Pilot Study of Ultrasound-Guided Corticosteroid Hip Injections by Emergency Physicians

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Introduction: Our objective was to assess the efficacy of ultrasound-guided hip injections performed by emergency physicians (EPs) for the treatment of chronic hip pain in an outpatient clinic setting.

Methods: Patients were identified on a referral basis from the orthopedic chronic pain clinic. The patient population was either identified as having osteoarthritis of the hip, osteonecrosis of varying etiologies, post-traumatic osteoarthritis of the hip, or other non-infectious causes of chronic hip pain. Patients had an ultrasound-guided hip injection of 4ml of 0.5% bupivacaine and 1ml of triamcinolone acetate (40mg/1ml). Emergency medicine resident physicians under the supervision of an attending EP performed all injections. Pain scores were collected using a Likert pain scale from patients prior to the procedure, and 10 minutes post procedure and at short-term follow-up of one week and one month. The primary outcome was patient-reported pain score on a Likert pain scale at one week.

Results: We performed a total of 47 ultrasound-guided intra-articular hip injections on 44 subjects who met inclusion criteria. Three subjects received bilateral injections. Follow-up data were available for 42/47 (89.4%) hip injections at one week and 40/47 (85.1%) at one month. The greatest improvement was at 10 minutes after injection with a mean decrease in Likert pain score from pre-injection baseline of 5.57 (95% CI, 4.76-6.39). For the primary outcome at one week, we found a mean decrease in Likert pain score from pre-injection baseline of 3.85 (95% CI, 2.94-4.75). At one month we found a mean decrease in Likert pain score of 1.8 (95% CI, 1.12-2.53). There were no significant adverse outcomes reported.


INTRODUCTION

Hip pain is a common complaint with a wide variety of etiologies. These range from the benign and chronic, such as osteoarthritis, to the acutely joint-threatening, such as septic arthritis. Chronic hip pain has an estimated prevalence of up to six percent and is a common cause of pain in patients presenting to orthopedic clinics and emergency departments (EDs).¹ Joint injections with corticosteroids are first-line recommended therapy by the American College of Rheumatology;² and the European League Against Rheumatism recommends intra-articular hip injections for flares of chronic hip osteoarthritis.³ A 2007 randomized controlled trial has also demonstrated clear efficacy without any complications.³ Ultrasound (US) guidance for either hip arthrocentesis...
or hip injections has since been described in the radiology, 
rheumatology, and orthopedic literature.\textsuperscript{5-9} Three trials in 
emergency medicine (EM) literature have shown efficacy of US 
to aid in diagnosis of hip effusions,\textsuperscript{10-13} but no EM trials to date 
have demonstrated US-guided hip injections as efficacious in the 
treatment of chronic hip pain. US-guided hip injections have 
been shown to be more safe and efficacious as compared to blind 
jagrams.\textsuperscript{6,7,13} However, these procedures are rarely performed in 
an ED setting, and many front-line practitioners who encounter 
patients with hip pain from degenerative diseases of the hip may 
not have the training to perform the procedure. Frequently the 
intrarticular corticosteroid hip injections are performed only 
in specialty clinics, limiting the access for optimal pain control 
in patients with non-infectious hip pathology. With adequate 
training and coordinated follow-up, clinicians could facilitate 
timely pain control (without overreliance on standard opioid and 
anti-inflammatory therapies) for these patients.

In a prospective cohort pilot study, we aim to analyze 
the effect of US-guided corticosteroid hip injections on pain 
scores as performed by EM trainees.

METHODS

Study Design and Setting

This is a prospective pilot study of US-guided 
intra-articular hip injections performed by EM trainees 
with bupivacaine and triamcinolone for hip pain due 
to osteoarthritis, avascular necrosis, and other chronic 
conditions. Patients were consecutively enrolled from an 
orthopedic surgery clinic at a busy, urban hospital and 
trauma center. The Institutional Review Board of Alameda 
County Medical Center approved this study.

Selection of Participants

Enrollment occurred from September 2012 to February 
2013. Adult patients (age\textgreater{}18) were eligible for inclusion if 
deemed to have chronic hip pain related to osteoarthritis, 
avascular necrosis, post-traumatic degenerative changes, late 
sequelae of septic arthritis, or hip dysplasia as determined by 
the referring orthopedic surgery attending. All patients were 
consented for the procedure and enrollment into the study. 
Exclusion criteria were any signs of systemic infection such as 
fever, recent illnesses in the past two weeks, contraindication 
or allergy to the injection agents, anticoagulant therapy other 
than aspirin, previous hip injection within the last four months, 
planned total hip arthroplasty in the coming four months, if 
an interpreter was not available for the consent process, or if 
the patient was receiving a diagnostic injection as part of a hip 
or back pain work up. Patients found ineligible for the study 
or who declined study enrollment still had the opportunity to 
receive a hip injection.

Interventions

All patients received the study injection solution of 4ml of 
0.5% bupivacaine and 1ml of triamcinolone (40mg). The dosing 
and medication selection for analgesia and steroid injection 
were selected based on prior research on fluoroscopically- 
guided hip injections.\textsuperscript{4} An EM attending physician, fellowship 
trained in emergency US, supervised all procedures. First and 
second-year EM resident physicians performed all procedures 
after receiving an instructional handout and a standardized five- 
minute bedside training session. All trainees had completed 
a one-month US rotation that included instruction on needle- 
guided procedures, such as US-guided central lines, peripheral 
venous access, and nerve blocks. They also received a five- 
minute bedside tutorial on the anatomy of the femoral neck and 
and anterior synovial recess. Trainees had also completed a 
one-month orthopedics rotation during which they performed 
landmark-based knee and shoulder injections as part of their 
clinical rotation. None of the residents had performed hip 
jjections prior to this study, and each resident performed 1-2 
jections during the clinic. An ultrasound fellowship-trained 
EM attending physician supervised all procedures.

Procedure

An ultrasound system (Sonosite M-Turbo; Bothell, WA) 
with a low frequency curvilinear probe (2-5MHz) was used 
to identify the hip joint. Local anesthesia over the injection 
site was applied using ethyl chloride spray. Using standard 
sterile procedure and local analgesia, a 10cc syringe filled 
with a 5cc mix of bupivacaine and 40mg of triamcinolone 
attached to a 20g 3.5 inch spinal needle was guided into 
the joint space with real-time in-plane ultrasound guidance 
(Figure 1) (Video). This solution was then injected after 
ultrasonographic confirmation that the needle tip was in the 
joint space (Figure 2). Patients were observed for a period of 
twenty minutes after the procedure.

![Figure 1. Set-up of ultrasound machine and patient for injection.](image)

Methods and Measurements

We used a standardized data collection tool to collect 
demographic and clinical information of all enrolled subjects, 
including age, sex, race, etiology for pain, whether or not the 
patient previously had a hip injection performed, and whether 
or not there were any complications or adverse events as
Three subjects received bilateral injections. Follow-up data were available for 42/47 (89.4%) hip injections at one week and 40/47 (85.1%) at one month. The median age was 56 years (IQR 45-62), and the majority of patients were female (63.6%). The age of patients ranged from 19 to 75 years. Additional demographic and clinical information of the cohort are available in Table 1. Osteoarthritis was the most common cause of chronic hip pain, present in 37/47 (78.2%) of included patients, followed by avascular necrosis 5/47 (10.6%), hip dysplasia 2/47 (4.3%), and other causes 3/47 (6.4%). Previous injections had been performed in 5/47 (10.6%) of patients.

Median pain scores at all-time intervals and mean changes in Likert pain score at the follow-up intervals are available in Table 2. We found clinically and statistically significant decreases in pain scores at all-time intervals, with the greatest improvement at 10 minutes after injection. For the primary outcome at one week, we found a mean decrease in Likert pain score from the pre-injection baseline of 3.85 with a 95% CI from 2.94-4.75. Additionally at one month, there was a mean decrease in Likert pain score from pre-injection baseline of 1.8 with a 95% CI from 1.12-2.53.

Graphical representation of individual pain scores for the 42 patients available for follow-up at the primary outcome of one week appears in Figure 3. Patients with both high and low levels of pre-procedural pain had improvement in their pain scores at one week. At one week follow-up one patient had an increase in his pain, three patients had no change in their pain, and seven patients had no pain at all after one week.

One patient reported transient dizziness after the procedure (lasting one minute). No other complications were described by the patient. Pain scores before and at various times after the hip injection were collected using a 0 to 10 Likert scale. A research assistant performed telephone follow-up to obtain pain scores after clinic discharge and determine whether or not any complications occurred according to a standardized questionnaire.

**Outcomes**

Our primary outcome was decrease in pain score at one week. We chose this as the primary outcome interval because the effect of local anesthetics will have worn off and the corticosteroid effect should have become evident. We additionally collected pain scores at five minutes, 10 minutes, and one month after the procedure.

**Analysis**

We collected descriptive statistics for our cohort and report median pain scores, with interquartile range (IQR), pre-injection, then post-injection at five minutes, 10 minutes, one week, and one month. We calculated mean change in pain scores within a 95% CI, from the pre-injection score to the post-injection intervals. Graphical and statistical methods were used to assess for normality of the pain score and pain score change distributions. Additionally, we determined mean changes in pain score stratified by pre-injection pain levels (mild<4, moderate 4-7, or severe >8). Finally, we used a multivariate linear regression model to assess whether covariates (age, sex, race, etiology of hip pain, or pre-injection pain score) were associated with reduction in pain score at one week. Statistical analyses were performed with Stata SE version 11 (StataCorp LP, College Station, TX).

**RESULTS**

We performed a total of 47 US-guided intra-articular hip injections on 44 subjects who met inclusion criteria.
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Multivariate analysis did not identify an association between age, sex, race, etiology of hip pain, or pre-injection pain score and change in hip pain at one week.

DISCUSSION

Our study demonstrates that EM trainees can effectively perform US-guided corticosteroid hip injections as a method for the treatment of chronic hip pain. We noted a statistically significant reduction in pain scores at one week, and while the effect waned somewhat, patients continued to have modest pain relief at one month.

Our analysis demonstrates that US-guided corticosteroid intra-articular hip injection as performed by EM trainees is effective at decreasing chronic hip pain acutely and over a one-week period. This is a well-established and safe procedure practiced in various settings, and has been reported in the ED setting. In our study, junior EM providers with modest comfort with point-of-care US were able to successfully perform a US-guided hip injection after a brief tutorial.

Our results suggest that the technical difficulty of the procedure is low. The trainees performing the study were supervised, though the attending physician did not intervene during any of the procedures. Successful performance of the procedure was evident due to the dramatic improvement in pain after just ten minutes consistent with expected analgesia using bupivacaine.

The American College of Rheumatology advises corticosteroid injections as a first-line therapy for osteoarthritis of the hip. To our knowledge, the American College of Emergency Physicians has no comment on intra-articular steroid injections. Integration of this treatment to the care of ED patients should be considered for a number of reasons. Degenerative diseases of the hip, as well as other painful chronic musculoskeletal conditions, may be treated primarily in the ED setting with non-steroidal anti-inflammatory drugs (NSAIDs), acetaminophen, opioid pain medications, and referral to specialists. There are significant risks associated with long-term use of NSAIDs and a growing concern over opioid drug abuse. In settings where access to primary care, pain management, and orthopedic specialists are limited, wait-times to see a practitioner may exceed months. This temporizing procedure may assist with chronic pain management, but, nonetheless, should only be performed in conjunction with appropriate consultation and referral to providers who will ultimately care for these patients. With this in mind, providing timely corticosteroid injections are an appealing approach to pain management.

Table 2. Pain scores and differences at established intervals.

<table>
<thead>
<tr>
<th>Outcome</th>
<th>N (hips)</th>
<th>Median pre-injection Pain score (IQR)</th>
<th>Mean decrease in pain score from pre-injection baseline (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-injection</td>
<td>47</td>
<td>8 (7-10)</td>
<td>-</td>
</tr>
<tr>
<td>5 minutes</td>
<td>47</td>
<td>2 (0-5)</td>
<td>4.98 (4.18-5.78)</td>
</tr>
<tr>
<td>10 minutes</td>
<td>47</td>
<td>1 (0-4)</td>
<td>5.57 (4.76-6.39)</td>
</tr>
<tr>
<td>1 week</td>
<td>42</td>
<td>4 (1-5)</td>
<td>3.85 (2.94-4.75)</td>
</tr>
<tr>
<td>1 month</td>
<td>40</td>
<td>5 (4-8)</td>
<td>1.8 (1.12-2.53)</td>
</tr>
</tbody>
</table>

Figure 3. Individual patient pain scores from pre-injection to one week after hip injection.*
*Includes 42/47 injections available for follow up at one week.
management for degenerative diseases of the hip to decrease repeat ED visits for chronic pain and opioid prescriptions. More specifically, similar attempts at moving towards multi-modal approaches to pain management have been shown to decrease the need for opioids, and ultimately may help curb the adverse effects and abuse associated with these and other controlled drugs. Integration of intra-articular hip injections into practice in the ED may not be successful without careful planning. The scope of this study does not assess the efficacy or feasibility in an ED and it if it is implemented the higher acuity and undifferentiated patient population needs to be considered. As with any procedure, clinicians should be adequately trained and comfortable and require the necessary credentialing. They should screen for patients with any signs of septic arthritis and should not perform the procedure in patients with signs of an infectious etiology for their pain, prosthesis hip joints, allergies to the medications, or overlying cellulitis. We would suggest that any patient with suspicion of a septic joint and a visualized effusion should have their joint aspirated using this same US-guided technique. In cases where the risks or benefits of an intra-articular hip injection are not clear, providers should not perform it or should obtain expert consultation. Additionally, patient consent is paramount. Known rare but significant complications of this procedure include avascular necrosis, post-procedural septic joint, and increased risk of post-operative septic joint when performed in proximity to surgery, and as such, patients must be informed and appropriately counseled. Finally, coordination of care beyond the ED is important, specifically with primary care and specialty clinic follow-up. Again, we reiterate that this procedure should not be done in lieu of appropriate referral. Rather, it can serve as an effective temporizing intervention for pain reduction, and should be followed by referral to an orthopedic surgeon, general practitioner, physical therapist, and/or pain specialist. Similarly, an understanding by the community of physicians who will be caring for these patients should be reached. It will be important, for example, that orthopedic surgeons seeing these patients in follow-up are aware of and comfortable with the performance of this procedure by emergency providers. In our experience at our medical center, there has been wide acceptance of intra-articular steroid injections of the hip by emergency providers, orthopedic specialists, and primary care physicians.

LIMITATIONS

There are inherent limitations to our study. Our non-blinded and uncontrolled method does not show that this method is superior compared to placebo, though this has already been established. As in most ultrasound studies, the issue of operator dependence is a limitation; however, EM providers may have extensive experience with US-guidance during procedures. There is no doubt that this procedure is unique as compared to vascular access or nerve blockade, and, as such, our training module and supervision by ultrasound fellowship-trained attending physicians was aimed at ensuring familiarity with the anatomy, needle insertion approach and angle, as well as appropriate injection of anesthetic and steroid. Training of providers on the unique technical aspects of this procedure will be paramount before adoption into clinical practice.

Additionally, for the purposes of this study, we enrolled patients directly from an orthopedic clinic. While it remains unclear if the benefit of this procedure would be as evident in ED patients, its benefit has been shown in various settings performed by providers of different specialties, suggesting that the benefit would also be realized in the ED setting. ED patients with hip pain are likely a more heterogeneous and higher acuity population in terms of presenting complaint, etiology of pain, or reliability for follow-up. While our clinic’s patient population represented a more heterogeneous population than previous studies in terms of etiology of disease, it may not approximate the heterogeneity likely to be seen in the ED. The patients in our study were identified by an orthopedic attending and selected as patients who would safely benefit from such a procedure, and as such we must temper our results in light of this limitation. Taken together, this limitation stresses the importance of careful ED patient selection and screening, and, in some instances, may support the use of consultation prior to performing the procedure.

CONCLUSION

This current study suggests that intra-articular hip injections for chronic pain conditions can be reliably taught to EM providers and, similar to previous studies, we have shown that it is safe and effective. It is entirely possible, however, that intra-articular hip injections may not be a feasible, efficient, or effective intervention when introduced into clinical practice in the ED. Prospective observational or randomized trials in this setting are warranted prior to wider acceptance of this procedure.

Future research should study the efficacy of this procedure in a larger population of ED patients. Analyses should specifically focus on proper patient selection, and identification of factors associated with higher efficacy (i.e., which etiology of hip pain benefits more from the injection). Meanwhile, educational resources should be made available to emergency providers who wish to learn this skill for their practice or study its use.
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Video. Hip Injection.

REFERENCES