PMS3.1 software was used to generate a random number and simple randomisation was used (p28)"
Allocation concealment (selection bias)	Low risk	Quote: "Make grouped allocation results into cards. Put them into sealed kraft paper envelopes then sequentially numbered them. (p28)
Blinding of participants and personnel (perfor- mance bias) All outcomes	Low risk	Participants blinded to group allocation through the use of sham control group.
Blinding of outcome as- sessment (detection bias) All outcomes	Low risk	Quote: "People who participate in random allocation do not participate in in- tervention and data statistics. This study was designed to blind the personnel who assess outcome and analyse data" (pg 28).
Incomplete outcome data (attrition bias) All outcomes	Low risk	Dropouts balanced and unrelated to intervention group
Selective reporting (re- porting bias)	Low risk	All outcomes stated in the PhD thesis methods reported
Other bias	Low risk	No baseline imbalance or other evidence of bias

AP: acupuncture point DSM IV: Diagnostic and Statistical Manual of Mental Disorders, Fourth Edition RSP: Daily Record of Severity of Problems scale HAT: hand acupuncture therapy HMT: hand moxibustion therapy ICD-10: International Statistical Classification of Diseases and Related Health Problems 10th Revision NICMAN: National Institute for Complementary Medicine Acupuncture Network NSAID: non-steroidal anti-inflammatory drug PMDD: premenstrual dysphoric disorder PMS: premenstrual syndrome PSST: Premenstrual Symptoms Screening Tool SF12 HRQOL: Short Form 12 - Health Related Quality of Life. TCM: traditional Chinese medicine WHO: World Health Organization

Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion					
Cao 2013	Control treatment (Celebrex) did not met the inclusion criteria.					
Carvalho 2013	Quasi-randomised (assigned alternately).					
Chen 2011	Control treatment (Celebrex) did not met the inclusion criteria.					

Study	Reason for exclusion							
Gu 2008	Control treatment did not met the inclusion criteria (eight methods of intelligent turtle combined with point selection treatment based on syndrome differentiation versus point selection treatment based on syndrome differentiation)							
Hong 2002	Quasi-randomised (by the visiting sequence).							
Hu 2010	Control treatment (Xiaoyao Pill) did not met the inclusion criteria.							
Jiang 2005	Quasi-randomised (by the visiting sequence).							
Koleini 2017	Control treatment (vitamin B6) did not meet the inclusion criteria.							
Li 2004	Not randomised, no control group.							
Li 2008	Not randomised, no control group.							
Li 2014	Control treatment (Xiaoyao Pill) did not met the inclusion criteria.							
Sun 2004	Quasi-randomised (by the visiting sequence). Control treatment was comprehensive treatment method (medroxyprogesterone acetate and diazepam).							
Sun 2008	Control treatment (ibuprofen) did not met the inclusion criteria.							
Wang 2003	Not randomised, no control group.							
Xu 2006a	Control treatment was comprehensive treatment method (medroxyprogesterone acetate and di- azepam).							
Xu 2006b	Control treatment was comprehensive treatment method (medroxyprogesterone acetate and di- azepam).							
Xu 2011	Control treatment was comprehensive treatment method (Vitamin B1 and Oryzanol and Xiaoyao Pills).							
Zhang 1994	Control treatment was comprehensive treatment method (Ciwujia Injection combined with di- azepam and Xiaoyao Pills).							
Zhao 2012	Control treatment did not met the inclusion criteria (fire-needles versus ordinary needles).							
Zheng 2001	Control treatment (vitamin B6) did not meet inclusion criteria.							

Characteristics of studies awaiting assessment [ordered by study ID]

Kim 2005

2005	
Methods	Acupuncture vs placebo acupuncture
Participants	20 participants with PMS screened by DRSP
Interventions	Experimental group received acupuncture at CV6 and SP6 plus other points based on diagnosis. Control group received acupuncture at SI5 and ST40. Up to 13 sessions over 8 weeks for both groups
Outcomes	Digital infrared thermal imaging (DITI), DRSP and progesterone levels.

Acupuncture and acupressure for premenstrual syndrome (Review)

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Kim 2005 (Continued)

Notes

Unable to contact author to determine randomisation status and group numbers.

Kurebayashi 2013

Methods	A randomised clinical trial aimed at evaluating the auriculotherapy
Participants	40 females aged 21 to 44 years
Interventions	Control group: no treatment Experimental group: Traditional Chinese medicine ear points: Kidney, Liver, Stomach, Brain Stem, Ovary and Uterus point with semi-permanent needles once per week for 8 sessions.
Outcomes	Premenstrual syndrome symptoms
Notes	Completed July 2012. Contacted CI (Leonice Fumiko Sato Kurebayashi) in May 2018 regarding data and publications. No response

DRSP: Daily Record of Severity of Problems scale

Characteristics of ongoing studies [ordered by study ID]

NCT02504515

Trial name or title	Effect of homeopathy, acupuncture or anthroposophic medicine in women's quality of life
Methods	A randomised clinical trial aimed to evaluating the effect of homeopathy, acupuncture or anthro- posophic medicine
Participants	906 females aged 18 years and over
Interventions	Homeopathy
	Acupuncture
	Anthroposophic medicine
	No further details provided
Outcomes	World Health Organization Bref (WHOQOL-BREF) questionnaire
Starting date	April 2015
Contact information	Rubens Lene Carvalho Tavares, Professor, Federal University of Minas Gerais
Notes	Estimated completion date December 2017

DATA AND ANALYSES

Comparison 1. Acupuncture vs sham acupuncture

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Overall premenstrual symptoms- Mood	1	67	Mean Difference (IV, Fixed, 95% CI)	-9.03 [-10.71, -7.35]
2 Overall premenstrual symptoms - Physical	1	67	Mean Difference (IV, Fixed, 95% CI)	-9.11 [-10.82, -7.40]
3 Adverse events	3	167	Risk Ratio (M-H, Fixed, 95% CI)	1.74 [0.39, 7.76]
4 Response rate	2	100	Risk Ratio (M-H, Random, 95% CI)	4.22 [0.45, 39.88]
5 Quality of Life	1	67	Mean Difference (IV, Fixed, 95% CI)	2.85 [1.47, 4.23]

Analysis 1.1. Comparison 1 Acupuncture vs sham acupuncture, Outcome 1 Overall premenstrual symptoms- Mood.

Study or subgroup	Acu	Acupuncture N Mean(SD)		Sham Acupuncture N Mean(SD)		Mean Difference				Weight	Mean Difference
	Ν					Fixed, 95% CI					Fixed, 95% CI
Zhang 2017	34	42.9 (3.3)	33	51.9 (3.8)						100%	-9.03[-10.71,-7.35]
Total ***	34		33		•					100%	-9.03[-10.71,-7.35]
Heterogeneity: Not applicable											
Test for overall effect: Z=10.52(P<0	.0001)							1	1		
			Favours	acupuncture	-10	-5	0	5	10	Favours sham	

Analysis 1.2. Comparison 1 Acupuncture vs sham acupuncture, Outcome 2 Overall premenstrual symptoms - Physical.

Study or subgroup	Acu	puncture	Sham A	cupuncture	Mean Difference			Weight	Mean Difference		
	Ν	Mean(SD)	Ν	Mean(SD)		Fix	xed, 95% Cl				Fixed, 95% CI
Zhang 2017	34	37 (3.5)	33	46.1 (3.7)						100%	-9.11[-10.82,-7.4]
Total ***	34		33		•					100%	-9.11[-10.82,-7.4]
Heterogeneity: Not applicable											
Test for overall effect: Z=10.42(P<0	.0001)					i.					
			Favours	acupuncture	-10	-5	0	5	10	Favours sham	

Analysis 1.3. Comparison 1 Acupuncture vs sham acupuncture, Outcome 3 Adverse events.

Study or subgroup	Acupuncture	Sham Acupuncture		Risk Ratio			Weight	Risk Ratio
	n/N	n/N	M-H	, Fixed, 9	95% CI			M-H, Fixed, 95% CI
Habek 2002	1/18	0/17	_		•	-	20.19%	2.84[0.12,65.34]
Yu 2006	0/33	0/32						Not estimable
Zhang 2017	3/34	2/33	1		<u> </u>	1	79.81%	1.46[0.26,8.16]
		Favours sham	0.002 0.1	1	10	500	Favours acupuncture	



Study or subgroup	Acupuncture	Sham Acupuncture		Risk Ratio				Weight	Risk Ratio
	n/N	n/N		M-H, F	ixed, 9	95% CI			M-H, Fixed, 95% Cl
Total (95% CI)	85	82						100%	1.74[0.39,7.76]
Total events: 4 (Acupuncture), 2 (Sham Acupuncture)								
Heterogeneity: Tau ² =0; Chi ² =	0.14, df=1(P=0.71); l ² =0%								
Test for overall effect: Z=0.72	(P=0.47)								
		Favours sham	0.002	0.1	1	10	500	Favours acupuncture	

Analysis 1.4. Comparison 1 Acupuncture vs sham acupuncture, Outcome 4 Response rate.

Study or subgroup	Acupuncture	Sham Acupuncture		Risk Ratio			Weight	Risk Ratio
	n/N	n/N		M-H, Random, 95	% CI			M-H, Random, 95% CI
Habek 2002	14/18	1/17					41.52%	13.22[1.94,89.96]
Yu 2006	29/33	15/32		-			58.48%	1.87[1.27,2.77]
Total (95% CI)	51	49					100%	4.22[0.45,39.88]
Total events: 43 (Acupuncture	e), 16 (Sham Acupuncture)							
Heterogeneity: Tau ² =2.21; Ch	i ² =5.43, df=1(P=0.02); l ² =81.5	8%						
Test for overall effect: Z=1.26	(P=0.21)					1		
		Favours sham	0.002	0.1 1	10	500	Favours acupuncture	

Analysis 1.5. Comparison 1 Acupuncture vs sham acupuncture, Outcome 5 Quality of Life.

Study or subgroup	Acu	puncture	Sham A	Sham Acupuncture Mean Diffe		n Difference	Weight	Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)		Fix	ed, 95% CI		Fixed, 95% CI
Zhang 2017	34	76.7 (3.1)	33	73.8 (2.6)				100%	2.85[1.47,4.23]
Total ***	34		33					100%	2.85[1.47,4.23]
Heterogeneity: Not applicable									
Test for overall effect: Z=4.05(P<0.0	001)								
				Favours sham	-5	-2.5	0 2.5 5	Favours acu	Ipuncture

Comparison 2. Acupuncture vs no treatment

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Overall premenstrual symp- toms	1	14	Mean Difference (IV, Fixed, 95% CI)	-13.6 [-15.70, -11.50]
2 Adverse events	1	20	Risk Ratio (M-H, Fixed, 95% CI)	0.0 [0.0, 0.0]

Study or subgroup	Acu	Acupuncture		No Treatment		Mean Difference			Weight	Mean Difference	
	Ν	Mean(SD)	Ν	Mean(SD)		Fixed, 95% CI				Fixed, 95% CI	
Shin 2009	7	3.9 (1.7)	7	17.5 (2.3)			+			100%	-13.6[-15.7,-11.5]
Total ***	7		7				•			100%	-13.6[-15.7,-11.5]
Heterogeneity: Not applicable											
Test for overall effect: Z=12.72(P<0	.0001)										
			Favours	Acupuncture	-100	-50	0	50	100	Favours No	Treatment

Analysis 2.1. Comparison 2 Acupuncture vs no treatment, Outcome 1 Overall premenstrual symptoms.

Analysis 2.2. Comparison 2 Acupuncture vs no treatment, Outcome 2 Adverse events.

Study or subgroup	Acupuncture	No Treatment		Risk Ra				Weight	Risk Ratio
	n/N	n/N		м-н,	Fixed, 95°	% CI			M-H, Fixed, 95% CI
Shin 2009	0/10	0/10							Not estimable
Total (95% CI)	10	10							Not estimable
Total events: 0 (Acupuncture), 0 (No	Treatment)								
Heterogeneity: Not applicable									
Test for overall effect: Not applicable	5								
	Fav	ours no treatment	0.01	0.1	1	10	100	Favours acupuncture	

Comparison 3. Acupressure vs sham acupressure

Outcome or subgroup title	No. of studies	No. of partici- pants	Statistical method	Effect size
1 Overall premenstrual symp- toms	1	90	Risk Ratio (M-H, Fixed, 95% CI)	0.64 [0.52, 0.79]
2 Quality of Life	1		Mean Difference (IV, Fixed, 95% CI)	Subtotals only

Analysis 3.1. Comparison 3 Acupressure vs sham acupressure, Outcome 1 Overall premenstrual symptoms.

Study or subgroup	Acupressure	Sham Acu- pressure			Risk Ratio			Weight	Risk Ratio
	n/N	n/N		м	I-H, Fixed, 95	% CI			M-H, Fixed, 95% CI
Bazarganipour 2017	37/60	29/30			+-			100%	0.64[0.52,0.79]
Total (95% CI)	60	30			•			100%	0.64[0.52,0.79]
Total events: 37 (Acupressure),	29 (Sham Acupressure)								
Heterogeneity: Not applicable									
Test for overall effect: Z=4.19(P-	<0.0001)								
		Favours sham	0.05	0.2	1	5	20	Favours acupressure	

Study or subgroup	Acu	pressure	Sham Acupressure			Mean Difference				Weight	Mean Difference
	N	Mean(SD)	Ν	Mean(SD)	Fixed, 95% CI		% CI			Fixed, 95% CI	
Bazarganipour 2017	60	69.5 (7.8)	30	52.3 (10)				+		0%	17.17[13.08,21.26]
Bazarganipour 2017	60	91.5 (13.4)	30	67.2 (17.5)				+		0%	24.3[17.18,31.42]
				Favours sham	-50	-25	0	25	50	Favours acu	pressure

Analysis 3.2. Comparison 3 Acupressure vs sham acupressure, Outcome 2 Quality of Life.

APPENDICES

Appendix 1. Cochrane Gynaecology and Fertlity specialised register search strategy

Procite platform

Searched 21 September 2017

Keywords CONTAINS "premenstrual " or "premenstrual dysphoria" or "premenstrual aggravation" or "premenstrual dysphoric disorder" or "premenstrual pain " or "premenstrual symptoms" or "Premenstrual Syndrome-Symptoms" or "PMS" or "dysphoria" or "PMDD" or Title CONTAINS "premenstrual " or "premenstrual dysphoria" or "premenstrual aggravation" or "premenstrual dysphoric disorder" or "premenstrual aggravation" or "premenstrual or "premenstrual dysphoria" or "PMDD" or "premenstrual aggravation" or "premenstrual or "PMDD" or "premenstrual aggravation" or "premenstrual or "PMDD" or "premenstrual aggravation" or "premenstrual or "premenstrual or "premenstr

AND

Keywords CONTAINS "acupoint" or "acupressure" or "acupressure-acupuncture therapy" or "acupuncture" or "electro-acupuncture" or "electro-magnetic" or "electroacupuncture" or "electrical stimulation" or "moxibustion" or "*acupuncture" or "auricular acupressure" or "auricular pressure" or "needle insertion" or Title CONTAINS "acupoint" or "acupressure" or "acupressure-acupuncture therapy" or "acupuncture" or "electro-acupuncture" or "electro-magnetic" or "electro-acupuncture" or "acupressure" or "acupressure" or "acupressure" or "acupressure" or "acupuncture" or "electro-acupuncture" or "electro-acupuncture" or "electro-acupuncture" or "acupressure" or "acupressure" or "acupuncture" or "electro-acupuncture" or "acupuncture" or "acupation" or "moxibustion" or "moxibustion" or "moxibustion" or "moxibustion" or "acupation" or "moxibustion" or "moxibustion" or "moxibustion" or "moxibustion" or "acupation" or "acupation

Appendix 2. CENTRAL Register of Studies Online (CRSO) search strategy

Web platform Searched 21 September 2017

#1 MESH DESCRIPTOR Premenstrual Syndrome EXPLODE ALL TREES 385

#2 (Premenstrua* adj5 Syndrome*):TI,AB,KY 655

#3 dysphor*:TI,AB,KY 781

#4 (PMS or PMT):TI,AB,KY 558

#5 (premenstrua* adj5 tension):TI,AB,KY 75

#6 premenstrual:TI,AB,KY 845

#7 (late luteal phase adj5 disorder*):TI,AB,KY 23

#8 PMD:TI,AB,KY 75

 $\#9\ \#1\ \text{OR}\ \#2\ \text{OR}\ \#3\ \text{OR}\ \#4\ \text{OR}\ \#5\ \text{OR}\ \#6\ \text{OR}\ \#7\ \text{OR}\ \#8\ 1828$

#10 MESH DESCRIPTOR Acupuncture Therapy EXPLODE ALL TREES 3646

#11 acupuncture:TI,AB,KY 8917

#12 acupressure:TI,AB,KY 628

#13 (Electroacupunctur* or electro-acupunctur*):TI,AB,KY 1426

#14 acupoint*:TI,AB,KY 2025



#15 meridian*:TI,AB,KY 603

#16 (non-meridian* or trigger*):TI,AB,KY 4884

#17 (moxibustion or moxa*):TI,AB,KY 1088

 $\#18\ \#10\ \text{OR}\ \#11\ \text{OR}\ \#12\ \text{OR}\ \#13\ \text{OR}\ \#14\ \text{OR}\ \#15\ \text{OR}\ \#16\ \text{OR}\ \#17\ 15644$

#19 #9 AND #18 42

Appendix 3. MEDLINE search strategy

Ovid platform Searched from 1946 to 21 September 2017

1 exp Premenstrual Syndrome/ (3983)

2 (Premenstrua\$ adj5 Syndrome\$).tw. (2446)

3 dysphor\$.tw. (4926)

4 pms.tw. (4737)

5 pmt.tw. (1661)

6 (premenstrua\$ adj5 tension\$).tw. (503)

7 premenstrual.tw. (4795)

8 (late luteal phase adj5 disorder).tw. (85)

9 llpd\$.tw. (38)

10 pmd\$.tw. (4411)

11 or/1-10 (18786)

12 exp Acupuncture/ (1506)

13 exp acupuncture therapy/ or exp acupressure/ or exp acupuncture analgesia/ or exp acupuncture, ear/ or exp electroacupuncture/ or exp meridians/ or exp moxibustion/ (21564)

14 acupuncture.tw. (18640)

15 acupressure.tw. (880)

- 16 exp Electroacupuncture/ (3258)
- 17 (Electroacupunctur\$ or electro-acupunctur\$).tw. (4176)
- 18 acupoint\$.tw. (4083)
- 19 meridian\$.tw. (4819)
- 20 ((meridian or non-meridian or trigger) adj10 point\$).tw. (2478)

21 exp Moxibustion/ (1542)

22 (moxibustion or moxabustion or moxa\$).tw. (3206)

23 or/12-22 (33736)

24 11 and 23 (60)



25 randomized controlled trial.pt. (481966)

26 controlled clinical trial.pt. (96881)

27 randomized.ab. (422264)

28 randomised.ab. (82875)

29 placebo.tw. (201990)

30 clinical trials as topic.sh. (190475)

31 randomly.ab. (292311)

32 trial.ti. (190079)

33 (crossover or cross-over or cross over).tw. (78725)

34 or/25-33 (1236937)

35 exp animals/ not humans.sh. (4585712)

36 34 not 35 (1140717)

37 24 and 36 (29)

Appendix 4. Embase search strategy

Ovid platform Searched from 1980 to 21 September 2017

1 exp premenstrual dysphoric disorder/ or exp premenstrual syndrome/ (6019)

2 (Premenstrua\$ adj5 Syndrome\$).tw. (2862)

3 dysphor\$.tw. (6118)

4 (pms or pmt).tw. (7991)

5 (premenstrua\$ adj5 tension\$).tw. (356)

6 premenstrual.tw. (5287)

7 (late luteal phase adj5 disorder).tw. (103)

8 (llpd\$ or pmd\$).tw. (6147)

9 or/1-8 (24195)

10 exp acupuncture analgesia/ or exp acupuncture/ or exp acupuncture needle/ (39566)

11 exp acupressure/ (1741)

12 (acupuncture or acupressure).tw. (26239)

13 exp electroacupuncture/ (5482)

14 (Electroacupunctur\$ or electro-acupunctur\$).tw. (5132)

15 acupoint\$.tw. (5137)



- 16 meridian\$.tw. (5511)
- 17 ((meridian or non-meridian or trigger) adj10 point\$).tw. (3607)
- 18 exp moxibustion/ (2335)
- 19 or/10-18 (48148)
- 209 and 19 (142)
- 21 Clinical Trial/ (947951)
- 22 Randomized Controlled Trial/ (469604)
- 23 exp randomization/ (75622)
- 24 Single Blind Procedure/ (29554)
- 25 Double Blind Procedure/ (140190)
- 26 Crossover Procedure/ (53240)
- 27 Placebo/ (299862)
- 28 Randomi?ed controlled trial\$.tw. (167370)
- 29 Rct.tw. (25692)
- 30 random allocation.tw. (1685)
- 31 randomly allocated.tw. (28302)
- 32 allocated randomly.tw. (2267)
- 33 (allocated adj2 random).tw. (784)
- 34 Single blind\$.tw. (19787)
- 35 Double blind\$.tw. (175378)
- 36 ((treble or triple) adj blind\$).tw. (713)
- 37 placebo\$.tw. (255724)
- 38 prospective study/ (402253)
- 39 or/21-38 (1804452)
- 40 case study/ (49880)
- 41 case report.tw. (338520)
- 42 abstract report/ or letter/ (1010607)
- 43 or/40-42 (1390853)
- 44 39 not 43 (1758453)
- 45 20 and 44 (63)



Appendix 5. PsycINFO search strategy

Ovid platform Searched from 1806 to 21 September 2017 1 exp Premenstrual Dysphoric Disorder/ (429) 2 exp Premenstrual Syndrome/ (1475) 3 (Premenstrua\$ adj5 Syndrome\$).tw. (1189) 4 dysphor\$.tw. (5709) 5 (pms or pmt).tw. (1684) 6 (premenstrua\$ adj5 tension\$).tw. (172) 7 premenstrual.tw. (2479) 8 (late luteal phase adj5 disorder).tw. (112) 9 (llpd\$ or pmd\$).tw. (1045) 10 or/1-9 (8817) 11 exp Acupuncture/ (1313) 12 exp Acupuncture/ (1313) 13 (acupuncture or acupressure).tw. (1882) 14 exp Electrical Stimulation/ (19571) 15 (Electroacupunctur\$ or electro-acupunctur\$).tw. (308) 16 acupoint\$.tw. (241) 17 meridian\$.tw. (781) 18 ((meridian or non-meridian or trigger) adj10 point\$).tw. (408) 19 or/11-18 (22509) 20 10 and 19 (40) 21 random.tw. (50922) 22 control.tw. (393763) 23 double-blind.tw. (20962) 24 clinical trials/ (10575) 25 placebo/ (4969) 26 exp Treatment/ (694599)

27 or/21-26 (1077010)

28 20 and 27 (35)



Appendix 6. AMED search strategy

Ovid platform Searched from 1985 to 21 September 2017

1 exp Premenstrual syndrome/ (107)

2 (Premenstrua\$ adj5 Syndrome\$).tw. (165)

3 dysphor\$.tw. (39)

4 (pms or pmt).tw. (94)

5 (premenstrua\$ adj5 tension\$).tw. (7)

6 premenstrual.tw. (171)

7 (late luteal phase adj5 disorder).tw. (0)

8 (llpd\$ or pmd\$).tw. (20)

9 or/1-8 (273)

10 exp Acupuncture/ (3292)

11 exp Acupressure/ or exp Acupuncture therapy/ (7689)

12 (acupuncture or acupressure).tw. (9807)

13 exp Electroacupuncture/ (878)

14 (Electroacupunctur\$ or electro-acupunctur\$).tw. (1070)

15 acupoint\$.tw. (1983)

16 meridian\$.tw. (671)

17 ((meridian or non-meridian or trigger) adj10 point\$).tw. (554)

18 exp Moxibustion/ (537)

19 or/10-18 (11126)

20 9 and 19 (16)

Appendix 7. CINAHL search strategy

Ebsco platform Searched from 1961 to 21 September 2017

#	Query	Results
S35	S20 AND S34	19
S34	S21 OR S22 or S23 or S24 OR S25 OR S26 OR S27 OR S28 OR S29 OR S30 OR S31 OR S32 OR S33	1,160,698
\$33	TX allocat* random*	7,195



Cochrane Database of Systematic Reviews

(Continued)		
S32	(MH "Quantitative Studies")	16,373
S31	(MH "Placebos")	10,354
S30	TX placebo*	47,288
S29	TX random* allocat*	7,195
S28	(MH "Random Assignment")	44,035
S27	TX randomi* control* trial*	131,386
S26	TX ((singl* n1 blind*) or (singl* n1 mask*)) or TX ((doubl* n1 blind*) or (dou- bl* n1 mask*)) or TX ((tripl* n1 blind*) or (tripl* n1 mask*)) or TX ((trebl* n1 blind*) or (trebl* n1 mask*))	906,331
S25	TX ((trebl* n1 blind*) or (trebl* n1 mask*))	210
S24	TX ((trebl* n1 blind*) or (trebl* n1 mask*))	210
S23	TX clinic* n1 trial*	211,140
S22	PT Clinical trial	80,028
S21	(MH "Clinical Trials+")	221,365
S20	S9 AND S19	51
S19	S10 OR S11 OR S12 OR S13 OR S14 OR S15 OR S16 OR S17 OR S18	18,602
S18	TX (moxibustion or moxabustion or moxa*)	1,198
S17	(MM "Moxibustion")	264
S16	TX meridian*	2,334
S15	TX acupoint*	983
S14	TX (Electroacupunctur* or electro-acupunctur*)	1,342
S13	TX Electroacupuncture	1,296
S12	TX acupressure	1,193
S11	TX acupuncture	16,542
S10	(MH "Acupuncture+") OR (MM "Acupuncture, Ear") OR (MM "Acupuncture Points") OR (MM "Acupuncturists") OR (MM "Acupuncture Anesthesia") OR (MM "Acupuncture Analgesia")	12,300
S9	S1 OR S2 OR S3 OR S4 OR S5 OR S6 OR S7 OR S8	3,658
S8	TX(late luteal phase N5 disorder)	11
S7	TX premenstrual	1,780



(Continued)		
S6	TX (premenstrua* N5 tension*)	46
S5	TX pmt	229
S4	TX pms	1,210
S3	TX dysphor*	1,292
S2	TX(Premenstrua* N5 Syndrome*)	1,444
S1	(MM "Premenstrual Syndrome+") OR (MM "Premenstrual Dysphoric Disor- der+")	1,074

WHAT'S NEW

Date	Event	Description
23 August 2018	Amended	Abstract text corrected.

HISTORY

Protocol first published: Issue 2, 2005 Review first published: Issue 8, 2018

Date	Event	Description
28 January 2015	Amended	Converted to new review format.

CONTRIBUTIONS OF AUTHORS

Mike Armour: performed the screening of the searches, ran the additional Google Scholar and PubMed searches, rated studies using the NICMAN scale, entered the data, performed the meta-analysis and interpretation, generated the GRADE tables, and took the lead in writing the review.

Carolyn Ee: performed the grey literature search, searched the trial registries, assisted MA with screening, performed data extraction, rated studies using the NICMAN scale and commented on each draft of the review.

Jie Hao: searched the Chinese literature, screened Chinese language studies, performed data extraction, rated studies using the NICMAN scale and commented on the final version of the review.

Tania Wilson: performed data extraction, rated studies using the NICMAN scale and commented on the final version of the review.

Sofia Yao: performed data extraction, rated studies using the NICMAN scale and commented on the final version of the review.

Caroline A Smith: performed data extraction, rated studies using the NICMAN scale and commented on the final version of the review.

DECLARATIONS OF INTEREST

MA: is an acupuncturist recently involved in clinical practice and until recently, the director of an acupuncture clinic. As a medical research institute, NICM receives research grants and donations from foundations, universities, government agencies and industry. Sponsors and donors provide untied and tied funding for work to advance the vision and mission of the Institute.



CE: is a practising acupuncturist and the director of an integrative health centre which provides acupuncture clinical services to the general public. As a medical research institute, NICM receives research grants and donations from foundations, universities, government agencies and industry. Sponsors and donors provide untied and tied funding for work to advance the vision and mission of the Institute.

CS: none known. As a medical research institute, NICM receives research grants and donations from foundations, universities, government agencies and industry. Sponsors and donors provide untied and tied funding for work to advance the vision and mission of the Institute.

JH: is a Chinese medicine practitioner and acupuncturist.

SY: none known

TW: holds dual registration as a psychologist and acupuncturist

SOURCES OF SUPPORT

Internal sources

• Mike Armour, Australia.

National Institute of Complementary Medicine, Western Sydney University

• Carolyn Ee, Australia.

National Institute of Complementary Medicine, Western Sydney University

• Caroline Smith, Australia.

National Institute of Complementary Medicine, Western Sydney University

External sources

• No sources of support supplied

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

- 1. The standard of 'assessment of quality of included studies' had changed, given in the new version of the *Cochrane Handbook of Systematic Reviews of Interventions* 5.1.0.
- 2. The method of control group of included studies had changed. In the protocol, control intervention could be Chinese medicine, western medicine, sham acupuncture, no intervention, and any other interventions, but in this review, the control group method changed to sham acupuncture, blank control (no intervention) or western medicine such as SSRIs, oral contraceptives, gonadotropin-releasing hormone agonists, danazol or estradiol which were mentioned by ISPMD. Because if the effect of one control intervention is uncertain, it is meaningless to compare another intervention with it.
- 3. Acupressure was added as an appropriate intervention for inclusion in this review and title changed to "Acupuncture and acupressure for premenstrual syndrome" to reflect this.
- 4. NICMAN scale scores were added to provide a measure of the quality of the acupuncture or acupressure intervention delivered.
- 5. Response rates were added as a secondary outcome measure as this allowed inclusion of data not reported in a standard validated scale.

INDEX TERMS

Medical Subject Headings (MeSH)

*Acupressure; *Acupuncture Therapy; *Quality of Life; Luteal Phase [*psychology]; Premenstrual Dysphoric Disorder [*therapy]; Premenstrual Syndrome [*therapy]; Randomized Controlled Trials as Topic

MeSH check words

Female; Humans