Healthy Hip Joints Have Different Macroscopic and Microscopic Capsular Nerve Architecture Compared With Hips With Osteoarthritis, Femoroacetabular Impingement Syndrome, and Developmental Dysplasia of the Hip: A Systematic Review

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Purpose: To perform a systematic review to identify macroscopic and microscopic patterns and differences in hip capsule innervation between normal hips and hips with osteoarthritis (OA), femoroacetabular impingement (FAI) syndrome, and developmental dysplasia of the hip (DDH).

Methods: A systematic review was performed using Preferred Reporting Items for Systematic reviews and Meta-Analyses guidelines. Multiple databases were searched for both clinical and basic science laboratory studies on hip capsule innervation. Non-innervation capsule and non-human animal studies were excluded. Macroscopic and microscopic differences in capsular innervation between normal hips, and hips with OA, FAI, and DDH were analyzed. Methodological quality assessment of all studies included in this review was completed using the Methodological Index for Non-randomized Studies.

Results: Ten articles were analyzed (263 specimens; 211 patients, 52 cadavers; mean Methodological Index for Non-randomized Studies 10/16). The hip capsule is innervated by the sciatic and superior gluteal nerves posterosuperiorly, nerve to quadratus femoris and inferior gluteal nerve posteroinferiorly, and femoral and obturator nerves anteriorly. The anterior-superior capsule between 1:00 and 2:30 o’clock on a right hip is a safe internervous zone. The superolateral capsule has the greatest density of mechanoreceptors and sensory fibers. OA is associated with a greater expression of nerve fibers compared with normal hips but does not correlate with pain or disability. No significant differences were found in nerve fiber expression among patients with DDH, FAI, or normal hips. A negative correlation is seen with aging and pain fiber expression.

Conclusions: The hip capsule has a complex macroscopic and microscopic innervation pattern with varying nerve fiber expression from at least 6 separate peripheral nerves. OA is associated with a greater expression of nerve fibers, although nerve fiber expression does not correlate with painful pathology.

Level of Evidence: IV, Systematic review of level I-IV studies.

Pain is an essential symptom of hip osteoarthritis (OA), femoroacetabular impingement syndrome (FAI), and developmental dysplasia of the hip (DDH). Hip pain may be severe, equated to a pain “worse than death.” Nociception (peripheral noxious tissue injury detected by nociceptors) and pain (higher level central perception and interpretation of that peripheral tissue damage) are not the same entity, as nociception can...
occur without pain. This is an important distinction in the hip, considering the high prevalence of asymptomatic imaging abnormalities. Because of these complexities within pain and nociception, pain may be a singular indication for surgical intervention in patients with hip pathology. Therefore, the most important aim of treatment is pain relief and improvement of physical function and overall quality of life.

Hip pain results from the stimulation of free endings of nociceptors of slow-conducting (unmyelinated axons, C-fibers) and faster conducting (thinly myelinated axons, A-delta fibers) nerve fibers. The perception of pain in the hip may be transmitted by autonomic nerve supply of the intraosseous blood vessels, innervation of the synovium and capsule, and periarticular muscles. This effect can be amplified by biochemical agents, such as bradykinin or histamine, which may activate the endings of nerve fibers, triggering pain, whereas other agents, such as substance P or prostaglandins, mainly enhance the sensitivity to applied stimuli. Moreover, a persistent noxious, mechanical, or thermal stimulation input can be involved in developing or increasing central sensitization and release processes, leading to or supporting chronic pain.

The hip capsule is a complex anatomical structure that plays an important role in native hip stability. The capsule is a condensation of the iliofemoral, pubofemoral, and ischiofemoral ligaments and the zona orbicularis. While capsular management in hip-preservation surgery has evolved significantly over the recent past, the main interest lies in its contribution to joint stability. Although the issue is not “solved,” capsule preservation with repair is more of a rule now, rather than an exception. Unfortunately, the controversial issues have not investigated the capsule’s innate role in pain generation. The presence and quantity of neuronal elements in the joint capsule may implicate the capsule itself in the pathogenesis of pre- or post-treatment pain—too loose, too tight, too irregular, too thick, too thin, too much adhesion formation (capsulolabral adhesions), or foreign body irritation (suture). A thorough understanding of the normal and diseased hip joint capsule neuroanatomy can help guide treatment, especially with surgical intervention that involves capsulotomy, capsulectomy, capsular repair, or capsular plication. The purpose of this study was to perform a systematic review to identify macroscopic and microscopic patterns and differences in hip capsule innervation between normal hips and hips with OA, FAI syndrome, and DDH. It was hypothesized that normal hips would have different macroscopic and microscopic capsular nerve architecture compared with hips with OA, FAI and DDH.

**Methods**

**Design and Search Strategy**

A systematic review was registered on PROSPERO on April 18, 2018 (registration ID: CRD 42018094248). PRISMA (Preferred Reporting items for Systematic reviews and Meta-Analyses) guidelines were followed. Under the PROSPERO registration, similar previous systematic reviews and meta-analyses were sought, and none were identified. Separate electronic searches of the following databases were conducted: MEDLINE, Cochrane Central Register of Controlled Trials, SCOPUS and Sport Discus. The searches were performed on April 26, 2020. To ensure a stringent search strategy of relevant literature, key words including “hip capsule,” “innervation,” “hip osteoarthritis,” “femoroacetabular impingement,” and “developmental dysplasia” were combined with Boolean operators to develop a search protocol (Appendix Table 1). A hand search of the included reference lists also was performed to further minimize unintentional exclusion of relevant studies.

**Eligibility Criteria**

Levels I, II, III, and IV evidence (based on Oxford Centre for Evidence Based Medicine) clinical outcome therapeutic, diagnostic, or prognostic studies and basic science laboratory studies (including human cadaver specimens) published in English language that involved normal or diseased human hip capsule innervation were eligible for inclusion. Differences in macroscopic and microscopic differences in innervation of the normal hip and hips with OA, FAI, and DDH were sought, extracted, and analyzed. Macroscopic innervation studies were defined as any investigation of nerves visible with the naked eye without microscopic amplification. Microscopic innervation studies were defined as any investigation reporting outcomes of microscopic analysis of innervation of the hip capsule. Hips with arthritis were defined in a variety of ways: the study explicitly reported “osteoarthritis,” joint space width less than 2 mm, Tönnis grade 2 or 3, or the presence of osteophytes, subchondral sclerosis, and subchondral cysts. Hips with FAI were defined based on patient symptoms, clinical signs on physical examination, and radiographic features consistent with either cam (alpha angle greater than 55° on any view, 3-dimensional head-neck junction asphericity) or pincer (lateral center edge angle greater than 40°, anterior center edge angle greater than 40°, protrusio acetabulae, posterior wall and ischial spine signs [acetabular retroversion], or crossover sign with computed tomography evidence of focal loss of cranial acetabular anteverision) morphology. Hips with DDH were defined based on patient symptoms, clinical signs of physical examination, and radiographic features consistent with dysplasia (broken Shenton’s line, lateral center edge angle less than 15 to 20°, anterior center edge angle less than 15 to 20°, Tönnis angle greater than 10 to 15°, femoral head extrusion index greater than 25%, or computed tomography evidence of anterior, lateral, posterior, or global femoral head uncoverage). Non-human animal model, level V evidence, studies that did not focus on hip capsular innervation, and non-English
language studies or reports were excluded. The search results were reviewed for duplicates and the inclusion criteria to determine articles that were included in the final analysis (Fig 1).

**Data Extraction and Analysis**

Two of the authors independently reviewed, and data were extracted from all articles using methodology recommended by Harris et al. The study type, design, methods, and level of evidence were identified. Methodological quality assessment of all studies included in this review was completed using the Methodological Index for Non-randomized Studies. For clinical studies, the levels of evidence were assigned based on the Oxford Centre for Evidence-Based Medicine. Because of the heterogeneity of outcome measures and low level of evidence, a best-evidence synthesis was used instead of a meta-analysis.

**Results**

Ten studies were included in the systematic review (Fig 1). There were 4 cadaveric and 6 in vivo studies describing 263 specimens (Table 1). Mean Methodological Index for Non-randomized Studies was 10.0 ± 3.5.

**Macroscopic Innervation of the Hip Capsule**

Three cadaveric studies describing 44 cadavers investigated macroscopic innervation of normal hip capsules to find that the hip capsule is innervated by the sciatic and superior gluteal nerves posterosuperiorly (ischiofemoral ligament), nerve to quadratus femoris and inferior gluteal nerves posteroinferiorly (ischiofemoral ligament), and femoral and obturator nerves anteriorly (iliofemoral and pubofemoral ligament) (Fig 2). One study demonstrated that the arch in the anterosuperior right hip capsule between 1:00 and 2:30 o’clock is a safe internervous zone for capsular incision during surgery.

**Microscopic Innervation of the Hip Capsule**

One cadaveric and 6 in vivo studies (8 cadavers, 211 live humans) performed microscopic analyses of hip capsule innervation. One study used gold chloride staining in 8 cadavers without a known hip pathology to map nerve fiber distributions within the
| Article* | Publication Year | Country | Specimen Type | Mean Age, y (Range) | Specimen, n | Capsular Area | Study Modality | Staining Technique | Tissue Analysis | Nerves Identified | Disease |
|----------|------------------|---------|----------------|---------------------|-------------|--------------|---------------|------------------|-----------------|----------------|---------|
| Rabinowicz and Jacqueline 19 | 1990 | Switzerland | Live humans | 61.3 (30-90) | 32 | Unspecified | Histologic analysis | Hematoxylin, eosin, Gieson, Luxol-fast-blue, Loyez, Gleys, Tibor-pap, Gomori | Microscopic | Unspecified | OA, AS, AVN, FNFx |
| Birnbaum et al. 1997 | 1997 | Germany | Cadaver | NS | 11 | Unspecified | Gross dissection | None | Macroscopic | FN, ON, AON, SGN, NQF, SN | Unspecified |
| Gáspár et al. 2004 | 2004 | Hungary | Live humans | Adults, age NS | 22 | Unspecified | Immunohistochemistry | Anti-NF, Anti-SP, Anti-CGRP, Anti-NK1 | Microscopic | None | OA |
| Kampa et al. 2007 | 2007 | United Kingdom | Cadaver | 81 (54-107) | 20 | Unspecified | Gross dissection | None | Macroscopic | FN, ON, AON, IGN, SGN, NQF, SN | Unspecified |
| Saxler et al. 2007 | 2007 | Germany | Live humans | 68.7 (52-80) | 9 | Unspecified | Immunohistochemistry | Anti-Sp, Anti-CGRP | Microscopic | Unspecified | FNFx, OA, painless failed THA |
| Gerhardt et al. 2012 | 2012 | USA | Cadaver | 76.5 (68-93) | 8 | Anterior, posterior, inferior, superolateral | Histologic analysis | Gold chloride | Microscopic | Unspecified | OA |
| Hauersath et al. 2013 | 2013 | Germany | Live humans | 55.6 (8-87) | 34 | Anterior, posterior, inferior, superolateral | Immunohistochemistry, histologic analysis | Anti-S100, Anti-Sp, Anti-NF, Anti-nociceptin, Anti-neuropeptide Y | Microscopic | Unspecified | AVN, OA, FAI |
| Desteli et al. 2014 | 2014 | Turkey | Live humans | 0.9 (0.5-1.5) | 60 | Unspecified | Immunohistochemistry | Anti-S100 | Microscopic | Unspecified | Normal, DDH |
| Grzegorzewski et al. 2014 | 2014 | Poland | Live humans | 9.5 (5-18) | 34 | Unspecified | Immunohistochemistry | Anti-S100, Anti-SP | Microscopic | Unspecified | CP, DDH |
| Short et al. 2018 | 2018 | Canada | Cadaver | 79.3 ± 11.9 | 13 | Anterior | Gross dissection, ultrasound | None | Macroscopic | FN, ON, AON | Unspecified |

AO, accessory obturator nerve; AS, ankylosing spondylitis; AVN, avascular necrosis; CGRP, calcitonin gene-related peptide; CP, cerebral palsy; DDH, developmental dysplasia of the hip; FAI, femoroacetabular impingement; FN, femoral nerve; FNFx, femoral neck fracture; IGN, inferior gluteal nerve; NF, neurofilament; NK1, neurokinin 1; NQF, nerve to quadratus femoris; NS, not specified; OA, osteoarthritis; ON, obturator nerve; SGN, superior gluteal nerve; SN, sciatic nerve; SP, substance P; THA, total hip arthroplasty.

*All included clinical studies were Level IV evidence.
hip capsule.\textsuperscript{24} This study demonstrated that the superolateral zone of the capsule had the greatest density of mechanoreceptors and sensory fibers followed by the anterior zone of the capsule. No sensory fibers were found in either the inferior or posterior zones of the hip capsule specimens.

A total of 94 pediatric patients were assessed by 2 in vivo studies.\textsuperscript{26,27} Thirty subjects had DDH. Both articles were level IV evidence. Both studies performed immunohistochemical analysis of hip capsule biopsies for neurogenic protein S-100 to demonstrate that pain and femoral head articular cartilage loss is associated with a greater expression of nerve fibers. They also demonstrated that patients with cerebral palsy had a significantly greater number of nerve fibers compared with patients with DDH. However, there were no statistically significant differences in nerve fiber expression between DDH and normal hips.

A total of 117 adult patients were assessed by 4 in vivo studies using light microscopy histology and immunohistochemistry. Eleven adults had FAI and 7 adults had dysplasia. The remaining subjects had OA, avascular necrosis (AVN), femoral neck fracture, or inflammatory arthritides. All 4 articles were level IV evidence. Two studies analyzed nerve fiber densities within the hip capsule among patients with OA.\textsuperscript{21,23} Saxler et al.\textsuperscript{23} found that patients with OA have significantly greater nerve expression and increased inflammatory mediators such as prostaglandins, neuropeptides, bradykinin, and cytokines.\textsuperscript{29,30} However, there are limited number of studies that investigate the quantity and distribution pattern of nerve fibers in relation to pain. Among the studies included in this systematic review, Grzegorzewski et al.\textsuperscript{27} demonstrated a correlation between pain and the number of nerve fibers, specifically among pediatric patients with cerebral palsy. Saxler et al.\textsuperscript{23} found that patients with OA have significantly greater nerve fiber densities compared with hips without OA. However, Gáspár et al.\textsuperscript{21} found that this high nerve fiber density does not correlate with pre- and postoperative pain or disability. Another study performed a histologic analysis of nerve fibers among patients with OA, rheumatoid arthritis, ankylosing spondylitis, femoral neck fracture, and AVN.\textsuperscript{19} They demonstrated that in AVN, rheumatoid arthritis, and femoral neck fractures, the number of nerve fibers within the capsule decreases over time and are also transformed into fibrous tissue. They found that these transformed fibrous tissues do not transmit pain leading to less painful pathologies when compared with OA and ankylosing spondylitis, for which nerve fibers do not transform into fibrous tissues. Another study investigated capsule innervation among patients with AVN, OA, and FAI to find that there is a negative correlation with aging and pain fiber expression.\textsuperscript{25} They demonstrated that patients with AVN had a greater distribution of pain-associated nerve fibers in the superior aspect of the capsule but found no significant difference in nerve fiber distribution between patients with FAI and OA.

**Discussion**

The most important finding of this systematic review of 4 cadaveric and 6 in vivo studies was that normal and diseased hip capsule innervation is complex. Macroscopic analyses of cadaveric studies were able to demonstrate that a normal hip capsule is innervated by the sciatic and superior gluteal nerves posterosuperiorly, nerve to quadratus femoris and inferior gluteal nerve posteroinferiorly, and femoral and obturator nerves anteriorly. These studies also identified the existence of an internervous safe zone in the anterior-superior capsule, a common location where capsular incisions are made during hip-preservation surgery. Microscopic analyses of both cadaveric and in vivo studies were able to demonstrate that nerve fiber expression did not correlate with pain. This information can be used to guide hip-preservation surgery capsular management.

This investigation studied a complex subject regarding the unique innervation pattern of the hip capsule and its association with pain in hip pathology. Previous studies have demonstrated that joint pain is associated with increased activation of free nerve endings by inflammatory mediators such as prostaglandins, neuropeptides, bradykinin, and cytokines.\textsuperscript{29,30} However, there are limited number of studies that investigate the quantity and distribution pattern of nerve fibers in relation to pain. Among the studies included in this systematic review, Grzegorzewski et al.\textsuperscript{27} demonstrated a correlation between pain and the number of nerve fibers, specifically among pediatric patients with cerebral palsy. Saxler et al.\textsuperscript{23} found that patients with OA have significantly greater nerve fiber densities compared with hips without OA. However, Gáspár et al.\textsuperscript{21} found that this high nerve fiber density does not correlate with pre- and postoperative pain or disability. Rabinowicz and Jacqueline\textsuperscript{19} and Haversath et al.\textsuperscript{25} also found that aging is associated with decreased pain fiber expression and increased fibrous tissue formation. These studies demonstrate that pain seen among patients with hip pathology is multifactorial. Greater expression of nerve fibers more significantly affects pain level among pediatric patients. However, due to the negative correlation between age and nerve fiber density as well as fibrous tissue formation over time, adult patients are less affected by nerve fiber density. Instead, increased release of inflammatory mediators as previously described by Schaible et al.\textsuperscript{25} and Perrot and Guilbault et al.\textsuperscript{30} and likely play the major role in pain among patients with OA and FAI.
The studies analyzed in this review suggest that the hip capsule has a complex macroscopic and microscopic innervation that contributes to the overall hip pain found in hip pathologies. While several studies negate the role of nerve fiber quantity in relation to the amount of pain production, evidence from this review strongly suggest that higher nerve fiber expression does not correlate with painful hip pathology, particularly in adults. However, this systematic review was unable to determine the effects of nerve fiber manipulation on pain regardless of the number (or density) of nerve fibers. Nerve tension via stretch has been shown to cause pain, which is a likely contributing factor in the pain perceived due to joint effusion with capsular stretch. In addition to direct fiber transection from capsular incisions (interportal, T, H, or L capsulotomies), a separate similar mechanical stimulus may occur with capsular repair instrument penetration or suture strangulation. Capsular adhesions (e.g., capsulolabral scar, peripheral compartment adhesions to the femoral neck, or overlying musculotenindinous units [iliopsoas, rectus femoris, glutaeus minimus, iliacus, piriformis]) also may induce a nerve stretch instigating pain. Capsular deficiency (and the subsequent femoral head instability) may similarly stimulate capsular nerve fibers, causing pain. Thus, it is clear that capsular management plays a role in pre- and postoperative pain.

Limitations
There are limitations to this review. First, the heterogeneity of the included studies including study subjects (i.e., cadaveric vs in vivo) and methods used to evaluate microscopic innervation (types of staining) limited direct comparisons of results. Second, the included clinical studies were primarily retrospective level IV evidence. In addition, we sought to investigate the hip capsule in hip-preservation surgery subjects. Unfortunately, there was limited evidence (1 study, 11 subjects) in FAI syndrome and dysplasia (1 study, 7 subjects) in adults. Further, it is possible that our stringent search protocol and limiters may have excluded other relevant studies on this topic, including those published in languages other than English.

Conclusions
The hip capsule has a complex macroscopic and microscopic innervation pattern with varying nerve fiber expression from at least 6 peripheral nerves. OA is associated with a greater expression of nerve fibers, although nerve fiber expression does not correlate with painful pathology.

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Appendix Table 1. Search Strategy of Relevant Literature

- Search 1: (capsule[Title/Abstract] OR hip osteoarthritis[Title/Abstract] OR developmental dysplasia[Title/Abstract] OR femoral acetabular impingement[Title/Abstract]) AND (Hip[All Fields] AND innervation[All Fields]).
- Search 2: (capsule[Title/Abstract] OR hip osteoarthritis[Title/Abstract] OR developmental dysplasia[Title/Abstract] OR femoral acetabular impingement[Title/Abstract]) AND (hip osteoarthritis[All Fields] AND innervation[All Fields]).
- Search 3: (capsule[Title/Abstract] OR hip osteoarthritis[Title/Abstract] OR developmental dysplasia[Title/Abstract] OR femoral acetabular impingement[Title/Abstract]) AND (developmental dysplasia[All Fields] AND innervation[All Fields]).
- Search 4: (capsule[Title/Abstract] OR hip osteoarthritis[Title/Abstract] OR developmental dysplasia[Title/Abstract] OR femoral acetabular impingement[Title/Abstract]) AND (femoral acetabular impingement[All Fields] AND innervation[All Fields]).