A double blind randomised trial of low power laser treatment in rheumatoid arthritis

J K Heussler, G Hinchey, E Margiotta, R Quinn, P Butler, J Martin, A D Sturgess

Abstract

Objectives—To define the value of low power laser treatment in small joint rheumatoid arthritis.

Methods—Twenty five women with active disease were recruited. The metacarpophalangeal and proximal interphalangeal joints of one hand were treated with 12 J/cm² for 30 s with a gallium–aluminium–arsenate laser. The other hand received a sham laser treatment designed so that neither therapist nor patient could distinguish the active laser from the sham laser. Each patient received 12 treatments over four weeks. The following parameters were measured: pain as assessed by visual analogue scale; range of joint movements; grip strength; duration of early morning stiffness, joint circumference, Jębsen’s hand assessment; drug usage; total swollen joint counts; Arthritis Impact Measurement Scales; three phase bone scans; haematological and serological tests.

Results—A total of 72% of patients reported pain relief but this reduction was reported equally in both hands. No significant changes were seen in other clinical, functional, scintigraphic, or laboratory features. Neither patients nor staff were able to detect which hand was treated with the active laser.

Conclusion—When this specific laser and dose regimen was used, low power laser treatment had no objective effect on patients with rheumatoid arthritis. It did appear to produce analgesia through a powerful placebo effect.

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Low power laser treatment is increasingly used as a modality of physical therapy in the management of a variety of musculoskeletal diseases, including rheumatoid arthritis (RA).1–11 A variety of laser sources are used, such as gallium–aluminium–arsenate, helium–neon, gallium–arsenate, with power outputs of 50 mW or less. No thermal energy is delivered, and the laser may be within the visible light spectrum or invisible.

In patients with RA, previous workers have reported that laser treatment produces both local and systemic changes, including decreased joint pain, decreased joint swelling, increased functional activity, decreased duration of early morning stiffness, and improved laboratory parameters including erythrocyte sedimentation rate.1 3 No mechanism of action has been defined but photochemical reactions, particularly in the mitochondria, have been suggested.12–13 The optimal wavelength and dosage for the treatment of RA is unknown as, depending on the experimental system, either inhibition or stimulation of selected cell functions has been reported.14 A wide range of differing wavelengths and doses has been used in published reports making it difficult to compare studies (table).

In our hospital, treatment with a gallium–aluminium–arsenate laser appeared to produce significant benefit in patients with active RA in the hands. We therefore designed a double blind trial to investigate local changes in the hands of patients with RA, and to document any systemic changes. A protocol employing sham laser treatment to one hand was preferred rather than the use of a non-treatment group, so that each patient had an opportunity to benefit from active treatment, and each patient could compare changes in their two hands.

Patients and methods

Twenty eight women with RA and bilateral involvement of their metacarpophalangeal and proximal interphalangeal joints were selected from the St George’s Hospital rheumatology outpatients department, and by means of a small notice in the local paper. Patients were excluded if they were having an acute inflammatory episode, had arthropathies in either joint, were pregnant, or had evidence of digital vasculitis. Each patient was randomly allocated to one of two groups. Group 1 received active laser treatment to their right hand, whereas the left hand received sham laser treatment. Group 2 were treated in the reverse manner.

Of the 28 selected, three withdrew from the study: one because of an acute inflammatory episode and social reasons, another for social reasons alone, and the third because of admission to hospital for a non-related illness and surgery. Only the 25 patients who completed the trial were included in the analysis, giving 25 hands treated with active
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<td>2</td>
<td>Nd-YAG</td>
<td>λ=1060 nm; E=15 J/cm²; T=30 s</td>
<td>31 patients, laser to MCP/PIP of one hand. Sham laser to other hand. Once weekly for 10 weeks</td>
<td>Hand function</td>
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<td>λ=633 nm; E=6 J/cm²; P=10 mW; T=300 s</td>
<td>17 patients, laser to MCP of one hand. Sham laser to 2nd MCP of other hand. Thrice weekly for 3 weeks</td>
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<td>3</td>
<td>Ga-Al-As</td>
<td>λ=820 nm; E=3-5 J/cm²; P=15 mW; T=60 s</td>
<td>35 patients, laser to MCP of one hand. Sham laser to control group. Thrice weekly for 4 weeks</td>
<td>Hand function</td>
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<td>22</td>
<td>Helium-neon</td>
<td>λ=633 nm; E=3-6 J/cm²; P=5 mW; T=30-480 s</td>
<td>36 patients, laser to one group. Sham laser to a second group. Thrice weekly for 10 weeks with escalating treatment doses</td>
<td>Pain (VAS)</td>
</tr>
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λ= wavelength; Ga-Al-As=gallium-aluminium-arsenate; MCP=metakarposphalangeal joint; PIP=proximal interphalangeal joint; VAS=visual analogue scale.

Laser and 25 hands treated with sham laser. All patients were right hand dominant. The two groups were well matched, with group 1 patients having a mean age of 64.8 years (43-77) and a disease duration of 14.2 years (5-30), and group 2 patients having a mean age 62.5 years (40-80) and a disease duration 12.7 years (5-30).

A course of 12 laser treatments was given over a four week period. The active and sham laser probes were identical in external appearance except for a code label, and no visible light, heat, or vibration was detectable from either the active or the placebo probe. The code was kept secret by an uninvolved staff member until all data were analysed.

The treatment was delivered by means of medial and lateral approaches and with the joint held in a loose packed position—that is, 45° of flexion or neutral position. The treated joints were the metakarposphalangeal, proximal interphalangeal, and the interphalangeal joint of the thumb. A total of 12 J/cm² was delivered to each joint via a gallium–aluminium–arsenate laser with the following operating parameters: average power 50 mW; beam spot size 0.126 cm²; pulse width 0.16 ms; pulse frequency 5000 Hz; and wavelength λ 820 nm. The dosage was decided upon empirically after our physiotherapy staff had noted improvements after treatment of patients with this protocol. All treatments were conducted according to the Australian Physiotherapy Association standards, and the techniques and dosages were checked by the St George Hospital Laser Safety Officer. All patients signed an informed consent form and ethical approval was given by the area health service ethics committee.

Weekly measurements were taken, consisting of visual analogue scales used for each hand to assess pain; the duration of early morning stiffness in each hand; joint circumference via a Fröben meter; range of motion for each treated joint via a finger goniometer; grip strength (gross: Jamar hand dynamometer, pinch: pinch gauge, tripod: pinch gauge), index to thumb distance (span) via a goniometer ruler, and the pulp tip to the distal palmar crease distance via a goniometer ruler. In addition, before and after the four week treatment period, the following assessments were done: Jeberries hand assessment; drug usage; total hand swollen joint counts; Arthritis Impact Measurement Scales (AIMS); three phase bone scans and haematological and serological tests.

Four indices of the AIMS were selected for evaluation: pain; dexterity; activities of daily living; and household activities. In accordance with recommended guidelines, a significant change in the AIMS was said to occur when three out of four indices had demonstrated a change of greater than 30%.

Haemoglobin, erythrocyte sedimentation rate, differential white cell count, and C reactive protein (CRP) were assessed by automated counters, latex agglutination, and flow cytometry using standard techniques.

**Results**

**Bone Scintigraphy**

Patients were scanned one day before and two days after treatment. On both occasions, patients were injected with 800–1000 MBq of technetium-99m labelled methylene disphosphonate (Radpharm, Australia) with blood pool images taken two to four minutes after injection and delayed images taken three hours after injection. Palmar and dorsal views of the hands were obtained in both blood pool and delayed studies. Imaging was performed using a gammacamera with a small field of view connected to a dedicated nuclear medicine computer (GE Starcam, Milwaukee, WI, United States) with 256x256 matrix acquisition, low energy, all purpose collimator, and 10 minutes of acquisition time per image. All joints in each hand were scored on both blood pool and delayed studies by two nuclear medicine physicians. The scoring system was –1 for decreased joint activity, 0 for normal activity, and +1 for increased activity. Differences between observers were reviewed and a consensus decision was recorded.

The scans were scored in random order by physicians who were unaware of whether they were pretreatment or posttreatment or which hand had received active treatment.

For statistical analysis, the changes in the treated hands were compared with the changes in the sham treated hands, using Wilcoxon sign rank tests. Correction for multiple measures indicated that a p value of <0.003 was required for statistical significance.

**Clinical Parameters**

Eighteen of 25 patients (72%) reported improvement in pain, but pain (as measured by the visual analogue scale) was reduced in both treated and sham treated hands after laser intervention (p<0.001 for comparison before and after treatment). The figure shows the mean values for pain per week, where weeks 0 and 5 are assessment only. There was no difference in the extent of pain relief in the treated hands and the sham treated hands. To assess the patient's ability to discriminate between active laser and placebo they were
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Each patient assessed pain in each hand separately using a visual analogue scale (VAS) from 0 to 10. The mean of these values is shown. Pain assessment fell in both the treated hand and the sham treated hand. There was no difference between the degree of improvement in the two groups.

asked to identify which, if either, hand they felt had improved with treatment. Only five of 25 (20%) identified the treated hand as the one that had improved, whereas five thought that the sham treated hand had improved, eight thought both hands had improved equally, six noted no change in either hand, and one reported that both hands had worsened. Despite the reported lessening of pain, drugs for pain relief did not change.

FUNCTIONAL HAND ASSESSMENTS
There were no significant differences between the treated hand and the sham treated hand over the period of the trial in the duration of early morning stiffness, total swollen joint count, the joint circumference index, range of motion, pulp tip to distal palmar crease distance, grip strength measures, or the Jelenk Activities Index.

LABORATORY AND SCINTIGRAPHIC PARAMETERS
There were no significant haematological changes, and no significant differences in blood pool or the delayed images of bone scans were identified.

SIDE EFFECTS
No significant side effects were seen, although nausea was noted occasionally in the initial treatments.

Discussion
Compared with the large number of reported drug trials in rheumatoid arthritis, there are relatively few controlled clinical trials validating the use of physical or electrophysical agents. Despite this, such agents as ultrasound, interferential treatment, shortwave diathermy, and low power laser treatment are widely used. Baxter et al. surveyed 397 physiotherapy practices in Northern Ireland, and found 54-2% were using laser treatment, emphasising how widespread use of this technique has become. Previous published studies of the efficacy of laser treatment in RA have yielded conflicting results, and we therefore designed this double blind trial to document any clinical or laboratory changes. We expected a strong placebo effect from the intensive contact with the therapist, combined with the 'high tech' appearance of the laser equipment, and we were therefore careful to ensure that both patient and therapist were ignorant of which treatment was being given.

The impressive change was the reduction in pain which was reported by 18/25 (72%) subjects, but these changes were reported equally in both hands. Laboratory studies were unchanged making it unlikely that a systemic effect from laser treatment of one hand could have reduced pain in the other. Moreover, detailed function measures, including functional assessments of each hand, showed no changes in either the treated or sham treated hands, despite the reported reduction in pain. A placebo effect is clearly the most likely explanation for the reported analysis. Depending on the type of intervention, placebo responses varying from 6-2% to 80% have been described.

Typically, uncontrolled studies of laser treatment in RA have claimed dramatic improvement with pain reduction of more than 90%. One such paper conceded, however, that measures which were less placebo-sensitive, such as range of movement, 'did not always show rates of improvement similar to effectiveness of pain removal'. At least four double blind trials have been previously published. In each study a strong placebo effect was noted, with improvement in sham treated joints, usually in pain measurement, but sometimes in strength and range of movement. Palmgren et al. used the same type of laser as in our study but the dose (3.6 J/cm²) was slightly smaller. They showed significant improvements in grip strength, movement, joint swelling, and the duration of early morning stiffness compared with control patients who had sham treatment. Pain improved in both their patients’ hands, but more so in their treated hands. Both erythrocyte sedimentation rate and total leucocyte counts fell with a significant trend. In Palmgren's study half the patients received active laser treatment and in half, placebo laser was used. We treated one hand and sham treated the other, and it is possible that, in our study, some systemic effect occurred that 'flowed over' from the actively treated to the placebo treated hand. Arguing against this possibility was the lack of change in erythrocyte sedimentation rate, or any other parameter measured. In addition, we found no change in any objective measurement or functional assessment. Despite the differences in design and dosage, it is difficult to reconcile our negative results with Palmgren's dramatic improvements.

Bliddal et al. and Goldman et al. used the same design as our study, but with different lasers. Bliddal's helium–neon laser produced a burning sensation in the actively treated joints of some patients, which would invalidate the blind element of the trial. Even so, less than half their patients reported pain relief, and there was no change in duration of early morning stiffness or joint range of movement.
No changes in laboratory parameters were found. The authors concluded that laser treatment was of limited value.

Goldman’s study used a Q-switch neodymium laser which delivers pulsatile irradiation at a fairly high output (15 J/cm² in 30 ns). All patients reported stinging from the laser, and more than 10% reported erythema or hyperpigmentation at the actively treated joints, invalidating efforts at making the trial blind. Both the treated and sham treated hands showed improvement for a range of measures: heat; erythema; pain; swelling; and tenderness. Erythema and pain were reported to have improved more in the treated hand than the placebo treated hand. Grasp and pinch strength were reported as more objective measures of improvement in the treated hands, but the probability values were not corrected for multiple measures and were, in each case, p=0.02 and therefore unlikely to be significant. Similarly, the reported change in platelet aggregation (p=0.025), which was proposed to indicate a systemic effect, would not be significant if the rate for each experiment was considered.

Walker et al reported significant pain reductions only in the treated patients, but this required nine weeks of treatment. Conceivably our actively treated patients might have improved to a significant degree with more prolonged treatment. Photochemical reactions, particularly in vitamin D metabolism, melanin, and in the retina, are well described and there seems to be no a priori reason why laser treatment could not exert an effect on one or more of the many pathways that lead to pain and inflammation in the arthritic hand. We found no objective benefit with the dose regimen we used, but cannot exclude the possibility of efficacy with another regimen. The gallium–aluminium–arsenate lasers are safe, and have been efficacious in at least one well controlled trial, and it therefore seems prudent to use such lasers in further trials. We have demonstrated the absolute necessity of placebo controls in low power laser treatment research.

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