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Injection with methylprednisolone proximal to the carpal tunnel: randomised double blind trial

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Abstract

Objective To assess the effect of a 40 mg methylprednisolone injection proximal to the carpal tunnel in patients with the carpal tunnel syndrome.

Design Randomised double blind placebo controlled trial.

Setting Outpatient neurology clinic in a district general hospital.

Participants Patients with symptoms of the carpal tunnel syndrome for more than 3 months, confirmed by electrophysiological tests and aged over 18 years.

Intervention Injection with 10 mg lignocaine (lidocaine) or 10 mg lignocaine and 40 mg methylprednisolone. Non-responders who had received lignocaine received 40 mg methylprednisolone and 10 mg lignocaine and were followed in an open study.

Main outcome measures Participants were scored as having improved or not improved. Improved was defined as no symptoms or minor symptoms requiring no further treatment.

Results At 1 month 6 (20%) of 30 patients in the control group had improved compared with 23 (77%) of 30 patients the intervention group (difference 57% (95% confidence interval 36% to 77%). After 1 year, 2 of 6 improved patients in the control group did not need a second treatment, compared with 15 of 23 improved patients in the intervention group (difference 43% (23%) to 63%). Of the 28 non-responders in the control group, 24 (86%) improved after methylprednisolone. Of these 24 patients, 12 needed surgical treatment within one year.

Conclusion A single injection with steroids close to the carpal tunnel may result in long term improvement and should be considered before surgical decompression.

Introduction

The carpal tunnel syndrome is caused by compression of the median nerve at the wrist and is a common cause of pain in the arm, particularly in women. Injection with corticosteroids is one of the many recommended treatments.

One of the techniques for such injection entails injection just proximal to (not into) the carpal tunnel. The rationale for this injection site is that there is often a swelling at the volar side of the forearm, close to the carpal tunnel, which might contribute to compression of the median nerve. Moreover, the risk of damaging the median nerve by injection at this site is lower than by injection into the narrow carpal tunnel. The rationale for using lignocaine (lidocaine) together with corticosteroids is twofold: the injection is painless, and diminished sensation afterwards shows that the injection was properly carried out.

We investigated in a double blind randomised trial, firstly, whether symptoms disappeared after injection with corticosteroids proximal to the carpal tunnel and, secondly, how many patients remained free of symptoms at follow up after this treatment.

Participants and methods

Participants

The participants were patients referred to the Medical Centre Alkmaar with signs and symptoms of the carpal tunnel syndrome of more than 3 months' duration confirmed by electrophysiological tests. In those with bilateral symptoms, the arm with the most severe symptoms was chosen, and treatment of this arm was randomised. We excluded patients aged under 18 years or patients who had already been treated for symptoms of the carpal tunnel syndrome.
The trial was approved by the medical centre’s ethics committee. Patients gave written informed consent. The ethics committee required an interim analysis after inclusion of half of all participants.

**Intervention**

The injections were given by one neurologist (JWHHD). They contained 10 mg lignocaine or 10 mg lignocaine and 40 mg methylprednisolone. The site of injection was at the volar side of the forearm 4 cm proximal to the wrist crease between the tendons of the radial flexor muscle and the long palmar muscle. Injections were given with a 3 cm long 0.7 mm needle (fig 1). The angle of introduction of the needle depended on the size of the wrist. In participants with a thin wrist the median nerve is close to the skin. In these participants the angle was 10°. The angle was larger, about 20°, in those with a thick wrist. In participants with well developed muscles, the pronator quadratus muscle may push up the median nerve, so in a thick muscular arm the angle of introduction was also flat, between 10° and 20°. The needle was introduced slowly, and the participant was instructed to say stop if he or she felt pins and needles or pain in the fingers. If a resistance was felt the needle was withdrawn a few millimetres then repositioned. The injection was given without much pressure. After injection, the 1 ml fluid bolus was gently massaged towards the carpal tunnel.

**Outcome**

One month after injection the randomised participants visited the outpatient department and were asked by another neurologist (MMV) whether they had no symptoms or only minor symptoms that they considered so much improved that they felt no further treatment was necessary. Further visits were planned for 3, 6, 9, and 12 months after the injection or earlier if the participant felt this was necessary. At these visits, participants were asked the same question. If treatment was necessary the decision to treat was taken first, and then the trial code was broken. If a patient had not been treated with methylprednisolone this treatment was offered, otherwise surgical decompression was performed.

**Assignment and blinding**

Using a random number table, the hospital pharmacist prepared the trial drug in blocks of 20. The syringes for injection were sent from the pharmacy to the outpatient department, where it was impossible to distinguish the syringes containing methylprednisolone plus lignocaine from those containing lignocaine as paper was glued around the syringes. To further ensure blinding, the assessments were carried out by another neurologist (MMV). Neither the doctor nor the participant, therefore, knew what treatment was given. The doctors and participants remained blind to treatment during the assessments at follow up.

**Sample size**

The sample size calculation was based on the assumption that after 1 month 80% of the participants in the intervention group would respond to treatment versus 50% in the control group. With a power of 80% and a significance level of 5% two sided, this meant that at least 80 participants needed to be included.

**Results**

After the ethics committee had seen the results of the interim analysis (after 40 participants had been recruited) it withdrew permission for further randomisation. Meanwhile a further 20 participants had entered the study. The final analysis of the results is on all 60 randomised patients. None of the participants was lost at follow up. Figure 2 shows the participant flow.

Table 1 shows the baseline characteristics. No significant differences existed between the groups. After 1 month 23 of the 30 participants in the intervention

**Table 1** Baseline characteristics of 60 participants in trial

<table>
<thead>
<tr>
<th>Variable</th>
<th>Intervention</th>
<th>Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>No of participants</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Mean age (years)</td>
<td>52</td>
<td>51</td>
</tr>
<tr>
<td>No of females</td>
<td>24</td>
<td>26</td>
</tr>
<tr>
<td>No of participants with pain in arm at night</td>
<td>27</td>
<td>26</td>
</tr>
<tr>
<td>No of participants with swelling near carpal tunnel</td>
<td>19</td>
<td>26</td>
</tr>
<tr>
<td>Average duration of symptoms (months)</td>
<td>32</td>
<td>25</td>
</tr>
</tbody>
</table>

**Table 2** Treatment response at follow up

<table>
<thead>
<tr>
<th>Period after treatment</th>
<th>Intervention group (n=30)</th>
<th>Control group (n=30)</th>
<th>% observed difference (95% confidence interval of difference)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 month</td>
<td>23 (77)</td>
<td>6 (20)</td>
<td>57 (36 to 77)</td>
</tr>
<tr>
<td>3 months</td>
<td>19 (63)</td>
<td>2 (7)</td>
<td>56 (37 to 76)</td>
</tr>
<tr>
<td>6 months</td>
<td>17 (57)</td>
<td>2 (7)</td>
<td>50 (30 to 70)</td>
</tr>
<tr>
<td>9 months</td>
<td>16 (53)</td>
<td>2 (7)</td>
<td>46 (27 to 67)</td>
</tr>
<tr>
<td>12 months</td>
<td>15 (50)</td>
<td>2 (7)</td>
<td>43 (23 to 63)</td>
</tr>
</tbody>
</table>
Street noises

In this “free” country everyone seems to be at perfect liberty to make whatever discordant noise he likes in the streets at any time of day or night. Vendors of milk, of cat’s meat, and of newspapers, German bands and barrel organs, all do their utmost to outvie one another in the creation of the city’s unnecessary turmoil, which is highly detrimental to everyone engaged in brain work, to the sick and weary, and to all who sleep lightly. The report suggests that indiarubber tyres should be adapted to many light vehicles, especially mail carts and milk carts, and that rubber rings should be compulsorily placed on the floors of milk carts to prevent the rattling of cans; that newspaper boys should desist from shouting, and instead thereof should exhibit “contents sheets,” or that newspapers shall be sold from kiosks on the pavements, as in Paris. In fact, street noises should be made the subject of reasonable regulations, and the people who make noises should be taught to be orderly and systematic. London is indeed far behind other places in the matter of by-laws for the good government of the public thoroughfares. In Eastbourne, for instance, only two barrel-organ grinders are allowed in the place, and they on sufferance; whilst the milkmen do not make a sound. In Croydon, Bath, Brighton, and the counties of Kent and Surrey no crying of goods or street music is permitted on Sunday. But although the metropolis is at present far in arrear in this matter there is prospect of amendment, as the London Government Act lately passed empowers the new municipalities to make by-laws for regulating and suppressing street noises and nuisances, and this act will come into force in November, 1900. (BMJ 1899;i:796)