High volume image-guided injections and structured rehabilitation improve greater trochanter pain syndrome in the short and medium term: a combined retrospective and prospective case series

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Summary

Background: the aim of this study was to measure the effects of high volume image-guided injections and structured rehabilitation (HVIGI&SR) for greater trochanter pain syndrome (GTPS).

Methods: 31 consecutive subjects were recruited (23 retrospectively; 8 prospectively) over 5 months. GTPS was diagnosed based on history and examination findings, alongside radiological examination. The HVIGI used a 22-gauge spinal needle to administer 10ml of 0.5% Marcaine and 50 mg hydrocortisone just deep to the periosteum underlying the gluteal tendon insertion under ultrasound guidance, followed by structured rehabilitation. A visual analogue scale (VAS) for pain was used as the main outcome measure.

Results: the mean VAS improved from 81.7 mm (±17.6) to 42.3 mm (±28.3), (p<0.05) in the prospective subjects at a mean of 6 weeks, considered clinically significant. In the retrospective subjects the mean VAS had improved from 74.6 (±10.9) mm to 38.2(±31.2) mm at two weeks (p<0.01) and 31.3 (±27.6) mm at the final time point, a mean of 60 weeks (p<0.01). The Hip and Groin Outcome Score in the prospective group showed a non-significant increase from 173.2 to 296.1 (p=0.12).

Conclusion: HVIGI&SR should be considered when short- and medium-term pain-relieving treatment for GTPS is required. Controlled studies are warranted to fully establish effectiveness, and assess long term effects.

Level of evidence: case series.

KEY WORDS: injection, greater trochanter pain syndrome, ultrasound, VAS scale.

Introduction

Of the hip complaints presenting to primary care, 10-20% are thought to be due to greater trochanter area pain¹. Within the US military an incidence of 2.03 per 1000 person-years was found, with a significantly greater incidence in women compared to men ². The causes of trochanteric pain are numerous including trochanteric bursitis, tendinopathy, muscle tears, iliotibial band disorders and bursinflammation³⁻⁵. As a result of this, and the difficulty in making a precise diagnosis, the term Greater Trochanteric Pain Syndrome (GTPS) has been implemented in clinical practice ⁶. The majority of patients with GTPS are managed conservatively and respond well to therapies that include progressive exercise and education ⁶. Corticosteroid injections are also commonly used as a non-operation option with reported responses ranging from 60-100%⁷. However the literature also suggests that the longevity of the corticosteroid injection effect is limited and is less effective than home training or shock wave therapy ⁸. Indeed the site of the corticosteroid injection may be important, as one study suggested that greater trochanteric bursa injections are superior to deep to gluteus medius bursa injections⁹. A recent review suggested that despite its common use in clinical practice there are very few studies on its effectiveness, and also minimal evidence comparing blind injections to image guided injections⁷.

Surgery is used in very recalcitrant cases, however the evidence base does not allow definitive conclusions about the best form of treatment for different presentations to be made with confidence⁶. Many people suffer recalcitrant and recurrent problems with significant impact on health¹⁰; indeed the effect of greater trochanteric pain syndrome on a patient’s life has been found by Fearon et al. to be similar in some respects to late stage hip osteoarthritis¹¹.
High volume image-guided injections with structured rehabilitation (HVIGI&SR) have been shown to be effective in improving pain and function for both Achilles and patellar tendinopathy\textsuperscript{12-14}. More recently they have been shown to be effective in the short term reduction of pain and improvement in function in shoulder impingement syndrome, another syndrome encompassing many diagnoses\textsuperscript{15}. The mechanism behind the effect of the HVIGI is not well understood but Chan et al.\textsuperscript{13} hypothesise that there is disruption of the neovascularisation seen in tendinopathy. In shoulder impingement syndrome, which encompasses other non-tendinopathy conditions. Morton et al.\textsuperscript{15} hypothesise that there is disruption of the scar tissue or separation of tissues, although it is recognised that more work is required to confirm this. Therefore, the aim of this research was to provide preliminary evidence to clinicians about the short- and medium-term treatment effectiveness of HVIGI&SR for GTPS, using both retrospective and prospective data.

### Materials and methods

#### Subjects

Every patient (n=8) attending one specialist MSK radiology clinic over a five month period who met the inclusion and exclusion criteria (Tab. 1) were recruited prospectively, completing a visual analogue scale (VAS) for pain and the Hip and Groin Outcome Score (HAGOS) at the appointment and then again at six weeks post-HVIGI&SR\textsuperscript{16}. Retrospective patients (n=23) were identified from a database covering the previous two years and sent the above questionnaires. Ethical clearance was obtained from Queen Mary University of London Ethics of Research Committee and consent was obtained via the questionnaire. The subjects’ characteristics can be seen in Table 2. The research followed the guidelines as laid out by Padulo et al.\textsuperscript{17}. Please see Appendix 1 for copies of the questionnaire (both retrospective and prospective).

### Table 1. Inclusion and Exclusion Criteria.

<table>
<thead>
<tr>
<th>Inclusion</th>
<th>Exclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>18-80 years</td>
<td>Another cause for hip pain is suspected</td>
</tr>
<tr>
<td>Clinical diagnosis of GTPS</td>
<td>Injection contraindicated e.g. allergy to the content</td>
</tr>
<tr>
<td>Local tenderness over greater trochanter</td>
<td></td>
</tr>
<tr>
<td>Radiological examination to rule out alternative causes for their lateral hip pain such as osteoarthritis</td>
<td></td>
</tr>
<tr>
<td>Recalcitrant to rehabilitation</td>
<td></td>
</tr>
</tbody>
</table>

### Table 2. Subject Characteristics.

<table>
<thead>
<tr>
<th></th>
<th>Retrospective subjects</th>
<th>Prospective subjects</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age range</strong></td>
<td>46-55</td>
<td>56-65</td>
</tr>
<tr>
<td><strong>Gender (M:F)</strong></td>
<td>2:12</td>
<td>3:4</td>
</tr>
<tr>
<td><strong>Level of sport/activity</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No activity:</td>
<td>4</td>
<td>0</td>
</tr>
<tr>
<td>Recreational:</td>
<td>7</td>
<td>5</td>
</tr>
<tr>
<td>Local club:</td>
<td>0</td>
<td>2</td>
</tr>
<tr>
<td>Elite national/international (amateur):</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Elite national/international (prof):</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><strong>Side of injection</strong></td>
<td>Right: 8</td>
<td>Right: 4</td>
</tr>
<tr>
<td></td>
<td>Left: 5</td>
<td>Left: 3</td>
</tr>
<tr>
<td></td>
<td>Bilateral: 1</td>
<td>Bilateral: 0</td>
</tr>
<tr>
<td><strong>Mean number of clinicians seen</strong></td>
<td>4.38</td>
<td>2.86</td>
</tr>
<tr>
<td><strong>% of subjects that have undergone investigations:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Blood tests:</td>
<td>38%</td>
<td>14%</td>
</tr>
<tr>
<td>XR:</td>
<td>54%</td>
<td>71%</td>
</tr>
<tr>
<td>Magnetic Resonance Imaging (MRI):</td>
<td>69%</td>
<td>57%</td>
</tr>
<tr>
<td>CT:</td>
<td>8%</td>
<td>0%</td>
</tr>
<tr>
<td>US:</td>
<td>100%</td>
<td>100%</td>
</tr>
<tr>
<td>Other:</td>
<td>8%</td>
<td>0%</td>
</tr>
<tr>
<td><strong>Mean length of symptoms prior to HVIGI&amp;SR:</strong></td>
<td>3.09 years</td>
<td>1.29 years</td>
</tr>
<tr>
<td>(1 data set unavailable)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>% of subjects that had undergone previous treatments:</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Oral pain killers</td>
<td>86%</td>
<td>29%</td>
</tr>
<tr>
<td>Topical pain killers</td>
<td>43%</td>
<td>0%</td>
</tr>
<tr>
<td>Physiotherapy</td>
<td>43%</td>
<td>29%</td>
</tr>
<tr>
<td>Steroid injection (singular)</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Steroid injection (multiple)</td>
<td>14%</td>
<td>14%</td>
</tr>
</tbody>
</table>
HVIGI&SR for GTPS

Procedure

All ultrasound scanning and interventions were carried out by the same skilled MSK radiologist. The injection consisted of 10 ml 0.5% Marcaine and 50 mg hydrocortisone. The area of greatest pain over the greater trochanter was marked. A 22-gauge spinal needle was introduced just deep to the periosteum using an aseptic technique (Fig. 1). The injection was performed with ultrasound guidance using a 13 MHz probe (Elegra; Siemens, Erlangen, Germany), but not real-time guidance.

Structured rehabilitation

Following the injection, the patients were reviewed by a consultant physiotherapist or his deputy, who prescribed a standardised structured rehabilitation programme. Patients were initially advised to have relative rest for 3 days and 400 mg ibuprofen up to three times a day for the first three days as required, if not contraindicated. An exercise programme was developed that met their activity-specific targets. The primary intervention was educational, with advice given to avoid positions that put the gluteal tendons into compressive positions, such as postural advice to avoid lateral slouch standing and excessive adduction postures in sitting\textsuperscript{18, 19}. Non-impact cardiovascular exercise was advised if not already undertaken, optimally three times a week with an interval training component if tolerable. Assessment was based on the clinical examination principles outlined by Grimaldi (2011) and Reiman et al. (2014)\textsuperscript{20, 21}. The specific exercise element focussed on developing isometric and concentric-eccentric endurance tolerance in weight bearing during the first two weeks of the rehabilitation programme; progressing to increased load tolerance over 4 weeks, with power and impact being included in the programme last and only to the level required for function\textsuperscript{19}. Exercises were assessment based, but typically included hip extensor and abductor activation and endurance training alongside core body control exercise that focussed on lateral trunk control, as the hip abductors are recognised to be key in core stabilisation\textsuperscript{19}. Adductor or horizontal plane rotation exercises were less commonly prescribed.

Analysis

The change in the VAS pain scale was used to assess pain effects. The HAGOS was used to describe hip-specific functions\textsuperscript{16}. Data was analysed using SPSS (SPSS statistics version 20, IBM, USA). The data was found to be normally distributed. A paired t-test was used to analyse both the pre and post-injection VAS and HAGOS scores. Significance was set at \( p < 0.05 \).

Results

Six of the eight prospective subjects completed the follow-up questionnaire. The mean follow-up time was 43.5 days. In the retrospective study fourteen of the 23 subjects completed the questionnaires, a total response rate of 63\% in the retrospective group. Overall the response rate was 65\%. The mean time between the injection and completion of the questionnaire in the retrospective group was 60 weeks. Thirty-five per cent of all subjects had experienced pain for over 2 years prior to their attendance for HVIGI&SR.

VAS Pain Scale Results

In the prospective study, the mean VAS scores were reduced from 81.7 mm (±17.6) pre-injection to 42.3 mm (±28.3), a change of 39.4 mm, at a mean time of 43.5 days follow-up (\( p=0.03 \)). In the retrospective study, the mean VAS also reduced from 74.6 mm (±10.9) pre-HVIGI&SR to 38.2 (±31.2) mm at 2 weeks, which is both statistically and clinically significant (\( p<0.01 \)). The overall reduction in the VAS score of the retrospective group was a change of 43.3 mm from 74.6 (±10.9) mm to 31.4 (±27.6) mm at a mean time of 60 weeks (\( p<0.01 \)), showing a maintained reduction in the VAS score. 62\% of retrospective subjects had returned to their normal level of activity at the mean follow up time of 60 weeks. Four out of the 8 prospective subjects reported that they

Figure 1. Insertion of the 22-spinal gauge needle under ultrasound guidance and administration of HVIGI.
had returned to their normal levels of activity at the mean time of 43.5 days.

Hip and Groin Outcome Score

For the prospective group the mean HAGOS, for the six who completed the follow-up questionnaire, increased from 173.2 (±103.3) pre-HVIGI&SR to 294.4 (±163.2) at follow-up, which was not statistically significant (p=0.13). The majority of improvement occurred in the quality of life element of the questionnaire, which was statistically significant (p=0.04). The changes for each element are shown in Figure 3. Twenty five percent (n=5) of all subjects reported short term pain as a side effect of the injection. One subject reported short term stiffness as a result of the injection. No other side effects were described or observed.

Of the fourteen retrospective subjects two went on to have surgery in the time period following their injection and prior to being followed up by this study (one had iliotibial band lengthening, the other unknown); one other subject had acupuncture. Of these subjects all had had their pain for >18 months prior to the injection. None of the prospective subjects had any additional therapy in the time in which they were followed up.

Discussion

Both the retrospective and prospective groups showed a statistically significant reduction in pain following a HVIGI into the periosteum overlying the greater trochanter in patients with GTPS followed by structured rehabilitation. A reduction of 9 mm on the VAS pain scale is considered clinically significant in an acute pain setting and therefore the reduction of 43.3 mm for the retrospective group and 39.4 mm for the prospective group is likely to be considered a good clinical improvement in the presence of chronic symptoms. Shbeeb et al. have found that a single local corticosteroid injection resulted in a greater than 60% improvement in the short and medium-term, which is similar to the 62% who had returned to their normal activity level in this study. Rompe et al. showed that a home training regime showed the greatest long term improvement in comparison to corticosteroid injections, although in the short-term corticosteroid injections were most successful. It is therefore likely that the combination of both the HVIGI and the standard rehabilitation programme in this study will ensure patients benefit in the short term from the injection and, in the longer term, from the rehabilitation programme; longer follow up is required to confirm this.

The change observed in the HAGOS in the prospective group showed a trend towards significance (p =0.13), likely due to the small group size. There was a statistically significant (p=0.04) improvement in the questions relating to quality of life, which is likely to be clinically important to those receiving any treatment. However, due to the small group size and inter-patient variability, no strong conclusions can be drawn from the HAGOS data, and further confirmation with larger numbers is required.

Unlike for patellar and Achilles tendinopathy, the proposed mechanism of effect for the HVIGI is not disruption of neovascularisation, nor is it likely to be disruption of scar tissue as none was visualised on ultrasound. It is however hypothesised that the injectate lifts the periosteum, causing a local reaction which stimulates a healing response. This mechanistic theory requires further study. It has also been hypothesised that the chemical effects of Marcaine may cause some local denervation so that pain is diminished. Further radiological studies such as magnetic resonance imaging, and also surgical examination, may be useful to evaluate the potential mechanism of the injection. Whatever the underlying mechanism it is believed that the rehabilitation programme is required to ensure long term benefit and to decrease the risk of recurrence.
It remains unclear as to why some patients have a good response to the HVIGI&SR, while others experience little or no effect. When one is considering HVIGI&SR, adherence to the rehabilitation programme is likely to be important in the overall outcome, along with other psychosocial factors such as pain beliefs and everyday use of the hip, for example at work. Future studies would therefore benefit from establishing patients’ experiences of a HVIGI&SR, especially in comparison to other treatment modalities. Also, GTPS covers so many pathologies that it may be some respond better than others and again this requires further investigation, with other imaging modalities perhaps being useful in assessing which conditions will benefit more from such an injection\textsuperscript{3-5}. One suggestion is to use MRI pre-HVIGI and follow the subsequent changes to soft tissues and the bone on MRI over a suitable follow-up period, to allow clinicians to visualise the effects on the soft tissue and to establish whether certain initial changes seen on MRI predict the response to the HVIGI&SR. This would allow the HVIGI&SR to be targeted to those in whom it is likely to be clinically effective. The side effects of pain and stiffness described by the subjects are similar to the findings in a study that used steroid injections in GTPS\textsuperscript{8}. With this being a temporary phenomenon and with the lack of any other side effects a HVIGI&SR in GTPS can provisionally be considered safe subject to confirmation in larger cohort controlled studies.

**Study Limitations**

The main limitation of this study was the size of the study, despite having both the retrospective and prospective data. This is especially noticeable within the prospective group and a longer recruitment period would be beneficial in the future. A sample size calculation using an 80\% power and 5\% significance suggests a sample size of 16 is required to show significance, 10 more than in this study\textsuperscript{25}. It should however be noted that the majority of patients attending the had already failed conservative management, as shown in Table 2 by the number of clinicians patients had seen prior to their injection, and this is likely to affect the ability to recruit patients as it is normally a referral for the HVIGI&SR. The follow up time of just over six weeks in the prospective group is also not ideal, especially as the majority of the subjects had experienced pain for over two years. However, the retrospective follow-up at 60 weeks appears to show maintenance of the pain relieving effect, with 62\% of the retrospective subjects reporting that they had returned to their normal level of activities. This is in contrast to a study which showed that the effect of a local corticosteroid injection alone declined after 1 month\textsuperscript{6}, therefore suggesting that the HVIGI&SR combination avoids this decline. Out of the retrospective subjects only three had an additional treatment; 2 underwent surgery and 1 had acupuncture suggesting the majority responded to the HVIGI and physiotherapy rehabilitation alone. It seems therefore likely that the findings in the prospective group would be maintained, although confirmation of this is required. It would also be useful to study whether adherence to the physiotherapy programme improved the outcomes associated with the HVIGI as it is likely that the injection primarily allows a pain-free period in which to start the rehabilitation programme correctly.

**Future Research**

As described above, further research is required to determine the mechanism of action of the HVIGI&SR in GTPS. More work is also required to establish the duration of the effect of a HVIGI&SR in GTPS. Anecdotally, 8 of the patients had more than one HVIGI. A randomised control trial would be the gold standard to establish the effects of the HVIGI&SR, although in the short term a larger prospective study would be beneficial.

**Conclusions**

A high volume image guided injection into the periosseous tissue of the greater trochanter followed by a structured physiotherapy-led rehabilitation programme should be considered as an effective treatment in the short and medium term for GTPS, when conservative measures have previously failed. Further controlled studies are warranted to conclusively determine the long-term effects of a HVIGI&SR, along with comparison to other treatment modalities.

**References**


Appendix 1. Questionnaires administered.

<table>
<thead>
<tr>
<th>Questionnaire for Retrospective Study</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Reference Code:</strong></td>
</tr>
<tr>
<td><strong>Date:</strong></td>
</tr>
<tr>
<td><strong>1. Age (in years)</strong></td>
</tr>
<tr>
<td>18-25</td>
</tr>
<tr>
<td>26-35</td>
</tr>
<tr>
<td>36-45</td>
</tr>
<tr>
<td>46-55</td>
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<tr>
<td>56-65</td>
</tr>
<tr>
<td>66-75</td>
</tr>
<tr>
<td>76-80</td>
</tr>
<tr>
<td>81+</td>
</tr>
<tr>
<td><strong>2. Gender</strong></td>
</tr>
<tr>
<td>Male</td>
</tr>
<tr>
<td>Female</td>
</tr>
<tr>
<td><strong>3. What sports/exercise do you take part in? (please list)</strong></td>
</tr>
<tr>
<td><strong>4. To what level do you participate? (please state for each sport/exercise)</strong></td>
</tr>
<tr>
<td>For own enjoyment/fitness</td>
</tr>
<tr>
<td>Club level</td>
</tr>
<tr>
<td>Regional level</td>
</tr>
<tr>
<td>National/international level</td>
</tr>
<tr>
<td>Amateur</td>
</tr>
<tr>
<td>Semi-professional</td>
</tr>
<tr>
<td>Professional</td>
</tr>
<tr>
<td><strong>5. How many hours/week of sport/exercise did you participate in before your hip pain started?</strong></td>
</tr>
<tr>
<td>Less than 5 hours</td>
</tr>
<tr>
<td>5-10 hours</td>
</tr>
<tr>
<td>11-15 hours</td>
</tr>
<tr>
<td>16-20 hours</td>
</tr>
<tr>
<td>More than 20 hours</td>
</tr>
<tr>
<td><strong>6. How many hours/week of sport/exercise did you participate in after your hip pain started?</strong></td>
</tr>
<tr>
<td>Less than 5 hours</td>
</tr>
<tr>
<td>5-10 hours</td>
</tr>
<tr>
<td>11-15 hours</td>
</tr>
<tr>
<td>16-20 hours</td>
</tr>
<tr>
<td>More than 20 hours</td>
</tr>
<tr>
<td><strong>7. Which is your dominant side (ie left or right handed)?</strong></td>
</tr>
<tr>
<td>Left</td>
</tr>
<tr>
<td>Right</td>
</tr>
<tr>
<td><strong>8. What is the diagnosis given for your hip pain?</strong></td>
</tr>
<tr>
<td>Bursitis</td>
</tr>
<tr>
<td>Tendinopathy</td>
</tr>
</tbody>
</table>

(to be continued)
**Questionnaire for Retrospective Study**

<table>
<thead>
<tr>
<th><strong>Muscle tear</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Greater Trochanter Pain Syndrome</td>
</tr>
<tr>
<td>You are unsure of the diagnosis but you have been given one</td>
</tr>
<tr>
<td>Doctor/physio is unsure of diagnosis</td>
</tr>
<tr>
<td>Other (please state):</td>
</tr>
</tbody>
</table>

9. Please describe the mechanism of injury (how it occurred)

10. How long did you have hip pain before the high volume injection?

| Less than 6 months |
| More than 6 months, less than 12 months |
| More than 12 months, less than 18 months |
| More than 18 months, less than 2 years |
| More than 2 years (please state) |

11. Do you have other medical conditions or previous operations? (please list)

12. How many clinicians have you seen with your hip pain? (state number of each)

| Doctor |
| Physiotherapist |
| Surgeon |
| Osteopath |
| Masseur |
| Other (please state) |

13. Which investigations have you received for your hip pain? (please tick all that apply)

| Blood tests |
| XRay |
| CT scan |
| MRI scan |
| Ultrasound scan |
| Other (please state) |

14. Which treatments for your hip pain did you use before the high volume injection? (please tick all that apply)

| Oral painkillers |
| Topical painkillers |
| Physiotherapy |
| Steroid injection (please state number) |
| Shock wave therapy (please state number) |
| Surgery |
| Other (please state): |

**Following the injection:**

15. Did you experience any side effects of the high volume injection?

| Yes |
| No |

*(to be continued)*
### Appendix 1. (cont.)

#### Questionnaire for Retrospective Study

If yes, please state side effects

16. Have you been able to carry out the exercise programme as advised by the physiotherapist?
- Yes
- No

If no, please state what difficulties you had

17. Have you received any other treatments for your hip pain since the high volume injection?
- Yes
- No

If yes, please state which treatments

18. How many hours/week of sport/exercise are you currently participating in?
- Less than 5 hours
- 5-10 hours
- 11-15 hours
- 16-20 hours
- More than 20 hours

19. Have you returned to your normal activities (regardless of current level of ability)?
- Yes
- No

20. Have you returned to your normal level of ability?
- Yes
- No

21. Any other comments:

---

**For practitioner to complete:**

Date of HVI:

Volume and content of injection:

**Level of pain**

Please mark with a cross on the line your level of pain during everyday activities before having the high volume injection.

No Pain [ ] Max Pain [ ]

Please mark with a cross on the line your current level of pain during everyday activities.

No Pain [ ] Max Pain [ ]

---

**(to be continued)**
Appendix 1. (cont.)

Initial Patient Questionnaire for Prospective Study

2. Gender
   Male
   Female

3. Height

4. Weight

5. What sports/exercise do you take part in? (Please list)

6. To what level do you participate? (please state for each sport/exercise)
   For own enjoyment/fitness
   Club level
   Regional level
   National/international level
   Amateur
   Semi-professional
   Professional

7. How many hours/week of sport/exercise did you participate in before your hip pain started?
   Less than 5 hours
   5-10 hours
   11-15 hours
   16-20 hours
   More than 20 hours

8. How many hours/week of sport/exercise have you participated in since your hip pain started?
   Less than 5 hours
   5-10 hours
   11-15 hours
   16-20 hours
   More than 20 hours

9. Which is your dominant side (i.e. left or right handed)?
   Left
   Right

10. What is the diagnosis given for your hip pain?
    Bursitis
    Tendinopathy
    Muscle tear
    Greater Trochanter Pain Syndrome
    You are unsure of the diagnosis but you have been given one
    Doctor/physio is unsure of diagnosis
    Other (please state):

11. Please describe the mechanism of injury (how it occurred)

12. How long have you had your hip pain?
    Less than 6 months

(to be continued)
Appendix 1. (cont.)

Initial Patient Questionnaire for Prospective Study

More than 6 months, less than 12 months
More than 12 months, less than 18 months
More than 18 months, less than 2 years
More than 2 years (please state)

13. Do you have other medical conditions or previous operations? (Please list)

14. How many clinicians have you seen with your hip pain? (state number of each)
   Doctor
   Physiotherapist
   Surgeon
   Osteopath
   Masseur
   Other (please state)

15. Which investigations have you received for your hip pain? (please tick all that apply)
   Blood tests
   XRay
   CT scan
   MRI scan
   Ultrasound scan
   Other (please state):

16. Which treatments have you previous used for your hip pain? (please tick all that apply)
   Oral painkillers
   Topical painkillers
   Physiotherapy
   Steroid injection (please state number)
   Shock wave therapy (please state number)
   Surgery
   Other (please state):

17. Any other comments:

For practitioner to complete:

Date of HVI:

Volume and content of injection:

Current level of pain

Please mark with a cross on the line your current level of pain during everyday activities.

No Pain  Max Pain

Follow-up Patient Questionnaire for Prospective Study

Reference Code:

Date:

1. Did you experience any side effects of the high volume injection?
   Yes

(to be continued)
Appendix 1. (cont.)

Initial Patient Questionnaire for Prospective Study

No
If yes, please state side effects

2. Have you been able to carry out the exercise programme as advised by the physiotherapist?
   Yes
   No
   If no, please state what difficulties you had

3. Have you received any other treatments for your hip pain since the high volume injection?
   Yes
   No
   If yes, please state which treatments

4. How many hours/week of sport/exercise are you currently participating in?
   Less than 5 hours
   5-10 hours
   11-15 hours
   16-20 hours
   More than 20 hours

5. Have you returned to your normal activities (regardless of current level of ability)?
   Yes
   No

6. Have you returned to your normal level of ability?
   Yes
   No

7. Any other comments:

Current level of pain

Please mark with a cross on the line your current level of pain during everyday activities.

No Pain __________ Max Pain __________

Appendix 2. HAGOS questionnaire used for prospective subjects:

HAGOS

Questionnaire concerning hip and/or groin problems

Today’s date: __________/________/________
Reference Code: _______________

INSTRUCTIONS: this questionnaire asks for your view about your hip and/or groin problem. The questions should be answered considering your hip and/or groin function during the past week. This information will help us keep track of how you feel, and how well you are able to do your usual activities.

Answer every question by ticking the appropriate box. Tick only one box for each question. If a question does not pertain to you or you have not experienced it in the past week please make your "best guess" as to which response would be the most accurate.

Symptoms

These questions should be answered considering your hip and/or groin symptoms and difficulties during the past week.

(to be continued)
### HVIGI&SR for GTPS

#### Appendix 2. (cont.)

<table>
<thead>
<tr>
<th>Question</th>
<th>None</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
<th>Extreme</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1 Do you feel discomfort in your hip and/or groin?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S2 Do you hear clicking or any other type of noise from your hip and/or groin?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S3 Do you have difficulties stretching your legs far out to the side?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S4 Do you have difficulties taking full strides when you walk?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S5 Do you experience sudden twinging/stabbing sensations in your hip and/or groin?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Stiffness**

The following questions concern the amount of stiffness you have experienced during the past week in your hip and/or groin. Stiffness is a sensation of restriction or slowness in the ease with which you move your hip and/or groin.

<table>
<thead>
<tr>
<th>Question</th>
<th>None</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
<th>Extreme</th>
</tr>
</thead>
<tbody>
<tr>
<td>S6 How severe is your hip and/or groin stiffness after first awakening in the morning?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>S7 How severe is your hip and/or groin stiffness after sitting, lying or resting later in the day?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

**Pain**

<table>
<thead>
<tr>
<th>Question</th>
<th>None</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
<th>Extreme</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1 How often is your hip and/or groin painful?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P2 How often do you have pain in areas other than your hip and/or groin that you think may be related to your hip and/or groin problem?</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The following questions concern the amount of pain you have experienced during the past week in your hip and/or groin. What amount of hip and/or groin pain have you experienced during the following activities?

<table>
<thead>
<tr>
<th>Question</th>
<th>None</th>
<th>Mild</th>
<th>Moderate</th>
<th>Severe</th>
<th>Extreme</th>
</tr>
</thead>
<tbody>
<tr>
<td>P3 Straightening your hip fully</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P4 Bending your hip fully</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P5 Walking up or down stairs</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P6 At night while in bed (pain that disturbs your sleep)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

(To be continued)
Appendix 2. (cont.)

P7 Sitting or lying
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

The following questions concern the amount of pain you have experienced during the past week in your hip and/or groin. What amount of hip and/or groin pain have you experienced during the following activities?

P8 Standing upright
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

P9 Walking on a hard surface (asphalt, concrete, etc.)
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

P10 Walking on an uneven surface
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

Physical function, daily living

The following questions concern your physical function. For each of the following activities please indicate the degree of difficulty you have experienced in the past week due to your hip and/or groin problem.

A1 Walking up stairs
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

A2 Bending down, e.g. to pick something up from the floor
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

A3 Getting in/out of car
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

A4 Lying in bed (turning over or maintaining the same hip position for a long time)
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

A5 Heavy domestic duties (scrubbing floors, vacuuming, moving heavy boxes etc.)
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

Function, sports and recreational activities

The following questions concern your physical function when participating in higher-level activities. Answer every question by ticking the appropriate box. If a question does not pertain to you or you have not experienced it in the past week please make your “best guess” as to which response would be the most accurate. The questions should be answered considering what degree of difficulty you have experienced during the following activities in the past week due to problems with your hip and/or groin.

SP1 Squatting
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

SP2 Running
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

SP3 Twisting/pivoting on a weight bearing leg
None Mild Moderate Severe Extreme
☐ ☐ ☐ ☐ ☐

(to be continued)
Appendix 2. (cont.)

SP4 Walking on an uneven surface
None □ □ □ □ □
Mild □ □ □ □ □
Moderate □ □ □ □ □
Severe □ □ □ □ □
Extreme □ □ □ □ □

SP5 Running as fast as you can
None □ □ □ □ □
Mild □ □ □ □ □
Moderate □ □ □ □ □
Severe □ □ □ □ □
Extreme □ □ □ □ □

SP6 Bringing the leg forcefully forward and/or out to the side, such as in kicking, skating etc.
None □ □ □ □ □
Mild □ □ □ □ □
Moderate □ □ □ □ □
Severe □ □ □ □ □
Extreme □ □ □ □ □

SP7 Sudden explosive movements that involve quick footwork, such as accelerations, decelerations, change of directions etc.
None □ □ □ □ □
Mild □ □ □ □ □
Moderate □ □ □ □ □
Severe □ □ □ □ □
Extreme □ □ □ □ □

SP8 Situations where the leg is stretched into an outer position (such as when the leg is placed as far away from the body as possible)
None □ □ □ □ □
Mild □ □ □ □ □
Moderate □ □ □ □ □
Severe □ □ □ □ □
Extreme □ □ □ □ □

**Participation in physical activities**

The following questions are about your ability to participate in your preferred physical activities. Physical activities include sporting activities as well as all other forms of activity where you become slightly out of breath. When you answer these questions consider to what degree your ability to participate in physical activities during the past week has been affected by your hip and/or groin problem.

**PA1 Are you able to participate in your preferred physical activities for as long as you would like?**
Always □ □ □ □ □
Often □ □ □ □ □
Sometimes □ □ □ □ □
Rarely □ □ □ □ □
Never □ □ □ □ □

**PA2 Are you able to participate in your preferred physical activities at your normal performance level?**
Always □ □ □ □ □
Often □ □ □ □ □
Sometimes □ □ □ □ □
Rarely □ □ □ □ □
Never □ □ □ □ □

**Quality of Life**

**Q1 How often are you aware of your hip and/or groin problem?**
Never □ □ □ □ □
Monthly □ □ □ □ □
Weekly □ □ □ □ □
Daily □ □ □ □ □
Constantly □ □ □ □ □

**Q2 Have you modified your life style to avoid activities potentially damaging to your hip and/or groin?**
Not at all □ □ □ □ □
Mildly □ □ □ □ □
Moderately □ □ □ □ □
Severely □ □ □ □ □
Totally □ □ □ □ □

**Q3 In general, how much difficulty do you have with your hip and/or groin?**
None □ □ □ □ □
Mild □ □ □ □ □
Moderate □ □ □ □ □
Severe □ □ □ □ □
Extreme □ □ □ □ □

**Q4 Does your hip and/or groin problem affect your mood in a negative way?**
Not at all □ □ □ □ □
Rarely □ □ □ □ □
Sometimes □ □ □ □ □
Often □ □ □ □ □
All the time □ □ □ □ □

**Q5 Do you feel restricted due to your hip and/or groin problem?**
Not at all □ □ □ □ □
Rarely □ □ □ □ □
Sometimes □ □ □ □ □
Often □ □ □ □ □
All the time □ □ □ □ □

Thank you very much for completing all the questions in this questionnaire.