Arthroscopic medial release for medial compartment osteoarthritis of the knee

THE RESULT OF A SINGLE SURGEON SERIES WITH A MINIMUM FOLLOW-UP OF FOUR YEARS

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The outcome of arthroscopic medial release of 255 knees in 173 patients for varying grades of osteoarthritis involving the medial compartment is reported. All operations were performed by a single surgeon between January 2001 and May 2003. The Knee Society score for pain and the patient’s subjective satisfaction were used for the outcome evaluation. Overall, satisfactory outcome was reported for 197 knees (77.3%) and the mean Knee Society score for pain improved from 17.6 (95% confidence interval, 16.7 to 18.5), pre-operatively to 39.4 (95% confidence interval, 37.9 to 41.1) (p < 0.001). There were minor manageable complications of persistent effusion in 16 knees and prolonged wound discomfort in 11. In total, 15 of the 21 knees with poor results were converted to total knee replacements and two other patients (three knees) were offered this option after a mean period of 16 months.

Based on these observations arthroscopic medial release is an effective treatment for osteoarthritis of the medial compartment of the knee joint and can be expected to reduce the pain in the majority of patients for at least four years post-operatively.

Arthroscopic techniques for the management of degenerative conditions of the knee include joint lavage, debridement, abrasion arthroplasty and microfracture. Nevertheless, their effectiveness has not been proven in prospective trials and the mechanism by which they improve the course of degenerative conditions of the knee has not been established. Therapeutic arthroscopy performed in the osteoarthritic knee also remains a source of controversy because of a lack of a clear consensus on the indications and surgical technique, as well as the unpredictability of the outcome. In 2007, Siparsky et al performed a retrospective, evidence-based review of the literature on the arthroscopic treatment of osteoarthritis of the knee and found limited support for its use in this condition.

Medial release as part of the surgical treatment for medial compartment osteoarthritis of the knee has been reported occasionally since 1940. In 1960, Loeffler described a procedure which included resection of the medial collateral ligament for patients with both early and late degenerative changes of the medial compartment. Success rates ranging from 72.5% to 90% have been reported following this procedure. In 2001, Leon, Blanco and Guthrie reported an arthroscopic intervention they referred to as arthroscopic decompressive medial release. They divided the medial third of the capsule and the medial collateral ligament (MCL) transversely 1.5 cm proximal to the medial meniscus to unload the medial compartment and achieved satisfactory results in a small group of 38 patients. In 2004, Moriya et al reported the clinical outcomes of a similar arthroscopic concept of posterior-medial release. The medial compartment was released by separating the medial capsule and MCL from the medial tibial plateau after meniscectomy. In their opinion, relatively advanced osteoarthritis of the knee for which arthroscopic debridement has conventionally been contraindicated, could be treated with this procedure in carefully selected patients. However, none of these salvage procedures have become popular.

During arthroscopic debridement in patients with medial compartment osteoarthritis the author has often observed unusual degenerative changes affecting the medial femoral condyle with adjacent inflammatory synovial tissue and a medial plica. The common presence of a plica in these circumstances and its positive correlation in severity to the pathological changes seen on the medial femoral condyle encouraged its resection with an associated medial release. We hypothesised that the eradication of the interplay between
the inflamed medial plica and the medial femoral condyle would be beneficial. Subsequently, the procedure has been refined. During the late 1990s it became referred to as arthroscopic medial release and was standardised. In this retrospective study, the details of this procedure and its clinical results are reported.

Patients and Methods
Between January 2001 and May 2003, 215 patients underwent arthroscopic surgery for osteoarthritis involving the knee at the Tzu-Chi Dalin General Hospital. Within this group of patients, 267 arthroscopic medial releases were performed in 182 patients (153 female and 29 male) with medial compartment osteoarthritis. The mean age of the patients was 61 years (38 to 75). A total of 85 patients received bilateral treatment: 79 were treated with simultaneous operations and six were treated after an interval of six to nine months. The procedure was offered to patients with osteoarthritis of any stage that was confined mainly to the medial compartment. All patients had been treated conservatively for more than one year without success prior to the arthroscopic treatment. The exclusion criteria were the presence of modified Outerbridge\textsuperscript{27} grade III or IV osteoarthritis of the lateral or patellofemoral compartments, unstable meniscal fragments, instability due to previous ligament injury, or post-traumatic osteoarthritis. Age, radiological grading according to Kellgren and Lawrence,\textsuperscript{28} and the pain domain of the Knee Society scoring system\textsuperscript{29} (KSS) were recorded pre-operatively.

The anteroposterior standing radiographs of these patients were evaluated for the severity of medial compartment osteoarthritis. The four-grade classification of Kellgren and Lawrence\textsuperscript{28} was used, and is as follows: grade 1, doubtful narrowing of joint space and possible osteophytic lipping; grade 2, definite osteophytes and possible narrowing of joint space; grade 3, moderate multiple osteophytes, definite narrowing of joint space, some sclerosis and possible deformity of bone ends; grade 4, large osteophytes, marked narrowing of joint space, severe sclerosis, and definite deformity of bone ends. The radiological grading was performed by the author, who also performed all of the procedures.

Surgical procedure. All procedures were performed under spinal or general anaesthesia with the use of a tourniquet.

Routine arthroscopic examination was performed through an inferolateral portal and the condition of articular cartilage, menisci and ligaments was evaluated. If osteoarthritis was found to involve the lateral or patellofemoral compartments, unstable meniscal fragments, or ligamentous deficiency was identified, the patient was excluded from the study.

The condition of the medial plica was studied in detail. The type and grade of severity of the plica according to its gross appearance, was classified as previously described.\textsuperscript{26} In summary, a type A plica has no direct contact with the medial femoral condyle, a type B rides onto but not beyond half the width of the medial femoral condyle, and a type C plica extends over more than half of the width of the medial femoral condyle. A grade I plica looks like a membrane, its margin is semi-transparent and is soft in consistency when palpated by a probe. The grade II plica is not transparent but is hypertrophied and thickened while retaining a soft consistency. A grade III plica looks like a fibrotic band and is thicker than a grade II but is elastic in consistency. A grade IV plica exhibits the additional sign of wear (caused by abrasion with the femoral condyle) as well as looking fibrotic. Its margin is frail and fibrillated. A grade V plica represents an inflamed version of a grade IV, with focal synovitis adjacent to it. In most cases synovectomy is required in this grade to enable clear visualisation of the plica. The observation for the classification of the plica is performed through the inferolateral portal with the knee in full extension.

First-stage debridement. The inflammatory tissue occupying the inferomedial region of the patella, including the ligamentum mucosum, fibrotic or inflamed synovium, capsule and distal part of the medial plica, was removed using a power shaver passed through a superolateral portal, with the debridement extended as medially as possible to eradicate all the inflammatory tissue found in this space (Fig. 1).

Release of the medial gutter. The obliterated space of the medial gutter was released by resection of the fibrotic synovium, capsule and proximal part of the medial plica little by little using either an electrical shaver or a punch. When complete, the tightened medial patellofemoral joint had been released and the medial gutter cleared of any fibrotic or inflammatory tissue. In most cases, this stage of the procedure could be performed through the superolateral portal.

**Fig. 1**

Intra-operative photograph showing that the debridement is carried out through the superolateral portal as medially as possible to eradicate all the inflammatory tissue (IT) occupying the inferomedial region of the patella (P).
Intra-operative photograph showing that the obliterated space of the medial gutter (MG) is gradually released by resection of the fibrotic synovium, capsule, and proximal part of the medial plica. In most cases this stage could be performed through the superolateral portal.

(Fig. 2). However, in some difficult cases it was more practical to use an inferomedial portal.

**Final assessment of the release.** The edge of the resection at the synovio-meniscal junction was smoothed to avoid impingement against the medial femoral condyle (C) when the knee joint is moves.

(Fig. 3). The adequacy of the medial release was confirmed by passing the arthroscope under the patella and verifying that the previously tightly closed medial patellofemoral joint space could be easily opened and the medial retinaculum visualised when the knee was held in full extension (Fig. 4).

The target of the capsular release was layer III, the so-called ‘true capsule’ of the three-layer medial structure described by Warren and Marshall. As shown in Figure 5, the capsulectomy was extended superiorly to the midline of the suprapatellar pouch and inferiorly to the upper margin of the medial meniscus. Anteriorly, it was extended to the medial margin of the patella, and posteriorly it was continued until a portion of the conjoined part of layer II and III was removed and the gracilis tendon could be seen. Only the deep medial ligament was severed by this procedure, so that medial stability was not disturbed.

**Synovectomy and chondroplasty.** Any focal synovitis or loose chondral flaps on the cartilaginous surface were debrided as conventional arthroscopic treatment for osteoarthritis of the knee, but no bony procedure, such as drilling or microfracture, was performed. At the end of the procedure, thorough irrigation was performed to remove any debris from the knee.

**Post-operative management.** A suction drain was used for 24 hours. The leg was protected by an elastic bandage for one week. The patient was allowed to mobilise freely as comfort permitted without restriction on weight-bearing. The patient was discharged the day after the operation. Active range of movement and quadriceps exercises were encouraged at home.

**Follow-up and evaluation of outcome.** The patients were reviewed monthly for three months. Thereafter, they
returned annually for assessment. The 50-point pain domain of the KSS was used to evaluate the pain both pre- and post-operatively, where 50 points indicate no pain and 0 points indicate severe pain. Subjective satisfaction was assessed by direct questioning using a categorical scale prepared for this study (Table I). The outcome was regarded as satisfactory if subjective satisfaction was rated as ‘excellent’ or ‘good’. The inquiry into subjective satisfaction and the evaluation of KSS were conducted by nursing specialists. All investigations focused on individual knees in bilateral cases.

**Statistical evaluation.** Results were presented as means, SD and 95% confidence intervals (CI) as appropriate. Comparisons were made using one-way analysis of variance (ANOVA) to detect differences in the distribution of patient age and the type and grade of severity of the medial plica in each grade of osteoarthritis. Comparison of pre- and post-operative Knee Society pain scores was performed using the paired t-test. Post-operative Knee Society scores > 40 and an increase in the score of > 20 were defined as a successful outcome. Logistic regression tests were used to evaluate the correlation of size of the plica, the pathological severity of the plica and the severity of osteoarthritis with the patient’s age. A p-value < 0.05 was considered to be statistically significant. All statistical analysis was carried out using the statistical discovery software, JMP (version 5.0.1.2, SAS Institute Inc., Cary, North Carolina).

**Results** All the knees in the study were found to have a medial plica. The type of plica and severity of the osteoarthritis were examined in relation to the mean age of the patient. The size of the plica was negatively correlated with the patient’s age (r = 0.29, p < 0.0001). The mean age and distribution of different grades of plica and the type and severity for each grade of osteoarthritis are shown in Table II. The severity of the plica (r = 0.23, p = 0.0018) and the severity of osteoarthritis (r = 0.58, p < 0.0001) were positively correlated with the patient’s age.

The mean follow-up period was 5.3 years (4 to 6.5). None of the nine grade I osteoarthritic knees were lost to follow-up; two of 90 knees (2.2%) with grade II osteoarthritis in two patients, six of 135 knees (4.4%) with grade III osteoarthritis in four patients, and four of 33 knees (12.1%) with grade IV osteoarthritis in three patients were lost to follow-up. There were 255 knees in 173 patients available with more than four years follow-up.

**Post-operative complications.** Of the 255 available knees, 16 (6.3%) in 12 patients experienced persistent effusion and discomfort for more than two weeks after the procedure. After initial treatment, including rest, compression bandaging and anti-inflammatory medication, the effusion subsided spontaneously within two weeks in 11 knees. The other five knees in four patients required aspiration of the knee after four weeks.
Most patients reported some pain, catching or instability during the first post-operative month. This resolved spontaneously. In 11 knees (4.3%) in nine patients there was discomfort due to wound irritation for more than three months. In five knees in four patients this settled with routine treatment within three months. A hydrocortisone injection was required in three other knees and two further patients (three knees) had persistent pain and instability despite this treatment. Their symptoms settled, however, following arthroscopy and adhesiolysis. A fibrotic band and tightness of the medial capsule was found in these knees.

**Outcome.** Of the 255 knees available for final review the subjective assessment was satisfactory in 197 (77.3%). The mean KSS for pain improved from 17.6 (95% CI 16.7 to 18.5) pre-operatively to 39.4 (95% CI 37.9 to 41.1) at final follow-up. The improvement was statistically significant in all groups (Table III). For knees with an excellent outcome, the mean post-operative KSS for pain was 50, with a mean increase of 33.1 (95% CI 31.8 to 34.4), for knees with a good outcome it was 41.8 (95% CI 41.2 to 42.4), with a mean increase of 24.9 (95% CI 23.4 to 26.4), and for knees with a fair outcome it was 26.2 (95% CI 24.6 to 27.8), with a mean increase of 1.9 (95% CI 0.9 to 2.9). However, for knees with a poor outcome, the mean KSS for pain was 5.2 (95% CI 3.1 to 7.3) pre-operatively with a mean reduction of 6.9 (95% CI 4.9 to 8.9) post-operatively. Applying a post-operative KSS for pain of > 40 or the increase of scores of > 20 as the criteria for treatment success, for knees with grade I osteoarthritis the success rate was 100% (nine knees), for grade II, 87.5% (77 of 88 knees); for grade III, 62.8% (81 of 129 knees), and for grade IV, 41.2% (12 of
29 knees). Applying this evaluation with the KSS for pain the success rate of the whole series was 70.2% (179 of 255 knees).

Of the 21 knees with poor results three occurred in two patients who received a second arthroscopy and adhesiolysis. In total, 15 of the 21 knees with poor results were converted to total knee replacement after a mean period of 16 months (13 to 25). The other two patients (three knees) were also offered knee replacement. All the patients with poor results had grade III and grade IV osteoarthritic changes.

Discussion

Arthroscopic release for medial compartment osteoarthritis of the knee gave a satisfactory subjective outcome rate of 77.3%, and the success rate evaluated by KSS for pain was 70.2%. Generally, the results were best in the patients with the least advanced arthritic changes. Even in grade IV knees, 44.8% (41.2% by KSS for pain) of the patients were satisfied with this procedure. Although this outcome is not as good as that provided by total knee replacement, it still offers a benefit for many patients because of its simplicity and low morbidity.

The outcomes of commonly-used arthroscopic procedures for osteoarthritis of the knee, including joint lavage, debridement, abrasion arthroplasty and microfracture, are inconsistent. In most patients, short-term symptomatic relief can be expected with arthroscopic lavage and debridement. In 2002, Moseley et al8 performed a double-blind, randomised, placebo-controlled trial to compare the effectiveness of arthroscopic lavage and arthroscopic debridement versus a sham procedure and found that arthroscopy provided subjective pain relief via a placebo effect. Abrasion chondroplasty was popularised in the late 1980s. Rand,10 comparing arthroscopic partial meniscectomy with limited debridement and arthroscopic abrasion arthroplasty in patients with osteoarthritis, found that abrasion arthroplasty offered little benefit over partial meniscectomy and debridement. Concern exists over the durability of the fibrocartilage repair tissue in subchondral penetration procedures, and over thermal damage to subchondral bone and adjacent normal articular cartilage during this procedure. Microfracture is a modification of chondroplasty, but a deterioration of the results has been reported starting 18 months after surgery.9

The concept of medial release procedures is similar. The main purpose of Loeffler’s operation is to release the tight medial soft-tissue structures and excise hypertrophied synovial tissue, which may cause pain in degenerative knees. Leon et al24 and Moriya et al25 claim that their procedures of arthroscopic release unload the medial compartment by releasing the medial capsule and medial collateral ligament, which may permit a reduced adduction moment and a reduction of the external rotation restraint in extension found in more severely osteoarthritic knees.

The main rationale for our arthroscopic medial release is to eradicate the abrasion or impingement phenomenon between the tight, fibrotic and hypertrophied medial plica and the adjacent medial femoral condyle, which has been described in previous studies in patients with osteoarthritic knees.26,31,32 It has been reported that certain neuropeptides are responsible for the neural transmission of pain in the synovium and capsule of osteoarthritic knees.33 This may account for most of our patients obtaining early, rapid pain relief. This effect has also been reported by Ikeuchi, Takanashi and Tani,34 who successfully treated 19 osteoarthritic knees with localised synovial hypertrophy in the anteromedial compartment by arthroscopic partial synovectomy. In a one-year longitudinal arthroscopic study in 422 patients, Ayral et al35 found that abnormalities of the medial synovium were a common feature of painful medial osteoarthritis of the knee associated with more marked medial chondropathy. In addition, they suggested that an inflammatory aspect of the medial perimeniscal synovium could be related to a subsequent medial degradation. Therefore, the eradication of all the inflammatory synovium in this area in our procedure might have reduced or arrested the degenerative process in the medial compartment.

The prevalence of a medial plica varies from 19% to 95%.36,37 The universal presence of the medial plica in our patients with osteoarthritic knees, and the relationship between its size and gross appearance and the patients’ age, are in keeping with a previous study.26 Whether the medial plica plays a role in the pathogenesis of the osteoarthritis is worthy of further investigation.

A limitation of this study was the lack of radiological follow-up evaluation to correlate with the clinical outcome. The lack of independence in the grading system and no measures of reproducibility were also drawbacks of this study.

In conclusion, this experience with arthroscopic medial release suggests that it is not a safe release of the knee over a period of at least four years. Whether it is able to modify the disease process warrants more prospective studies, including imaging evaluation. The high incidence of a medial plica in patients with medial compartment degenerative disease also justifies further investigations for its role of this structure in the pathogenesis of osteoarthritis.

Supplementary Material

A further opinion by Mr T. Wilton is available with the electronic version of this article on our website at www.jbjs.org.uk

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References


