The value of the glenohumeral joint cross-sectional area as a morphological parameter of glenohumeral osteoarthritis

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Abstract
Glenohumeral joint (GHJ) space narrowing has been demonstrated to be an important morphologic parameter of glenohumeral osteoarthritis (GHO). However, the morphology of GHJ space is irregular because of degeneration of subchondral bone and articular cartilage. Thus, we devised GHJ cartilage cross-sectional area (GHJCCSA) as a new diagnostic morphological parameter to assess the irregular morphologic change of GHJ. GHJ samples were acquired from 33 patients with GHO and from 33 normal controls without evidence of GHO based on shoulder magnetic resonance imaging. T2-weighted coronal MRIs were collected at the GHJ level for all individuals. GHJCCSA and GHJ cartilage thickness (GHJCT) at the GHJ were measured on MRIs using a graphic measuring system. The GHJCCSA was measured as the whole cartilage cross-sectional area of the GHJ. The average GHJCCSA was 115.28 ± 17.36 mm² in normal individuals and 61.77 ± 13.74 mm² in the GHO group. The mean GHJCT was 2.06 ± 0.35 mm in normal individuals and 1.50 ± 0.28 mm in the GHO group. GHO patients had significantly lower GHJCCSA (P < .001) and GHJCT (P < .001) than normal individuals. Receiver operator characteristics curve analysis revealed that the optimal cutoff score of the GHJCCSA was 82.21 mm², with a sensitivity of 97.0%, a specificity of 97.0%, and an area under the curve of 0.99 (95% CI: 0.97–1.00). Although GHJCCSA and GHJCT were both significantly associated with GHO, the GHJCCSA was a more sensitive measurement parameter.

Abbreviations: AUC = area under the curve, CI = confidence interval, GHJ = glenohumeral joint, GHJCCSA = glenohumeral joint cartilage cross-sectional area, GHJCT = glenohumeral joint cartilage space thickness, GHO = glenohumeral osteoarthritis, ROC = receiver operating characteristic, S-MRI = shoulder magnetic resonance imaging.

Keywords: cartilage, cross-sectional area, diagnosis, glenohumeral joint, glenohumeral osteoarthritis, thickness

1. Introduction
Glenohumeral osteoarthritis (GHO) is defined by degeneration of subchondral bone and articular cartilage with narrowing of the glenohumeral joint (GHJ).[1–3] GHO causes functional limitation, disability, and pain, with an estimated prevalence of 4%. It is commonly understood that the GHJ space remains unchanged until the initiation of GHO, at which point joint space narrowing and progressive degenerative changes will occur.[4–6] Only a few trials have been performed to evaluate the predictable value of shoulder magnetic resonance imaging (S-MRI) findings in diagnosing symptomatic GHO. Previous researches have evaluated the GHJ space narrowing using a single measurement called GHJ cartilage space thickness (GHJCT) at the approximate “halfway” or “middle” of the GHJ.[7–9] Kircher et al have demonstrated that the GHJCT decreases with increasing age.[9] This age-dependent joint space narrowing is enhanced in patients with GHO. However, an irregular osteophyte formation can occur anywhere. Therefore, measurement inaccuracy could occur. It might be worthy to reconsider the predictable value of S-MRI findings obtained from a turbo spin echo coronal S-MRI in the diagnosis of symptomatic GHO.[10] To analyze irregular narrowing of the GHJ, we propose that the GHJ cartilage cross-sectional area (GHJCCSA) is a new diagnostic morphological parameter. Compared to the GHJCT, the GHJCCSA does not suffer from mistake of measurement because the GHJCCSA measures the whole cross-sectional area of the GHJ. We hypothesize that the GHJCCSA is a main morphologic parameter in GHO diagnosis. Therefore, the objective of this study was to used S-MRI images to compare GHJCCSA and GHJCT between GHO patients and control individuals.
2. Methods and Material

2.1. Patients

The retrospective research material used to support the findings of this research were approved by Catholic Kwandong Institutional Review Board. (IRB number: IS19RISI0061).

Individuals who visited the shoulder orthopedic clinic with shoulder pain from July 2017 to April 2019 and had taken S-MRI within 12 months of the last visit were reviewed retrospectively. Inclusion criteria of the GHO group were: a history of pain and tenderness in the GHJ; crepitation with movement of the joint that was often palpated anteriorly; persistent symptom (patients should be seen in follow-up at 6 to 8 weeks to reevaluate range of motion). We excluded subjects if patients had any of the following histories: inflammatory arthritis; humerus bone fracture; history of shoulder infection; acute clavicle fracture; or any history of shoulder surgery.

A total of 33 patients was confirmed by 2 board-certified experienced musculoskeletal radiologists. There were 16 (48.48%) men and 17 (51.52%) women with an average age of 60.55 ± 10.03 years (range, 43 to 80 years) (Table 1). To compare results between individuals with and without GHO, normal subjects were enrolled. Individuals who wanted to take S-MRI for the exact diagnosis of shoulder pain without evidence of GHO were enrolled for the normal group. In the normal group, 33 individuals (12 men and 21 women) were enrolled. Their average age was 58.73 ± 8.81 years (range, 40 to 83 years).

2.2. Imaging parameters

Shoulder assessment was performed using a 3T S-MRI system (Magnetom Skyra, Siemens, Medical Solutions, Germany) and a 3T Ingina scanners (Philips Healthcare, Eindhoven, Netherlands). For all S-MRI examinations, oblique coronal fat suppressed T2-weighted images were obtained with a slice thickness of 3 mm, slice gap of 0.9 mm, repetition time 4010-ms of/echo time 76-ms, 150 × 150 cm field of view, 512 × 256 matrix, and > 3 ETL.

2.3. Image analysis

GHJCCSA and GHJCT measurements were performed by one pain specialist who was blinded to the classification of shoulders. Oblique coronal T2-weighted S-MRIs were obtained at the GHJ. GHJCCSA and GHJCT were measured on S-MRI using an image analysis program (INFINITT PACS; Infinitt Medical Solutions, Incheon, Seo-gu, Republic of Korea) (Figs. 1, 2A and B). The GHJCT was measured at the narrowest GHJ. The GHJCCSA was measured as the whole cross-sectional area of the cartilage of GHJ.

2.4. Statistical analysis

Student’s t test was used to compare GHJCCSA and GHJCT between GHO and control groups. Receiver operating characteristic curve was applied to assess diagnostic values of GHJCCSA and GHJCT. Area under the curve (AUC) with 95% confidence intervals (CIs), cutoff points, sensitivity, and specificity were obtained (IBM/SPSS Inc., Chicago, IL).

3. Results

There were 15 right shoulders and 18 left shoulders in the control group. The average GHJCCSA was 115.28 ± 17.36 mm² in the control group and 61.77 ± 13.74 mm² in the GHO group. There were 18 right shoulders and 15 left shoulders in the GHO group. The mean GHJCT was 2.06 ± 0.35 mm in the normal group and 1.50 ± 0.28 mm in the GHO group. GHO patients had significantly lower GHJCCSA (P < .001) and GHJCT (P < .001) than normal subjects (Table 1). Regarding the diagnostic accuracy of both GHJCCSA and GHJCT as predictors
of GHO, receiver operating characteristic curves revealed that the best cutoff score of the GHJCT was 1.82 mm, with a sensitivity of 90.9%, a specificity of 87.9%, and an AUC of 0.92 (95% CI: 0.85-0.99) (Table 2, Fig. 3). The best cutoff point of the GHJCCSA was 82.21 mm², with a sensitivity of 97.0%, a specificity of 97.0%, and an AUC of 0.99 (95% CI: 0.97-1.00) (Table 3, Fig. 2).

4. Discussion

The GHJ is one of the most common affected large joints following the hip and knee. Although the true prevalence of GHO is difficult to analyze, population-based researches have reported that about 20% of elderly populations have radiographic evidence of GHO. Major risk factors for GHO include obesity and female sex. Secondary causes of GHO include crystalline or infectious arthropathy, avascular necrosis, prior trauma such as surgery and dislocation. Arthrogenic factors include presence of bony defects of either the humerus or glenoid, presence of a rotator cuff tear, and age at time of dislocation. Physical examination should be attempted to identify their etiologies within the shoulder such as bursitis and tendinosis and pathology of pain outside the shoulder should be excluded. Pain radiating down the arm and neck pain with provocative maneuvers such as Spurling’s test can rule out a cervical origin problem. A thorough neurovascular and neck examination should always be performed to diagnose properly.

The first step of treatment of primary GHO is nonoperative conservative management. Operative management of GHO is recommended for patients who have failed the first step of treatment. In elderly patients, total shoulder arthroplasty is a highly successful and reliable procedure. However, concerns regarding decreased activity levels and implant longevity have led orthopedic surgeons to pursue joint-preserving procedures in more active and younger patients. Radiograph is the first choice for staging and diagnosing GHO. As described above, posterior glenoid wear and joint space narrowing are commonly found. The presence of osteophytes and subchondral sclerosis from the humeral head is also an anticipated finding. Advanced imaging modalities are frequently necessary for exact diagnosis because they can provide important information for

| GHJCT (mm) | Sensitivity (%) | Specificity (%) |
|-----------|----------------|----------------|
| 1.11      | 9.1            | 100            |
| 1.23      | 24.2           | 93.9           |
| 1.45      | 42.4           | 93.9           |
| 1.82 *    | 90.9           | 87.9           |
| 1.84      | 93.9           | 84.8           |
| 1.99      | 100            | 60.6           |

GHJCT = glenohumeral joint cartilage thickness.
*The best cutoff point on the receiver operating characteristic curve.
staging disease progression, identification of concomitant rotator cuff or labral pathology, and preoperative planning.[5,8,11] Only a few trials have evaluated the predictable value of S-MRI findings in diagnosing symptomatic GHO. Schleich et al have provided normative information for comparison of GHJ cartilage changes in various pathological disorders such as early GHO.[27] Kircher et al have demonstrated that GHJCT decreases with increasing age. This age-dependent joint space narrowing is enhanced in patients with GHO.[9]

However, previous articles assessed the GHJ space narrowing using a single measurement called GHJCT at the approximate “single halfway” of the GHJ. Therefore, a measurement error frequently occurs. We think that it might be worthy to reconsider the predictable value of S-MRI findings obtained from a coronal, turbo spin echo T2-weighted, fat-suppressed S-MRI in the diagnosis of symptomatic GHO.

To analyze narrowing of the GHJ, we devised the GHJCCSA as a new morphological parameter. In contrast to the GHJCT, the GHJCCSA is not influenced by this measurement bias because the GHJCCSA measures the cross-sectional area of the whole GHJ. Eventually, we found that the GHJCCSA was better than the GHJCT as a diagnostic parameter of GHO. In the current original research, we found that the GHJCCSA had a sensitivity of 97.0%, a specificity of 97.0%, and an AUC of 0.99 (95% CI: 0.97–1.00) to predict GHO. In contrast, the GHJCT had a sensitivity of 90.9%, a specificity of 87.9%, and an AUC of 0.92 (95% CI: 0.84–0.99). These findings suggest that the GHJCCSA is a better predictor of GHO than the GHJCT. We believe that our results can be used to improve the quality of diagnosis of the GHO.

This study has some limitations. There are several isolated GHJ pathology in the symptomatic shoulder such as rotator cuff tear arthropathy. However, we focused on GHO. Second, some alternative methods such as ultrasound, plain X-ray, arthrocentesis have been shown to be effective at discriminating GHO.[27] Especially, ultrasound provides real-time imaging.[26–33] However, we only measured GHJCCSA and GHJCT on S-MRI. Third, there might be some bias associated with measuring GHJCCSA and GHJCT on S-MRI. Even though we tried to check these diagnostic parameters in the oblique coronal T2-weighted image that best showed the GHJ, coronal images could be irregular.

5. Conclusion

Although GHJCCSA and GHJCT are both significantly associated with GHO, GHJCCSA is a more sensitive diagnostic parameter for GHO than GHJCT. We demonstrated that the most suitable cutoff value of the GHJCCSA was 82.21 mm², with sensitivity and specificity of 97.0% both. The optimal cutoff point of the GHJCT was 1.82 mm, with a sensitivity of 90.9% and a specificity of 87.9%. Thus, physicians should carefully evaluate the GHJCCSA rather than the GHJCT when assessing patients with GHO.

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