Practice, practitioner, or placebo? A multifactorial, mixed-methods randomized controlled trial of acupuncture

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ABSTRACT

The nonspecific effects of acupuncture are well documented; we wished to quantify these factors in osteoarthritic (OA) pain, examining needling, the consultation, and the practitioner. In a prospective randomised, single-blind, placebo-controlled, multifactorial, mixed-methods trial, 221 patients with OA awaiting joint replacement surgery were recruited. Interventions were acupuncture, Streitberger placebo acupuncture, and mock electrical stimulation, each with empathic or nonempathic consultations. Interventions involved eight 30-minute treatments over 4 weeks. The primary outcome was pain (VAS) at 1 week posttreatment. Face-to-face qualitative interviews were conducted (purposive sample, 27 participants). Improvements occurred from baseline for all interventions with no significant differences between real and placebo acupuncture (mean difference −2.7 mm, 95% confidence intervals −9.0 to 3.6; P = .40) or mock stimulation (−3.9, −10.4 to 2.7; P = .26). Empathic consultations did not affect pain (3.0 mm, −2.2 to 8.2; P = .26) but practitioner 3 achieved greater analgesia than practitioner 2 (10.9, 3.9 to 18.0; P = .002). Qualitative analysis indicated that patients' beliefs about treatment veracity and confidence in outcomes were reciprocally linked. The supportive nature of the trial attenuated differences between the different consultation styles. Improvements occurred from baseline, but acupuncture has no specific efficacy over either placebo. The individual practitioner and the patient's belief had a significant effect on outcome. The 2 placebos were equally as effective and credible as acupuncture. Needle and nonneedle placebos are equivalent. An unknown characteristic of the treating practitioner predicts outcome, as does the patient's belief (independently). Beliefs about treatment veracity shape how patients self-report outcome, complicating and confounding study interpretation.

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1. Introduction

All complex therapeutic interventions achieve their outcomes through a combination of specific and nonspecific mechanisms, including the patient–therapist relationship, treatment process and rituals, expectation, suggestibility, conditioning, and practitioner bias [1,11]. Acupuncture studies show very large improvements from pretreatment to posttreatment [2,28], with mean improvements of 80% for symptom severity (across painful pathologies) and 62% for pain [6]. There is a significantly greater benefit from acupuncture than from standard conventional care [5], with acupuncture “placebos” routinely outperforming conventional care [3,12,25]. This suggests that the effectiveness of acupuncture is not point- or therapy-specific.

There is considerable interest in maximising treatment effects through understanding the placebo more generically [10,16]. Acupuncture's effectiveness coupled with marginal proven efficacy implies that contextual factors may have a profound influence on outcome. Kaptchuk et al. suggest that acupuncture needles create enhanced placebo effects [15], implying that different placebo rituals may demonstrate different relative benefits.

The aims of this study were to investigate whether there is an enhanced nonspecific effect associated with needling, to ascertain the effects of the consultation process and the practitioner, to evaluate the efficacy of acupuncture on severe osteoarthritic (OA) pain, and to enhance interpretation of the quantitative study through nested qualitative work.
2. Materials and methods

A randomised, single-blind (patient), multifactorial trial was conducted involving 3 interventions, 2 consultation types, and 3 practitioners with a nested qualitative study to elucidate the quantitative findings. Recruitment was via joint replacement waiting lists at Southampton General and Salisbury District Hospitals (Table 1).

2.1. Randomisation

After informed consent and 1-week baseline pain recording, a 2-stage randomisation occurred via an independent third party using a computer-generated list. Patients were first randomised to treatment type and consultation type, and then to specific practitioners.

2.2. Interventions

2.2.1. Real acupuncture

A Western acupuncture approach was used involving a flexible but prescribed range of points. Practitioners were free to pick appropriately from the list as clinically indicated. A mean of 6 points was used at each treatment, with deep needling, which lasted for 20 minutes during the 30-minute appointment twice per week for 4 weeks. Deqi (needle sensation) was elicited for each needle through needle rotation. The acupuncture treatment was designed to provide high-quality Western acupuncture of the type commonly used in UK clinics.

2.2.2. Placebo acupuncture

The format was exactly the same as those for real acupuncture (RA), but using streitberger needle (SN) nonpenetrating needles instead of RA needles. These work rather like a stage dagger, have been validated, and patients cannot distinguish them from RA needles [27].

2.2.3. Mock electrical stimulation

An electroacupuncture stimulator was used (NOMA Ltd., Southampton, UK) to provide mock transcutaneous electrical stimulation to acupoints via electrodes fixed to the surface of the patient’s skin. The cables were disconnected inside the output plug. All aspects of the intervention were exactly the same as with RA and SN, including time of treatment and checking on patients at regular intervals. This control has previously been well validated [22].

2.3. Consultation types

2.3.1. Empathic

Empathic (EMP) consultations were deemed to be normal pragmatic treatment sessions. Patients were greeted in a friendly, warm manner and were free to enter into conversation with their practitioner, who in turn would willingly do so. Practitioners did their utmost to comply with participants’ wishes, providing detailed answers to questions and emphasising patient comfort and well-being.

2.3.2. Non-EMP

This encounter was more “clinical” in nature. Patients were greeted in an efficient manner and quietly shown to the treatment cubicle. Practitioners would only discuss matters directly relating to the treatment to enable them to effectively carry out that treatment, e.g., pattern of pain and side effects. Necessary explanations were kept as short as possible, and if patients attempted to enter into any discussion, the practitioner would respond using the words “I'm sorry but because this is a trial I am not allowed to discuss this with you.” Between needle stimulations, patients were left on their own in a curtained cubicle.

2.3.3. Acupuncturists

Three qualified, experienced (range 3 to 10 years) practitioners were available and funded to provide treatments (physiotherapist, nurse, and licensed acupuncturist). They met frequently throughout the trial to ensure that treatments, including the acupoints used, and consultation types were as comparable and equivalent as possible.

2.4. Protocol/procedure

Patients were assessed and gave consent (including consent to differing consultation types). Ethics approval was gained from the Southhampton and South West Hampshire and the Salisbury and South Wiltshire Research ethics committees (approval number 170/03/7).

The information sheet explained that “a proportion of patients will receive placebo (dummy) treatment and so the acupuncture points will not be stimulated; i.e., the needle will not penetrate your skin or the machine will not deliver any current.”

Patients were given a daily pain diary (100 mm visual analogue scale VAS) to complete for 7 (pretreatment) days, which included recording their analgesia. A minimum mean weekly score of 30 of 100 was an inclusion requirement. Patients were then randomised and treated. Throughout treatment, patients recorded their weekly pain and analgesia with a final extra weeks’ diary after treatment completion.

Face-to-face open-ended narrative qualitative interviews were conducted with 27 randomized controlled trial (RCT) participants purposively sampled to obtain representation from all treatment groups (EMP and non-EMP, all 3 interventions and the 2 practitioners not conducting the qualitative study), and both responders and nonresponders. Interviews took place 4 to 8 weeks after treatment completion. There was no topic guide, apart from the opening question in which participants were asked to talk about their experiences of taking part in the trial. Interviews lasted between 30 and 120 minutes (typically 60 minutes), and were audio-taped, transcribed, and annotated with the interviewer’s field notes. Summaries were sent to interviewees to provide a member check on the analysis. Patients were not interviewed by their treating acupuncturist (Table 2).

2.5. Outcomes

Pain measured on a 100-mm VAS in the 7 days immediately after treatment completion. The primary outcome was the differences between groups in percentage change in pain. Differences of >30% improvement from baseline were defined as clinically important [8,9]. Secondary outcomes were Nottingham Health Profile (part 1) and Western Ontario and McMaster University Osteoarthritis Index and pain over time (weekly VAS).
2.6. Potential confounders

To assess potential confounding, we recorded treatment credibility [7], attitudes to complementary medicine holistic complementary and alternative medicine questionnaire (HCAMQ) [13], empathy consultation and relational empathy (CARE) questionnaire [19], analgesic intake (tablet count), and needling sensation [20]. Patients were asked at treatment completion, “Do you think the treatment you had was real” and required to give a yes or no answer.

2.7. Sample size and analyses

Assuming a 13% dropout, 96 subjects were required per arm for 80% power to detect a 20% difference between each group on VAS (assuming SD = 38 mm) for our primary outcome. To avoid an accumulated type I error from multiple testing, we required a P value of .01 or less, creating a sample size of 288.

An analysis of covariance (ANCOVA) for pain (VAS) estimated main effects and interactions between factors and controlled for baseline pain, stratifiers, and confounders. If interactions were not found, the main effects with confidence intervals were reported. Missing data were dealt with by either taking a mean of the values on either side of the missing value or by carrying forward the last entry as appropriate. Our per-protocol analysis examined needling sensation, belief in treatment, and attitudes to complementary medicine, as well as treatment type, the empathic nature of the consultation, and practitioner.

Framework analysis [24] incorporating deductive and inductive elements was used to relate the qualitative data to the quantitative trial. An index was used by 2 authors (F.B., G.L.) to identify talk that related to the 4 main predetermined elements of the framework: outcomes, intervention characteristics, practitioners, and empathy. Talk that related to each of these themes was explored inductively to identify subthemes. Predetermined comparisons were then made across intervention groups by examining how each subtheme manifested in the different participants’ accounts. These comparisons suggested consistent explanations for the main RCT findings. Table 3 shows how the RCT findings were used to determine the 4 key themes and how data mapped to themes, and were explored inductively to identify subthemes. The findings are presented with illustrative quotes (using pseudonyms) selected as typical and/or particularly clear examples.

3. Results

Patients (279) were assessed and completed baseline pain recordings (01/2004 to 08/2007) with 221 patients recruited (Fig. 1). Eleven patients (4.9%) dropped out (Table 4); 3 completed some pain scores, and their data were carried forward per intention-to-treat analysis and included in the final analysis. Eight did not complete any outcomes and were excluded from analysis.

3.1. Demographics and baseline measurements

Age (mean = 66.75 years, SD = 8.29) and sex were balanced across all treatments, consultation types, and practitioners; 57.5% were female; 59.7% knee joints were treated. Table 5 shows baseline data and mean posttreatment scores.

The ANCOVA showed that treatment credibility and HCAMQ scores had no effect on outcome, implying that equipoise between groups was achieved. The final empathy questionnaire demonstrated statistically significant differences between consultation types, with mean scores of 45.73 (95% confidence interval [CI] 44.73 to 46.73) (empathic) and 36.38 (95% CI 34.34 to 38.42, Mann-Whitney U, P < .001) (nonempathic) demonstrating that this factor was delivered effectively. This difference in empathy for the CARE score, based on available data, also demonstrates a clinically significant effect for the EMP and non-EMP consultation types [18].

3.2. Primary outcome—pain

Pain diminished for all groups, with only RA achieving clinically relevant improvements at 29.5% (Table 5). SN reduced pain by 23.0% and mock electrical stimulation by 16.6%. Those allocated to nonempathic consultations achieved slightly (nonsignificant) greater analgesia than empathic consultations (25.2% vs 21.1%; mean VAS difference 3.00 mm, 95% CI –2.2 to 8.2, P = .26).

ANCOVA at week 5 showed no significant effect on immediate posttreatment pain of the factors practitioner, treatment, and consultation type, and their first-order interactions with all of the possible baseline confounders as covariates (age, sex, joint, previous knowledge of acupuncture, credibility, CARE score, and HCAMQ scores). The ANCOVA was repeated with the insignificant covariates removed. Group differences are shown in Table 6. Patients who thought that their treatment was real recorded pain scores 11.5 mm (95% CI 3.4 to 19.5, P = .005) lower than those who did not. Treatment type and consultation type had no significant effect on posttreatment pain (Table 6). There was a significant practitioner effect. Practitioner 3 achieved greater analgesia than practitioner 2 (mean difference 10.9 mm, 95% CI 3.9 to 18.0, P = .002) across all treatments and consultation types. This also neared significance against practitioner 1 (mean difference 5.8 mm, 95% CI –1.1 to 11.7, P = .05).

3.3. Secondary outcomes

Pain over time decreased for all treatments (Fig. 2). This was assessed using a repeated-measures ANCOVA and showed that there was a highly significant downward trend in pain scores over time, with an average reduction of 3.4 mm in pain scores per week on the VAS scales (P < .001), but there was not a significant difference between the downward trends for the different treatments. All scores for Western Ontario and McMaster University Osteoarthritis Index and Nottingham Health Profile improved in all groups, with no significant differences between treatment or consultation types. Practitioner 3 achieved the largest improvements.

3.4. Blinding

A high percentage of patients believed that their treatment had been real; 96% (RA), 93% (SN), and 75% (mock electrical stimula-
3.5.2. The practitioner effect

seen practitioner 3 as more authoritative than practitioner 1. “Doctor,” was never referred to by his first name, and was typically affectionately referred to as “girl” and a “young lady,” and some described her using affectionate terms such as “sweet.” Practitioner 3 was referred to as a “girl” and a “young lady,” and some described her using affectionate terms. One difference was identified: terminological differences between treatments were significant ($\chi^2 = 15.486, P = .001$). Three adverse events were recorded, none of which were related to treatment. There were 28 cases of minor side effects (Table 7).

3.5. Qualitative findings

The qualitative analysis provides insight into how perceptions of treatment veracity influenced outcomes, why practitioner 3 was more effective, and why consultation type had no effect on posttreatment pain.

3.5.1. Treatment veracity

Interviewees expressed uncertainty regarding the veracity of their treatment, which was accompanied by uncertainty regarding subjective outcomes. Beliefs about treatment veracity and confidence in outcomes were reciprocally linked. As people became more confident that they were experiencing benefits, such as decreased pain, they also became more confident that they were receiving real acupuncture. As they became more confident that they were receiving real acupuncture, they also became more confident that they were experiencing positive health changes. Confidence in health changes in turn impacted how participants perceived outcomes between the different consultation styles. In this trial finding Deductive framework, no effect of consultation type, and effect of perceiving RA vs SN. Treatment Condition (MES vs RA vs SN). These differences between treatments were significant ($\chi^2 = 15.486, P = .001$). Three adverse events were recorded, none of which were related to treatment. There were 28 cases of minor side effects (Table 7).

Table 3

<table>
<thead>
<tr>
<th>Framework analysis</th>
<th>Inductive thematic analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>RCT finding</td>
<td>Deductive framework</td>
</tr>
<tr>
<td>No effect of intervention condition, and effect of perceiving treatment to be real</td>
<td>Outcomes</td>
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<tr>
<td>Effect of practitioner</td>
<td>Treatment characteristics</td>
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<tr>
<td>No effect of consultation type</td>
<td>Practitioners</td>
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<td>No effect of consultation type</td>
<td>Inductive themes</td>
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<td></td>
<td>Comparisons</td>
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</tbody>
</table>

EMP, empathic consultation; MES, mock electrical stimulation; RA, real acupuncture; RCT, randomized controlled trial; SN, Streitberger needle.

3.5.3. Consultation type

Participants in empathic consultations described practitioners as caring, friendly, and communicative. Those in nonempathic consultations undertook a little more work to explain their similarly positive views of their practitioners. Thus interviewees who had received nonempathic consultations talked about how they concluded with the practitioner to obey the study rules and have limited personal interactions. They suggested that the practitioners were not really nonempathic, they were just acting that way for the sake of the trial. For example, “I had the feeling that she sort of felt that you know, not being able to converse properly, that she felt a bit awkward about it” (Betty, nonempathic). Other aspects of the trial beyond the consultation (friendly and polite reception staff, convenient appointments) were also described positively as evidence that the researchers and practitioners did actually care about the interviewees and so the interviewees described feeling “cared for” by all those involved in the study.

Dorothy was the only interviewee to describe her experiences of being in the trial in general and her experiences of her (nonempathic) practitioner. In negative terms: “I don’t think that the environment was conducive to her work when she was doing her job.” Dorothy described not knowing that her practitioner’s behaviour was part of the trial (despite being fully informed at the recruitment interview) and was also the interviewee for whom the nonempathic consultation could be described as most successful. If she had recalled her recruitment interview, the nonempathic consultation may have lost its potency: “If I had this young lady and I’d known that some of you were going to be just businesslike and some are going to be friendly, I would have thought I had got one of the businesslike ones and brushed it off. But not to know was I think, wrong.”

4. Discussion

All groups improved from baseline, with many achieving clinically significant improvements. There were no differences between verum and placebo treatments. The empathic nature of the consultation process had no effect on outcome, but believing in the treatment and a powerful practitioner effect significantly affected pain at 5 weeks. All primary and secondary outcomes showed similar trends. The qualitative study helps to explain these findings, suggesting that beliefs about treatment veracity and confidence in outcomes were reciprocally linked and the supportive nature of the study and consent process may have attenuated differences in outcomes between the different consultation styles. In this
A type II error is unlikely; there was underrecruitment but dropouts were approximately half that anticipated, so the main-effect comparisons were adequately powered. Using the smallest groups, the absolute difference between the mean pain improvement for practitioners 2 and 3 was 13 mm, which is less than our predefined clinically important difference of 20 mm, and corresponds to 80% power at 1% significance. Other main comparisons were based on much larger group sizes. Type I error is possible for the practitioner effect, but the level of significance and the supportive findings from the qualitative study make this unlikely.

This study did not employ a no-treatment arm, and therefore it is possible that improvements recorded might simply be due to the placebo effect.
natural history of the condition, i.e., leading to spontaneous remission (a regression to the mean). The effect of this cannot be measured from this study, and we cannot discount this possibility. However, this possibility might be attenuated by the population chosen for this study, i.e., all were chronic end-stage OA sufferers such that all were awaiting joint replacement. Conservative treatment options for this group had been exhausted and pain was still persistent and consistently high enough to warrant surgical intervention.

4.1. The effect of the consultation process and treatment context

Manipulating the consultation process was effective quantitatively and resulted in significantly different CARE scores across the groups. Mercer [18] suggest that small differences in CARE represent large differences in real terms because patients are reluctant to give very low scores to their doctors. The 10-point difference found in this trial is therefore substantial, reflecting clinically important differences that may be much larger differences in observer rated scores [18]. This did not result in a different therapeutic outcome in spite of the acknowledged importance of a patient-centred approach [17]. It is possible that CARE is failing to adequately capture patient centeredness [17], but the qualitative study suggests that this is not the case. It seems that patients appear to have made allowances and colluded when receiving non-empathic consultations, which might have subverted the trial process. They drew empathy from other sources (reception staff, the initial prerandomisation consent/screening session), thus implying it may be very difficult to deliver a nonempathic process to patients in a trial in which patients have received prior informed consent. Researchers and ethics committees should consider this carefully in relation to consent and subsequent trial debriefing.

The other more fundamental issue is that we do not yet fully understand the relationship between verbal and nonverbal communication in the consultation with respect to patient centeredness and therapeutic outcome [17]. Interpretations of outcome related to empathy should therefore be approached with great caution.

Table 5
Mean pretreatment and posttreatment scores for prime outcome by group.

<table>
<thead>
<tr>
<th>Treatment</th>
<th>RA</th>
<th>SN</th>
<th>MES</th>
<th>Practitioner 1</th>
<th>Practitioner 2</th>
<th>Practitioner 3</th>
<th>EMP</th>
<th>Non-EMP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Baseline pain VAS (SD)</td>
<td>60.5 (14.2)</td>
<td>58.6 (14.6)</td>
<td>58.3 (16.0)</td>
<td>60.3 (15.6)</td>
<td>59.6 (13.3)</td>
<td>56.6 (14.9)</td>
<td>59.1 (14.9)</td>
<td>59.2 (15.0)</td>
</tr>
<tr>
<td>Posttreatment pain VAS (SD)</td>
<td>43.5 (25.5)</td>
<td>44.0 (21.7)</td>
<td>49.2 (25.7)</td>
<td>47.0 (24.8)</td>
<td>53.2 (23.0)</td>
<td>37.0 (22.7)</td>
<td>47.4 (25.3)</td>
<td>43.6 (23.4)</td>
</tr>
<tr>
<td>Mean pain improvement (% of patients with clinically relevant improvement in pain)</td>
<td>29.5 (61.9)</td>
<td>23.0 (62.3)</td>
<td>16.6 (41.6)</td>
<td>22.2 (55.2)</td>
<td>10.3 (45.8)</td>
<td>34.9 (62.7)</td>
<td>21.1 (52.3)</td>
<td>25.2 (58.4)</td>
</tr>
<tr>
<td>Baseline analgesia (SD)</td>
<td>4.6 (3.9)</td>
<td>4.4 (3.6)</td>
<td>4.0 (3.74)</td>
<td>4.6 (4.0)</td>
<td>4.8 (3.7)</td>
<td>3.4 (3.2)</td>
<td>4.6 (4.3)</td>
<td>4.1 (3.0)</td>
</tr>
<tr>
<td>Posttreatment analgesia (SD)</td>
<td>3.8 (4.6)</td>
<td>4.2 (6.9)</td>
<td>3.7 (4.1)</td>
<td>4.4 (6.6)</td>
<td>4.2 (3.9)</td>
<td>2.7 (3.3)</td>
<td>3.8 (4.6)</td>
<td>4.0 (6.0)</td>
</tr>
</tbody>
</table>

Table 6
Adjusted means and differences for pain at week 5 from the analysis of covariance.

<table>
<thead>
<tr>
<th>Treatments</th>
<th>Mean</th>
<th>Standard error</th>
<th>95% confidence interval</th>
<th>Lower bound</th>
<th>Upper bound</th>
<th>P value</th>
</tr>
</thead>
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<tr>
<td>RA</td>
<td>43.5</td>
<td>2.2</td>
<td>39.2</td>
<td>47.8</td>
<td></td>
<td></td>
</tr>
<tr>
<td>SN</td>
<td>46.2</td>
<td>2.3</td>
<td>41.6</td>
<td>50.8</td>
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<td></td>
</tr>
<tr>
<td>SE</td>
<td>47.3</td>
<td>2.4</td>
<td>42.6</td>
<td>52.1</td>
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<tr>
<td>Differences</td>
<td>Mean</td>
<td>Standard error</td>
<td>Lower bound</td>
<td>Upper bound</td>
<td>P value</td>
<td></td>
</tr>
<tr>
<td>RA–SN</td>
<td>–2.7</td>
<td>3.2</td>
<td>–9.0</td>
<td>3.6</td>
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<tr>
<td>RA–SE</td>
<td>–3.9</td>
<td>3.3</td>
<td>–10.4</td>
<td>2.7</td>
<td>.25</td>
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<tr>
<td>SN–SE</td>
<td>–1.2</td>
<td>3.4</td>
<td>–7.8</td>
<td>5.5</td>
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<tr>
<td>Practitioner</td>
<td>Mean</td>
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<td>95% confidence interval</td>
<td>Lower bound</td>
<td>Upper bound</td>
<td>P value</td>
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<tr>
<td>1</td>
<td>45.9</td>
<td>1.8</td>
<td>42.4</td>
<td>49.430</td>
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<tr>
<td>2</td>
<td>51.0</td>
<td>2.6</td>
<td>45.8</td>
<td>56.201</td>
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<tr>
<td>3</td>
<td>40.1</td>
<td>2.4</td>
<td>35.4</td>
<td>44.745</td>
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<tr>
<td>Differences</td>
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<td>Upper bound</td>
<td>P value</td>
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<tr>
<td>1–2</td>
<td>–5.1</td>
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<td>1–3</td>
<td>5.8</td>
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<td>–0.1</td>
<td>11.7</td>
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<td>2–3</td>
<td>10.9</td>
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<td>1.8</td>
<td>43.7</td>
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<td>ME</td>
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<td>2.0</td>
<td>40.3</td>
<td>48.0</td>
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<tr>
<td>Difference</td>
<td>Mean</td>
<td>Standard error</td>
<td>Lower bound</td>
<td>Upper bound</td>
<td>P value</td>
<td></td>
</tr>
<tr>
<td>EM–ME</td>
<td>3.0</td>
<td>2.6</td>
<td>–2.2</td>
<td>8.2</td>
<td>.26</td>
<td></td>
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</table>

Covariates appearing in the model are evaluated at the following values: mean baseline pain = 58.86, baseline mean daily tablet count = 4.32, real = 0.88, posttreatment mean daily tablet count = 3.64.

EM, empathic; ME, minimal empathy; RA, real acupuncture; SN, Streitberger needle; VAS, visual analogue score.
Kaptchuk et al. suggest that enhanced consultations produced better outcomes [14]. The differences between our studies could relate to the diagnoses (OA pain vs irritable bowel syndrome) or to the differences in the initial consent process (Kaptchuk et al. did not require consent to different consultation types, whereas we did). We only manipulated empathy, and Kaptchuk et al. manipulated the acupuncture treatment (palpation), consultation time, and therapists’ confidence in treatment. There may also be an interaction between different components within acupuncture treatment (talking, listening, and diagnosis) that produce a clinical effect that is greater than the sum of its individual elements. Patterson and Dieppe [21] suggest that these factors, in contrast to drug trials, are integral to complex nonpharmaceutical interventions.

Belief in treatment veracity was significantly correlated to outcome, suggesting that improving the management of expectation could enhance outcome. This is supported by the qualitative analysis, which suggests that patients link positive outcomes and recorded improved symptoms when they believed that the treatment had been real and vice versa. Where patients were unsure as to the “reality” of their treatment, this was manifest in a reluctance to score symptom improvement, and therefore they were less willing to commit themselves when completing the outcome questionnaires. This has implications for all placebo-controlled trials insofar as the real effects of an unconvincing verum treatment might be underreported (a type II error). One solution might be to encourage participants to understand and believe (correctly) that placebos are powerful interventions that act through a variety of complex sociocultural, psychological, and neurological mechanisms [23]. They have a proven range of beneficial effects, including pain relief. This might encourage participants to report perceived improvements confidently despite being uncertain as to their treatment allocation. It might also minimise the distress that some participants experience on unblinding when discovering they have responded to a placebo [4] and perceive this as something deceitful.

4.2. Practitioner effect

Practitioner 3 produced better outcomes across all treatment and consultation types. This occurred in spite of the meticulous care and planning taken to ensure consistency of treatment delivery among the 3 practitioners. The qualitative data suggested that the interviewees perceived practitioner 3 as a paternalistic male authority figure. Practitioner 3, as the primary investigator, might have been seen by patients as the expert, consequently establishing higher expectations of success, which in turn influenced outcome. Although this is consistent with previous research [10,29] a larger explanatory study involving many practitioners is needed.

4.3. Control type

It has been suggested that that needling carries an enhanced placebo effect [14,15]. We are the first to compare 2 acupuncture placebo controls demonstrating no difference between needle and nonneedle placebos. The magnitude of the improvements shown in the placebo arms in this and other studies indicates that placebo acupuncture has powerful effects outperforming standard care [12,25]. This supports the hypothesis that “exotic” processes or rituals may have powerful therapeutic effects.

4.4. Efficacy of acupuncture

Efficacy studies of acupuncture for OA pain have not reported consistent results. Although our results concur those of with Scharf et al. [25], other large studies demonstrate a statistically significant effect for verum acupuncture in knee pain [2,26,28]. The reasons for these differences are unclear. We included patients with hip pain, and this might not respond as well to acupuncture; it is also possible that acupuncture is much less effective in this group of severely affected OA patients.

4.5. Conclusions

This trial uniquely reports a mixed-methods approach in a single report to enable a contextual interpretation of the quantitative data. Patients receiving acupuncture demonstrated clinically important improvements from baseline (i.e., a 29.5% reduction in pain), but despite this, acupuncture has no specific efficacy over placebo for this group of patients. The clinical effect of acupuncture treatment and associated controls is not related to the use of an acupuncture needle, nor mediated by empathy, but is practitioner related and may be linked to the perceived authority of the practitioner. Qualitative analysis indicated that beliefs about treatment veracity and confidence in outcomes were reciprocally linked, and this appears to affect how patients self-report treatment outcomes, leading to a greater risk of demonstrating type II errors. This suggests that the process of interpreting RCT data is complex and contextual.

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